Carbon uptake and biogeochemical change in the Southern Ocean, south of Tasmania
Pardo, Paula Conde1, Tilbrook, Bronte1,2, Langlais, Clothilde2, Trull, Thomas William1,2, Rintoul, Stephen Rich1,2

1Antarctic Climate and Ecosystem Cooperative Research Centre, University of Tasmania, Hobart, Australia
2Climate Science Centre, CSIRO Oceans and Atmosphere, Hobart, Australia

Correspondence to: Paula C. Pardo (paula.condepardo@csiro.au)

Abstract. Biogeochemical change in the water masses of the Southern Ocean, south of Tasmania, was assessed for the 16-year period between 1995 and 2011 using data from 4 summer repeats of the WOCE/JGOFS/CLIVAR/GO-SHIP SR03 hydrographic section (at ~140°E). Changes in temperature, salinity, oxygen, and nutrients were used to disentangle the effect of solubility, biology, circulation and anthropogenic carbon (CANT) uptake on the variability of dissolved inorganic carbon (DIC) for 8 water mass layers defined by neutral surfaces (γn). CANT was estimated using an improved back-calculation method. Warming (~0.0352 ± 0.0170 °C yr¹) of Subtropical Central Water (STCW) and Antarctic Surface Water (AASW) layers decreased their gas solubility, and accordingly DIC concentrations increased less rapidly than expected from equilibration with rising atmospheric CO₂ (~0.86 ± 0.16 μmol kg⁻¹ yr⁻¹ versus ~ 1 ± 0.12 μmol kg⁻¹ yr⁻¹). An increase in apparent oxygen utilisation (AOU) occurred in these layers due to either remineralization of organic matter or intensification of upwelling. The range of estimates for the increases of CANT were 0.71 ± 0.08 to 0.93 ± 0.08 μmol kg⁻¹ yr⁻¹ for STCW and 0.35 ± 0.14 to 0.65 ± 0.21 μmol kg⁻¹ yr⁻¹ for AASW, with the lower values in each water mass obtained by assigning all the AOU change to remineralization. DIC increases in the Sub-Antarctic Mode Water (SAMW, 1.10 ± 0.14 μmol kg⁻¹ yr⁻¹) and Antarctic Intermediate Water (AAIW, 0.40 ± 0.15 μmol kg⁻¹ yr⁻¹) layers were similar to the calculated CANT trends. For SAMW, the CANT increase tracked rising atmospheric CO₂. As a consequence of the general DIC increase, decreases in total pH (pHT) and aragonite saturation (ΩAR) were found in most water masses, with the upper ocean and the SAMW layer presenting the largest trends for pHT decrease (~ -0.0031 ± 0.0004 yr⁻¹). DIC increases in deep and bottom layers (~ 0.24 ± 0.04 μmol kg⁻¹ yr⁻¹) resulted from the advection of old deep waters to resupply increased upwelling, as corroborated by increasing silicate (~ 0.21 ± 0.07 μmol kg⁻¹ yr⁻¹), which also reached the upper layers near the Antarctic Divergence (~ 0.36 ± 0.06 μmol kg⁻¹ yr⁻¹) and was accompanied by an increase in salinity. The observed changes in DIC over the 16-year span caused a shoaling (~340 m) of the aragonite saturation depth (ASD, ΩAR = 1) within Upper Circumpolar Deep Water that followed the upwelling path of this layer. From all our results, we conclude a scenario of increased transport of deep waters into the section and enhanced upwelling at high latitudes for the period between 1995 and 2011 linked to strong westerly winds. Although enhanced upwelling lowered the capacity of the AASW layer to uptake atmospheric CO₂, it did not limit that of the newly forming SAMW and AAIW, which exhibited CANT storage rates (~ 0.41 ± 0.20 mol m⁻² yr⁻¹) twice that of the upper layers.
1 Introduction.

The Southern Ocean is a key region in terms of climate change and climate variability, influencing the Meridional Overturning Circulation and therefore modulating the global circulation and oceanic biogeochemical cycles (Sarmiento et al., 1998, 2004; Orr et al., 2005). Deep waters, formed in the North Atlantic, spread south and enter the Southern Ocean, where they mix with deep layers of the Antarctic Circumpolar Current (ACC) and ultimately upwell between the Southern ACC Front and the Polar Front. The upwelled waters are eventually transformed into bottom, intermediate and mode waters, which are exported from the Southern Ocean to ventilate the thermocline and bottom layers of the major ocean basins. Some of the Southern Ocean waters subducted into the ocean interior return to the North Atlantic to balance the southward flux of North Atlantic Deep Water (Speer et al., 2000; Lumpkin and Speer, 2007; Iudicone et al., 2008).

Within the eastward flow of the ACC major water exchange between the three ocean basins takes place. The circumpolar path of the ACC consists of various narrow jets associated with sharp fronts that separate waters with different characteristics (Orsi et al., 1995; Belkin and Gordon, 1996). These jets can reach deep layers and often meander, intensify, merge and split, conditioned by the topography of the ocean floor, the stratification of the ACC, and atmospheric variability (Moore et al., 1999; Sokolov and Rintoul, 2002, 2009; Peña-Molino et al., 2014). Movements in the jets enhance cross-stream transports and mesoscale activity that can result in local changes of water mass properties and this may complicate the computation of long-term changes in water mass properties (Rintoul and Bullister, 1999; Sallée et al., 2008; Peña-Molino et al., 2014).

Water mass formation and ventilation transport heat, salt, and dissolved gases from the atmosphere to the ocean interior and other basins (Sarmiento et al., 2004), with the Southern Ocean contributing ~40% to the anthropogenic CO$_2$ (C$_{ANT}$) inventory of the ocean (Sabine et al., 2004; Gruber et al., 2009; Khatiwala et al., 2009). Circulation and biological processes drive the redistribution of dissolved inorganic carbon (DIC) that ultimately affects the capacity of the waters to uptake more CO$_2$. The uptake of CO$_2$ by the Southern Ocean presents strong spatiotemporal variability (Lenton et al, 2013) and this can lead to conflicting results for observational studies, models and atmospheric inversions depending on the methodology used (Verdy et al., 2007; Lenton et al., 2012; Fay et al., 2014). Quantifying long-term changes in the carbon system is difficult due to the scarcity of data (Lenton et al., 2012; Kouketsu and Murata, 2014; Fay et al., 2014) and the influence of biological processes.

Notably, long term trends in C$_{ANT}$ concentration are difficult to estimate due to its small signal (~3%) with respect to that of DIC in the ocean.

Once CO$_2$ dissolves in the ocean (DIC) it begins the process of ocean acidification, i.e., decreases the pH and the saturation state of calcium carbonate (CaCO$_3$) minerals such as calcite and aragonite (Feely et al., 2004; Bates et al., 2014), with potential to disrupt ecosystems and biological processes (Doney et al., 2009).

Numerous studies have documented warming and freshening of deep and bottom layers of the Southern Ocean in recent decades (see reviews by Jacobs, 2006 and van Wijk and Rintoul, 2014). The abyssal waters of the Australian-Antarctic Basin (A-AB) show the greatest freshening in the last 40 years (van Wijk and Rintoul, 2014). This freshening was accompanied by a warming of the deep-bottom layers, leading to a contraction in bottom waters by more than half their volume in the basin.
(Purkey and Johnson, 2012; van Wijk and Rintoul, 2014). Subsurface to intermediate layers have also warmed and freshened south of Australia (Bindoff and Church, 1992; Wong et al., 1999; Aoki et al., 2005). The solubility of gases and its distribution in the ocean depends on the dynamics and properties of water masses, and the thermohaline changes that have occurred south of Tasmania (Fig. 1) have implications in the carbon system of the region.

A reduction of the carbon sink of the Southern Ocean was observed between the 1980s and the early 2000s (Le Quéré et al., 2007; Lovenduski et al., 2008), with more recent studies suggesting a recovery and even intensification of the CO₂ uptake by 2011-2012 (Zickfeld et al., 2008; Fay et al., 2014; Landschützer et al., 2015). DeVries et al. (2017), used a global inverse model to postulate that changes in circulation are responsible for most of the variability in the oceanic CO₂ uptake, with the weakening of the upper-ocean circulation being responsible for the increase in oceanic carbon uptake over the past decade.

Ocean acidification has been observed in the whole Southern Ocean (Lauvset et al., 2015) and locally in the Atlantic and Pacific sectors (Williams et al, 2015; Hauri, et al., 2015). South of Tasmania, McNeil et al., (2001) reported an increase in C<sub>ANT</sub> uptake between 1968 and 1996 for the region 45-50°S. These authors reported, for the first time, C<sub>ANT</sub> accumulation in the AABW and highlighted the importance of the formation of bottom and mode waters as a mechanism for transporting C<sub>ANT</sub> to the ocean. In terms of ocean acidification, we are not aware of any study about trends in ocean acidification in the water masses south of Australia.

Considering the lack of observational estimates for recent biogeochemical changes in the A-AB as well as the large changes in CO₂ uptake and storage suggested by recent atmospheric and surface observations in the Southern Ocean (e.g., Fay et al., 2014; Landschützer et al., 2015) there is a need to provide a full ocean depth observational perspective on how the ocean is changing. The aim of this paper is to provide the first estimates of biogeochemical change in the water masses south of Tasmania, for the period 1995-2011, disentangling the effects that solubility, circulation, biology and C<sub>ANT</sub> uptake have on the variability of DIC. We use data from four summer repeats of the WOCE/JGOFS/CLIVAR/GO-SHIP hydrographic section SR03 (Fig. 1; Table 1), one of the most revisited sections in the Southern Ocean. Trends in oxygen (O₂), nutrients, and the carbon system parameters, i.e., DIC, total alkalinity (TA), anthropogenic carbon (C<sub>ANT</sub>), total pH (pH<sub>T</sub>) and % aragonite saturation (Ω<sub>ar</sub>) were estimated for the period 1995-2011, when both DIC and TA measurements are available. C<sub>ANT</sub> estimates were obtained with a back-calculation method (Pardo et al., 2014). The changes were evaluated in the different water mass layers of the section defined by neutral surfaces (γ<sup>n</sup>, McDougall et al., 1987).

2 Hydrography of the region.

The dynamical structure of the region south of Tasmania (Fig. 1) is characterized by a number of fronts that separate the major water masses of the region (Sokolov and Rintoul 2002, 2007, 2009). At the northern end of the section, the presence of the weak Subtropical Front (STF, Fig. 1) separates warm, salty subtropical surface waters from cooler and fresher sub-Antarctic surface waters (Deacon, 1937). The northern end of the section is a complex mixing zone where waters transported down the east coast of Tasmania in a series of mesoscale eddies from the East Australian Current (EAC, Fig. 1) mix into the Subantarctic
Zone, and also meet Zeehan Current (ZC, Fig. 1) waters transported down the west coast of Tasmania (Boland and Church 1981; Baines et al., 1983; Speich et al., 2002; Davis, 2005; Ridgway et al., 2007; Sloyan et al., 2016). The EAC transported south of Tasmania forms a zonal jet towards the southeast Indian Ocean known as the Tasman Outflow that reaches the bottom of the Tasman slope and that is maintained all year round (Rintoul and Bullister, 1999; Ridgway et al., 2007). The encounter between these currents presents high variability at the northern end of the section (Fig. 1).

Farther south (Fig. 1), the Sub-Antarctic Front (SAF) and the Polar Front (PF) are regions of maximum transport in the ACC (Rintoul and Bullister, 1999; Sokolov and Rintoul 2002). North of the SAF, deep winter convection generates Sub-Antarctic Mode Water (SAMW), a relatively uniform water mass that occupies subsurface layers down to ~600m deep (McCartney, 1977; Rintoul and Bullister, 1999). SAMW constitutes the main part of the upper limb of the MOC, ventilating the thermocline of all the ocean basins (e.g., Speer et al., 2000; Sloyan and Rintoul, 2001). The SAF coincides with the deepening of the salinity minimum at intermediate depths (Whitworth and Nowlin, 1987), which is the signature of the Antarctic Intermediate Water (AAIW). The AAIW underlies the SAMW and also ventilates the global thermocline layers of the ocean. It is mainly formed in the southeast Pacific and it is continuously transformed on its way to the region south of Tasmania (Hanawa and Talley, 2001). South of the SAF, colder and fresher Antarctic Surface Water (AASW) covers the surface ocean. AASW originates from progressive warming of Winter Water (WW, Mosby, 1934), which can be perceived even in summer as a remnant layer of cold water at the base of the AASW (Rintoul et al. 1997).

The Southern ACC Front (SACCF, Fig. 1) is a deep front located south of the PF (Orsi et al., 1995) and can coincide with the southern boundary of the ACC, which is represented by the southern limit of the oxygen minimum (Orsi et al., 1995). The oxygen minimum is related to the advection of Upper Circumpolar Deep Water (UCDW) that originates in the Indian and Pacific Oceans (Callahan, 1972) from where it spreads south, mixes with deep layers of the ACC and ultimately upwells near the Antarctic continent as part of the lower cell of the MOC. Lower Circumpolar Deep Water (LCDW) is below UCDW and is identified by a salinity maximum and contributes to upwelling around Antarctica. The precursor of the LCDW is North Atlantic Deep Water (NADW), originated in the Labrador and Nordic seas of the North Atlantic polar region (Dickson and Brown, 1984) that flows southward to enter the ACC as part of the MOC (Callahan, 1972; Orsi et al., 1995; Johnson, 2008).

At the southern end of the section, the Antarctic Slope Front forms the boundary between cold and fresh shelf water and relatively warm and salty waters offshore (Jacobs, 1991). Polynyas along the Adélie and George V Land coast (Fig.1) contribute to the formation of Adélie Land Bottom Water (ALBW), which is a mixture of High Salinity Shelf Water (HSSW) resulting from brine rejection during ice formation and ultra-modified LCDW (Foster and Carmack, 1976; Rintoul, 1998; Marsland et al., 2004). ALBW constitutes ~ 25% of the total volume of water <0°C in the ocean (Rintoul, 1998). The bottom waters near the southern end of the section also contain a component of Ross Sea Bottom Water (RSBW) that originates to the east and is deflected westwards towards the A-AB as it is transported down the continental slope. The RSBW is modified when it arrives at the location of the SR03 section by mixing with deep layers of the ACC and recently formed ALBW (Gordon and Tchernia, 1972; Rintoul, 1998). The ALBW and the modified RSBW together ventilate the abyssal layers of the A-AB before spreading north to ventilate the Indian and Pacific basins (Mantyla and Reid, 1995; Fukamachi et al. 2010).
3 Data and Method.

3.1 Data.

The SR03 hydrographic section between Tasmania and Antarctica (Fig. 1) was occupied between 1991 and 2011. Measurements of total alkalinity (TA) were only available from the beginning of 1995 and our evaluation of biogeochemical changes is limited to four summer sections occupied for the period 1995-2011 (Table 1). A winter cruise in 1996 was not considered in this study in order to minimise seasonal biases.

Water column salinity, temperature, pressure and dissolved oxygen (O$_2$) were collected from the conductivity-temperature-depth (CTD) device with an accuracy of ±0.002 for salinity and temperature, ±0.015 % of full scale range for pressure and ±1 % for O$_2$, according to WOCE standards (Joyce, 1994). Samples from the Niskin bottles were analysed for dissolved inorganic carbon (DIC) by coulometry and TA by open cell potentiometric titration (Dickson et al, 2007). Certified reference material provided by A. Dickson, Scripps Institution of Oceanography, were used as reference standards for DIC and TA. The precisions of DIC and TA measurements improved slightly on more recent sections, and for all sections were better than ±2 µmol kg$^{-1}$, for both variables, based on analysis of duplicate samples and certified reference material. Samples for dissolved O$_2$ were measured using modified Winkler titrations (Hood et al 2010), with an estimated accuracy and precision of ± 0.3% for the sections. For the 1995-cruise, sensor based O$_2$ was used instead of sampled O$_2$ because of the poor quality of many of the Winkler measurements.

Other variables of the dissolved CO$_2$ system were calculated from the DIC and TA measurements. We calculate pH$_T$ and Ω$_{Ar}$ (from measured DIC and TA) using the CO2sys program from Lewis and Wallace (1998) adapted to MATLAB by van Heuven et al. (2011). We use the constants for the carbonic acid from Mehrbach et al (1973) refit by Dickson and Millero (1987), the CO$_2$ solubility equation from Weiss (1974), dissociation constants for sulphate from Dickson (1990) and borate constant from Uppstrom (1974). Aragonite saturation states (Ω$_{Ar}$) are calculated because it is a less stable form of CaCO$_3$ than calcite and is the predominant biogenic form of CaCO$_3$ precipitated by calcifying organisms.

Data presented here were interpolated to a regular grid, and the water mass layers were defined as layers between neutral surfaces ($\gamma^\nu$) determined using potential temperature and salinity and published literature values (Table 2, section 2). The first 50m of the water column were eliminated in order to reduce the short-time scale variability in surface properties. The use of $\gamma^\nu$ to identify the water mass layers reduces the variability due to isopycnal heave caused for example by eddies and internal waves (McDougall et al., 1987; Bindoff and McDougall, 1994; Jackett and McDougall, 1997). The upper ocean layers south of the SAF were divided into the AASW layer to the north of the PF, and the AASW$_{upw}$ layer that is composed of surface waters south of the PF (Fig.1). ADLBW and RSBW were included in the AABW layer (Table 2). The ACC fronts along the SR03 section (Table 3) were defined as a function of hydrographic variables following Sokolov and Rintoul (2002).
3.2 Estimates of anthropogenic carbon (C\textsubscript{ANT}).

C\textsubscript{ANT} was estimated using a back-calculation method (Chen and Millero, 1979; Gruber, 1998) combined with an Optimum Multi-Parameter (OMP) analysis (Tomczak, 1981), described by Pardo et al. (2014). This technique has the advantage of considering water mass mixing and the temporal variability of the air-sea CO\textsubscript{2} disequilibrium. The accuracy of the method is ±6 μmol kg\textsuperscript{-1} (Pardo et al., 2014). Back-calculation methods assume the ocean is in steady state for dynamical and biological processes and estimate C\textsubscript{ANT} (C\textsubscript{ANT,BC}) as an excess of DIC in the ocean resulting from the increase of atmospheric CO\textsubscript{2} due to anthropogenic emissions as:

\[
C_{\text{ANT,BC}} = DIC - DIC^{BIO} - DIC^{\pi}
\]  

(1)

where DIC\textsuperscript{BIO} is the biological contribution to DIC, and DIC\textsuperscript{\pi} is the preformed concentration of DIC in preindustrial times.

The term DIC\textsuperscript{BIO} is calculated (Chen et al., 1982; Ikegami and Kanamori, 1983) by:

\[
DIC^{BIO} = \frac{AOU}{R_C} + \frac{1}{2} \left[ TA - TA^0 + AOU \ast \left( \frac{1}{R_N} + \frac{1}{R_P} \right) \right]
\]  

(2)

where R\textsubscript{C}, R\textsubscript{N} and R\textsubscript{P} are the stoichiometric ratios of carbon, nitrate and phosphate respectively, referred to O\textsubscript{2} consumption by respiration/remineralization processes that are considered constant (1.45, 9, 125, respectively, Broecker, 1974; Anderson and Sarmiento, 1994; Martiny et al., 2013).

\( \frac{AOU}{R_C} \) represents the remineralization of organic matter, with the apparent oxygen utilization (AOU) defined as (AOU = O\textsubscript{SAT} - O\textsubscript{2}) the difference between the saturation of oxygen (O\textsubscript{SAT}) at the potential temperature (θ) and salinity of the measured O\textsubscript{2}. The term \( \frac{1}{2} \left[ TA - TA^0 + AOU \ast \left( \frac{1}{R_N} + \frac{1}{R_P} \right) \right] \), represents the dissolution/precipitation of CaCO\textsubscript{3}, with TA\textsuperscript{0}, the preformed alkalinity, obtained in the water formation sites using regional parameterizations as function of salinity, θ and phosphate (Pardo et al., 2011; Vázquez-Rodríguez et al., 2012; Appendix section A3, Table A3).

The preformed preindustrial term, DIC\textsuperscript{\pi}, is the total concentration of carbon dioxide in seawater saturated with respect to the preindustrial pCO\textsubscript{2} (DIC\textsubscript{SAT}\textsuperscript{\pi}) and corrected for an air-sea CO\textsubscript{2} disequilibrium (CDIS\textsuperscript{\pi}) term:

\[
DIC^{\pi} = DIC^{\pi}_{\text{SAT}} + CDIS^{\pi}
\]  

(3)

CDIS\textsuperscript{\pi} is time dependent:

\[
CDIS^{\pi} = CDIS - \delta CDIS
\]  

(4)

where CDIS is the current disequilibrium between the ocean and the atmosphere pCO\textsubscript{2}, and \( \delta CDIS \) is the change in the disequilibrium from preindustrial to current times. CDIS was obtained by similar parameterizations to those used for TA\textsuperscript{0}, combined with monthly mean values of atmospheric CO\textsubscript{2} values from the NOAA network (Dlugokencky, et al., 2016) (see section A3 of the appendix and Table A3). The \( \delta CDIS \) values were obtained from results of the 1/10° resolution carbon model OFAM3-WOMBAT (Appendix section A1).

Interior values of preformed variables (TA\textsuperscript{0} and DIC\textsubscript{SAT}\textsuperscript{\pi}) and CDIS\textsuperscript{\pi} were obtained using an optimum multiparameter (OMP) analysis to mix end members as described in Appendix sections A2 and A3, using the end members in Table A1. The OMP analysis is based on the assumption that a property measured in a certain point is the result of linear mixing between end
members, known as source water types (SWTs). A system of equations is created for each measurement point and is solved to obtain the fractions of the different SWTs (Appendix section A2). The application of the OMP analysis requires good regional hydrodynamic knowledge as the results are strongly dependant on the definition of the SWTs (Tomczak, 1981). We used 11 SWTs to characterize the biogeochemical properties of the waters in the SR03 section and the SWT properties were assumed to be constant with time (Appendix Table A1, Fig. A1).

Small negative values of C\text{ANT}_BC can occur due to an overestimation of the $CDIS\pi$ term (Eq. 4), that acts as measure of the age of the water mass and has high values in old deep layers (see Table A1 in the appendix). These negative values of C\text{ANT}_BC were found at some points in deep waters of the section (mainly UCDW and LCDW layers) and were small (between 0 and -2 \, \mu mol \, kg^{-1}, i.e., less than the accuracy of the methodology) and changed to zero for our analysis.

3.3 Changes in the carbon system.

Changes in carbon system parameters ($\partial DIC/\partial t$, $\partial TA/\partial t$, $\partial C\text{ANT}_BC/\partial t$, $\partial pH/\partial t$ and $\partial OA/\partial t$) were estimated using linear regressions with time for the period 1995-2011 in each one of the water mass layers defined by their $\gamma^a$ condition (Table 2). The trends were estimated using all the points in each water mass layer and only those linear trends with p<0.05 and r\geq0.2 were considered statistically significant for discussion. We show the value of the root mean square error (RMSE or square root of the variance of the residuals), which can be interpreted in large part as unexplained variance caused by short-time scale processes and the different seasonal timings of the cruises. RMSE has the same units as the response variable.

Respect to the total change of DIC ($\partial DIC/\partial t$), our goal is to disentangle the effects that solubility, circulation, biology and C\text{ANT} uptake have on the variability of DIC. The total change of DIC ($\partial DIC/\partial t$) in a water mass is due to changes in the atmosphere-ocean interchange, biological processes and circulation processes. In order to account of the change in DIC due the atmosphere-ocean interchange and biological processes, we compare $\partial DIC/\partial t$ to $\partial C\text{ANT}_BC/\partial t$ and $\partial DIC^{BIO}/\partial t$, Eq. (2). The change in DIC not explained by $\partial C\text{ANT}_BC/\partial t$ or $\partial DIC^{BIO}/\partial t$ will then be due to circulation processes.

In order to compare $\partial DIC/\partial t$ to $\partial C\text{ANT}_BC/\partial t$ we need to consider how the change in $\partial DIC^{BIO}/\partial t$ and $DIC^{\pi}(\partial DIC^{\pi}/\partial t)$ (terms of Eq (1), section 3.2) affect the changes in C\text{ANT}_BC.

The term $\partial DIC^{\pi}/\partial t$ can be expressed as:

$$\frac{\partial DIC^{\pi}}{\partial t} = \frac{\partial DIC^{\pi}_{SAT}}{\partial t} - \frac{\partial CDIS^{\pi}}{\partial t}$$

The terms $\frac{\partial CDIS^{\pi}}{\partial t}$ and $\frac{\partial DIC^{\pi}_{SAT}}{\partial t}$ reflect changes in the properties of the water masses over time, primarily temperature and salinity change due to mixing and heating/cooling. CDIS$^{\pi}$ and DIC$^{\pi}_{SAT}$ are defined at the ocean surface (in each of the SWTs: Table A1) and are calculated at each point in the ocean interior using the OMP analysis (Appendix sections A2 and A3). Because a change in temperature and/or salinity in the water is solved by the OMP analysis as a change in the SWTs fractions, this also
produces varying $CDI_S^{\pi}$ and $DIC^{\pi}_{SAT}$. No significant trends were obtained for $CDI_S^{\pi}$ in any of the layers. $C_{\text{ANT}_B}$ (Eq. 1) is not affected by the changes in solubility occurring from one voyage to another (thus neither is $\frac{\partial C_{\text{ANT}_B}}{\partial t}$), since any change in temperature or salinity is cancelled out by the subtraction of $DIC^{\pi}$ with respect to DIC in Eq. (1) (section 3.2) and by $O_2$ respect to $O_{\text{SAT}}$ in the $DIC^{BIO}$ term (Eq. 2). However, $\frac{\partial DIC}{\partial t}$ based on the measured DIC in the sections will be affected by changes in the solubility over time and this difference needs to be accounted for when comparing $\frac{\partial DIC}{\partial t}$ with $\frac{\partial C_{\text{ANT}_B}}{\partial t}$ to obtain a better approximation for the change in $C_{\text{ANT}}$: $\frac{\partial C_{\text{ANT}}}{\partial t} = \frac{\partial C_{\text{ANT}_B}}{\partial t} + \frac{\partial DIC^{\pi}}{\partial t}$.

The term $\frac{\partial DIC^{BIO}}{\partial t}$ can be influenced by changes with time of alkalinity due to changes in the rate of carbonate precipitation/dissolution and of AOU due to changes in the rate of remineralization and in circulation. In the present study only surface waters of the SR03 section present changes $DIC^{BIO}$ between 1995 and 2011. Numerous studies have reported a strong influence of biological communities in the seasonal cycle of dissolved $O_2$ in surface waters (Bender et al., 1996; Moore and Abbott, 2000; Sambrotto and Mace, 2000; Trull et al., 2001a). Interannual variability in $O_2$ in upper layers of the Southern Ocean have also been related to changes in the entrainment of deeper waters into the mixed layer due to the mixed layer depth variability (Matear et al., 2000; Verdy et al., 2007; Sabine et al., 2008; Sallée et al., 2012). Although some studies found long-term decreases in $O_2$ due to circulation in deep waters of the Weddell Sea (van Heuven et al., 2014) and for the first 1000 m of the global ocean (Helm et al. 2011), significant long-term trends in $O_2$ due to circulation and remineralization processes have not yet been reported for surface waters of the Southern Ocean. Thus, the term $\frac{\partial DIC^{BIO}}{\partial t}$ may also contribute to variation in $\frac{\partial C_{\text{ANT}_B}}{\partial t}$, since part of the changes in AOU with time reflect changes in circulation that we cannot separate from those in remineralization. We consider the best approximation for the change in $C_{\text{ANT}}$ as a range depending on the possible effect of biology and circulation processes on $\frac{\partial DIC^{BIO}}{\partial t}$. If the value of $\frac{\partial DIC^{BIO}}{\partial t}$ is due to the variability in the remineralization rates and the change in solubility is considered, the estimate $\frac{\partial C_{\text{ANT}_B}}{\partial t}$ will be the lower limit of the range, (lower limit of $\frac{\partial C_{\text{ANT}}}{\partial t} = \frac{\partial C_{\text{ANT}_B}}{\partial t} + \frac{\partial DIC^{\pi}}{\partial t}$). For the upper limit of the range, we consider that the value of $\frac{\partial DIC^{BIO}}{\partial t}$ is due to changes in circulation and the upper limit of the range is obtained by $\frac{\partial C_{\text{ANT}}}{\partial t} = \frac{\partial C_{\text{ANT}_B}}{\partial t} + \frac{\partial DIC^{\pi}}{\partial t} + \frac{\partial DIC^{BIO}}{\partial t}$. We assume that the changes in $DIC^{BIO}$ due to circulation do not affect the amount of DIC in the layer. This assumption is one of the caveats of the methodology, since we cannot know how much of the change in DIC is associated to changes in circulation, i.e., how much of the change in DIC is a change in non-anthropogenic DIC. We will discuss this more in section 6.2.

4 Results.

The different trends in biogeochemical properties are summarized in Table 4. The biogeochemical changes between 1995 and 2011 are presented for each of the water mass layers and the effect of changes in solubility, biological processes and
circulation in the estimates of $\frac{\partial C_{\text{ANT}}}{\partial t}$ and $\frac{\partial DIC}{\partial t}$ are considered along with changes in the aragonite saturation depth and $C_{\text{ANT}}$ storage.

### 4.1 Changes in DIC, $C_{\text{ANT}}$ and pH$_T$ (1995-2011).

#### 5 Upper ocean layers (STCW, AASW and AASW$_{upw}$).

In the STCW layer, DIC increased between 1995 and 2011 (Fig. 2a, b) at a rate of 0.86 ± 0.07 μmol kg$^{-1}$ yr$^{-1}$ ($\frac{\partial DIC}{\partial t}$, Table 4), leading to a decrease of pH$_T$ of $-0.0027 \pm 0.0001$ yr$^{-1}$ ($\frac{\partial pH_T}{\partial t}$, Table 4, Fig. 2e, f). The trend in pH$_T$ is similar to the one found by Lauvset et al. (2015) between 1991 and 2011 for the IO-STPS (Indian Ocean subtropical permanently stratified) biome ($-0.0027 \pm 0.0005$ yr$^{-1}$). We found a decrease of DIC$_\pi$ ($\frac{\partial DIC_{\pi}}{\partial t} = -0.34 \pm 0.06$ μmol kg$^{-1}$ yr$^{-1}$, Table 5) in the STCW layer due to a negative trend of DIC$_{\text{SAT}}$ resulting from a decrease in solubility that resulted from a temperature increase (calculated from the section data) in the STCW layer of 0.0335 ± 0.0130 °C yr$^{-1}$ (not shown). The increase in temperature agrees with the warming trend observed south of Tasmania of 0.2 to 0.3 °C decade$^{-1}$ obtained from satellite data (Armour and Bitz, 2015) and from combined data and models (0.5 °C / 30 yr$^{-1}$, Aoki et al., 2015). For θ=16 °C and S=35.1 (definition of SWT$_{STW16}$ in the OMP analysis, Table A1), a change in temperature of 0.03 °C yr$^{-1}$ would lead to a decrease in DIC$_{\text{SAT}}$ of $-0.27$ μmol kg$^{-1}$ yr$^{-1}$, which is similar to the value obtained for $\frac{\partial DIC_{\pi}}{\partial t}$ in the STCW layer (Table 5). The difference between these trends is related to the mixing of the different SWTs fractions within the STCW layer established from the OMP analysis (see section 3 and section A2 of the Appendix). When the solubility change is incorporated into $\frac{\partial C_{\text{ANT}}}{\partial t}$, i.e. $\frac{\partial C_{\text{ANT}}}{\partial t} B_C + \frac{\partial DIC_{\pi}}{\partial t}$ (Table 5, Fig. 2c, d), we obtain a value of $0.71 \pm 0.08$ μmol kg$^{-1}$ yr$^{-1}$.

There is an increase of DIC$_{\text{BIO}}$ in the STCW layer ($\frac{\partial DIC_{\text{BIO}}}{\partial t}$, Table 5), that also affects the estimates of $\frac{\partial C_{\text{ANT}}}{\partial t}$. The increase of DIC$_{\text{BIO}}$ is due to an increase in AOU (no changes were found in TA, Eq. (2) in section 3.2) due to a decrease of O$_2$ in the layer. We cannot separate the effects of circulation and biology on the AOU change and $\frac{\partial C_{\text{ANT}}}{\partial t}$ in Table 4 should be considered a range. If the changes in AOU are only due to the variability in the remineralization rates, the calculated lower limit of $\frac{\partial C_{\text{ANT}}}{\partial t}$ is $0.71 \pm 0.08$ μmol kg$^{-1}$ yr$^{-1}$ (Table 4, $\frac{\partial C_{\text{ANT}}}{\partial t} = \frac{\partial C_{\text{ANT}}}{\partial t} B_C + \frac{\partial DIC_{\pi}}{\partial t}$ in Table 5). If the changes in AOU are due to changes in circulation, the upper limit value of $0.93 \pm 0.11$ μmol kg$^{-1}$ yr$^{-1}$ ($\frac{\partial C_{\text{ANT}}}{\partial t} = \frac{\partial C_{\text{ANT}}}{\partial t} B_C + \frac{\partial DIC_{\pi}}{\partial t} + \frac{\partial DIC_{\text{BIO}}}{\partial t}$) will explain the increase of DIC in the STCW layer ($\frac{\partial DIC}{\partial t} \approx \frac{\partial C_{\text{ANT}}}{\partial t}$, Table 4). The increase of C$_{\text{ANT}}$ found in this layer is comparable to the range of increase ($0.8 – 1.3$ μmol kg$^{-1}$ yr$^{-1}$) found by Carter et al. (2017) in the Pacific Ocean (P16 WOCE, CLIVAR and GOSHIP lines) for the two past decades (1990s-2000s and 2000s-2010s).
Changes in the AASW layer are summarised in Table 4. DIC increased at a similar rate of 0.85 ± 0.14 μmol kg\(^{-1}\) yr\(^{-1}\) to the STCW layer and the trend is similar to the values found by Williams et al. (2015) for the AASW layer in the Pacific sector of the SO (12-18 μmol kg\(^{-1}\) for the period 1992-2011 and 3-5 μmol kg\(^{-1}\) for the period 2005-2011). The increase of DIC in the AASW layer results in a pH\(_T\) decrease of -0.0035 ± 0.0002 yr\(^{-1}\), close to Williams et al. (2015) estimates for surface waters (~ -0.0023 ± 0.0009 yr\(^{-1}\)) and Lauvset et al. (2015) estimates of -0.0021 ± 0.0002 yr\(^{-1}\) for the Southern Ocean seasonally stratified, SO-SPSS, biome. The AASW layer for our sections warmed at a similar rate (0.0369 ± 0.0109 °C yr\(^{-1}\)) to the STCW layer, reducing the solubility and influencing \(\frac{\partial\text{DIC}}{\partial t}\) (Table 5) to changes in \(\text{DIC}_{\text{SAT}}\). The \(\text{DIC}_{\text{BIO}}\) also increased with time (0.50 ± 0.16 μmol kg\(^{-1}\) yr\(^{-1}\), Table 5) due to an increase of AOU. Following the same reasoning as for the STCW layer and considering the trend in \(\text{C}_{\text{ANT}}\) of 0.70 ± 0.06 μmol kg\(^{-1}\) yr\(^{-1}\) obtained by the back-calculation method (\(\frac{\partial\text{C}_{\text{ANT}}}{\partial t}\) ; Table 5), the best estimation of \(\frac{\partial\text{C}_{\text{ANT}}}{\partial t}\) in the AASW layer is a range of 0.35 ± 0.14 to 0.85 ± 0.22 μmol kg\(^{-1}\) yr\(^{-1}\) (Table 4). Our values are within the range of values found by Williams et al. (2015) for the AASW in the Pacific sector between 2005 and 2011 and the upper limit is similar to a \(\text{C}_{\text{ANT}}\) increase of 0.73 - 0.86 μmol kg\(^{-1}\) yr\(^{-1}\) for waters South of Tasmania for the period 1968-1996 found by McNeil et al. (2001). DIC in the AASW\(_{\text{upw}}\) layer increased at a rate of 0.61 ± 0.10 μmol kg\(^{-1}\) yr\(^{-1}\), and the pH\(_T\) decreased -0.0015 ± 0.0004 yr\(^{-1}\) (Table 4). We were not able to detect a statistically significant trend in \(\text{DIC}_{\pi}\) (i.e., solubility) or \(\text{C}_{\text{ANT}}\) from the estimates of the back-calculation method (\(\frac{\partial\text{C}_{\text{ANT}}}{\partial t}\) , Table 5). However, we found an increase of \(\text{DIC}_{\text{BIO}}\) of 0.42 ± 0.28 μmol kg\(^{-1}\) yr\(^{-1}\) (Table 5) that is due to an increase in AOU. Considering the different drivers of the AOU increase (biology/circulation), the optimal estimation of \(\frac{\partial\text{C}_{\text{ANT}}}{\partial t}\) for this layer is a value between 0 and 0.42 ± 0.28 μmol kg\(^{-1}\) yr\(^{-1}\) (Table 4).

### 4.1.2 Mode waters and intermediate layers (SAMW and AAIW).

The increase in DIC in the SAMW layer (1.10 ± 0.14 μmol kg\(^{-1}\) yr\(^{-1}\)) for the period 1995-2011 is higher than that of upper ocean layers and pH\(_T\) decreases over the same period at -0.0031 ± 0.0003 yr\(^{-1}\) (Table 4). The DIC increase is explained almost entirely by \(\frac{\partial\text{C}_{\text{ANT}}}{\partial t}\) of 0.92 ± 0.09 μmol kg\(^{-1}\) yr\(^{-1}\). No significant trend was found in \(\text{DIC}_{\text{BIO}}\) or \(\text{DIC}_{\pi}\) (i.e, \(\frac{\partial\text{C}_{\text{ANT}}}{\partial t} = \frac{\partial\text{C}_{\text{ANT}}}{\partial t}\)). In the AAIW layer the DIC trend of 0.40 ± 0.15 μmol kg\(^{-1}\) yr\(^{-1}\) results in a pH\(_T\) decrease of -0.0017 ± 0.0002 yr\(^{-1}\) and is also explained by the increase of \(\text{C}_{\text{ANT}}\) (0.42 ± 0.06 μmol kg\(^{-1}\) yr\(^{-1}\), \(\frac{\partial\text{C}_{\text{ANT}}}{\partial t} = \frac{\partial\text{C}_{\text{ANT}}}{\partial t}\), Tables 4 and 5). As with SAMW, no changes in solubility (\(\frac{\partial\text{DIC}_{\pi}}{\partial t}\)) or biology/circulation processes (\(\frac{\partial\text{DIC}_{\text{BIO}}}{\partial t}\)) were detected in the AAIW layer. The values found in the SAMW and AAIW layers are very similar to the mean decadal changes found by Murata et al. (2007) between the 1990s and the 2000s in the subtropical Pacific Ocean (~1 μmol kg\(^{-1}\) yr\(^{-1}\) for the SAMW layer and 0.4 μmol kg\(^{-1}\) yr\(^{-1}\) for the AAIW). Waters et al., (2011) used data from the P18 line along ~110°W and estimated an increase in \(\text{C}_{\text{ANT}}\) of 0.89 ± 0.4 μmol kg\(^{-1}\) yr\(^{-1}\) for the SAMW and 0.64 ± 0.2 μmol kg\(^{-1}\) yr\(^{-1}\) in AAIW for the period 1994-2008, which are also comparable to our results.
4.1.3 Deep-bottom layers (UCDW, LCDW and AABW).

The UCDW layer shows an increase of DIC of $0.29 \pm 0.02 \ \mu \text{mol} kg^{-1} \text{yr}^{-1}$ between 1995 and 2011 and a change in pH$_T$ of $-0.0013 \pm 0.0001 \ \text{yr}^{-1}$ and are similar to the change in DIC ($0.20 \pm 0.02 \ \mu \text{mol} kg^{-1} \text{yr}^{-1}$) and pH$_T$ ($-0.0012 \pm 0.0002 \ \text{yr}^{-1}$) for LCDW. No statistically significant changes in time were detected for $C_{ANT, BC}$ or in the $DIC^{\text{Bio}}$ and $DIC^{\pi}$ terms for any of these two layers. The AABW layer also shows an increase of DIC ($0.24 \pm 0.02 \ \mu \text{mol} kg^{-1} \text{yr}^{-1}$) during the period 1995-2011 with an associated decrease of pH$_T$ of $-0.0013 \pm 0.0002 \ \text{yr}^{-1}$ (Table 4). The increase in $C_{ANT}$ ($\frac{\partial C_{ANT, BC}}{\partial t}$) of $0.07 \pm 0.01 \ \mu \text{mol} kg^{-1} \text{yr}^{-1}$ is low and this trend indicates an increase in $C_{ANT}$ of $\sim 1 \ \mu \text{mol} kg^{-1}$, which is less than the accuracy of the back-calculation method ($\pm 6 \ \mu \text{mol} kg^{-1}$).

4.2 Changes in the aragonite saturation ($\frac{\partial \Omega_{Ar}}{\partial t}$) and $C_{ANT}$ storage.

There are statistically significant decreases of $\Omega_{Ar}$ in the STCW, AASW and SAMW layers ($\sim -0.010 \pm 0.001 \ \text{yr}^{-1}$, Table 4) similar to the trends observed at open-ocean time series sites in recent decades (Bates et al., 2014). The decrease of $\Omega_{Ar}$ found for the AASW layer ($-0.61 \pm 0.19 \ % \ \text{yr}^{-1}$, Table 4) is also similar to the values obtained by Williams et al. (2015) for the Pacific sector of the Southern Ocean ($-0.47 \pm 0.10 \ % \ \text{yr}^{-1}$ for the period 1992-2011 and $-0.50 \pm 0.20 \ % \ \text{yr}^{-1}$ for the period 2005-2011). Accompanying the decrease of $\Omega_{Ar}$ with time along SR03, is the shoaling of the aragonite saturation depth (ASD, $\Omega_{Ar} = 1$, Fig. 3) at a mean rate of $-13 \pm 3 \ \text{m} \ \text{yr}^{-1}$. The shoaling of the ASD is not uniform over the section. North of the PF, the ASD shoals at a rate of $-6 \pm 4 \ \text{m} \ \text{yr}^{-1}$ while the rate is 3.5 times greater south of the PF ($-21 \pm 4 \ \text{m} \ \text{yr}^{-1}$). North of the PF the shoaling mostly affects the AAIW layer (Fig. 3a). South of the PF from $\sim 62^\circ$S, the movement of the ASD follows the upwelling path of the UCDW layer (Fig. 3) with a shoaling of $\sim 340 \ \text{m}$ over the 16-year period.

The storage rate of $C_{ANT}$ (Table 6) for the surface and intermediate water mass layers is obtained from $\frac{\partial C_{ANT}}{\partial t}$ (Table 4) with the most storage in SAMW and AAIW due to both their greater thickness and $\frac{\partial C_{ANT}}{\partial t}$ values. The rate of increase of the $C_{ANT}$ storage in the whole longitude band of the SR03 section is $0.30 \pm 0.24 \ \text{mol} m^{-2} \ \text{yr}^{-1}$, calculated by computing the mean of the storage rates of the layers weighted by the mean volume occupied by each of the layers for the period 1995-2011 (Table 6).

5 Discussion.

Our results are indicative of a scenario of increased transport of deep waters into the section and enhanced upwelling at high latitudes for the period between 1995 and 2011 linked to strong westerly winds. Several studies have reported a trend in the Southern Annular Mode (SAM) toward its positive phase from the 1960s until the 2000s (Thompson and Solomon, 2002; Marshall, 2002, 2003; Lenton and Matear, 2007; Sallée et al., 2008). According to these studies, the positive phase of the SAM is correlated with an intensification and southward movement of the subpolar westerly winds that ultimately lead to the
enhancement of northward Ekman transport, meridional overturning and upwelling south of the ACC. Also, surface warming and more intense and frequent pulses in the extension of the EAC at long-time scales have been related to a poleward movement of the westerly winds (Rintoul and Sokolov, 2001; Ridgway, 2007; Hill et al., 2011). From the 2000s on, the SAM index no longer presents a positive trend but, although exhibiting considerable interannual variability (Fig. A2 in the supplementary material), the SAM index remains in its positive phase, favouring strong winds over the region.

In the northern part of the SR03 section, the area occupied by the STCW has high variability due to the encounter between the EAC and the ZC in the North of the section (Ridgway et al., 2007; Herraiz-Borreguero and Rintoul, 2011; Sloyan et al., 2016). The warming of the STCW layer found in this study ($0.0335 \pm 0.0130 \, ^\circ\text{C yr}^{-1}$) could be linked to variability in the extension of subtropical waters but it could also be related to atmospheric warming. Aoki et al. (2015) related the 30-year warming found north of the SAF in the South Pacific and Indian oceans to the intensification of the subtropical gyres, which promote the arrival of warmer waters. In the AASW layer that extends approximately between the SAF and the PF we found a similar warming ($0.0369 \pm 0.0109 \, ^\circ\text{C yr}^{-1}$) to that of the STCW. This could indicate that the increase of temperature found in the upper layers of the section could be most likely due to ocean heat uptake and atmosphere warming.

Due to the surface warming, the increase of DIC found in the STCW layer (Table 4) is lower than expected from the increase in atmospheric CO$_2$ ($\sim 1 \pm 0.12 \, \mu\text{mol kg}^{-1} \, \text{yr}^{-1}$). Nevertheless, at least 83% of the increase of DIC in the STCW layer is explained by the increase in C$_{\text{ANT}}$ (Table 4). As for the AASW layer, our results indicate that temperature does affect the estimate of $\frac{\partial \text{DIC}}{\partial t}$, but the effect of the increase in $\text{DIC}^{\text{BIO}}$ (due to an increase in AOU) outweigh that of solubility (Table 5).

The seasonal to interannual variability of the AASW layer is also influenced by the variability of the positions of the SAF and PF (Fig. 1, Table 3), that is highly conditioned by the flow of the ACC over the South-East Indian Ridge (Fig. 1). A close relationship between phytoplankton blooms and regions where the ACC fronts interact with large topographic features has been noticed (Moore et al., 1999; Moore and Abbott, 2000). A variability in the remineralization rates due to phytoplankton blooms variability could explain the changes in $\text{DIC}^{\text{BIO}}$ observed in the AASW layer. Nevertheless, no changes in nutrients (nitrates or phosphates) are measurable in this layer that could indicate intense biological activity.

Furthermore, the AASW layer is also affected by the upwelling of deep waters south of the PF, and an intensification of the upwelling could increase the content of low-O$_2$ DIC-rich waters in the AASW layer leading to an increase in AOU. The increase in $\text{DIC}^{\text{BIO}}$ found in the AASW$_{\text{upw}}$ layer (Table 4), south of the PF, is similar to the increase obtained for the AASW layer, which indicates the likelihood that the upwelling of deep waters results in the increase in AOU. The increase in $\text{DIC}^{\text{BIO}}$ in the AASW$_{\text{upw}}$ layer coincides with an increase of salinity of $0.0029 \pm 0.0001 \, \text{yr}^{-1}$ (not shown), that is consistent with increased transport of saltier waters from the deep ocean to subsurface layers. Besides, we also found an increase in dissolved silicate of $0.36 \pm 0.06 \, \mu\text{mol kg}^{-1} \, \text{yr}^{-1}$ ($\frac{\partial \text{SiO}_4}{\partial t}$; Table 4) that could be related to the upwelling enhancement as well (Tréguer, 2014).

The influence of the upwelling on the DIC budgets (as non-anthropogenic DIC) is clearer in the AASW$_{\text{upw}}$ layer than in the AASW layer. For the AASW layer, the lower limit of $\frac{\partial \text{C}_{\text{ANT}}}{\partial t}$ (i.e., the change in $\text{DIC}^{\text{BIO}}$ is assumed to be due to biological processes, Table 4) indicates that at least 41% of the increase of DIC in the layer is explained by the increase of C$_{\text{ANT}}$ while
the lower limit of $\frac{\partial C_{\text{ANT}}}{\partial t}$ is zero for AASW_{upw} (Table 4), meaning that the effect of the upwelling over AASW is lower than over AASW_{upw}. Matear and Lenton (2008) using carbon models, concluded that the uptake of CO$_2$ by the waters north of the PF is more influenced by the wind variability than by other processes such as the upwelling. An intensification of the winds (due to a positive phase in SAM) could contribute to the increase in $C_{\text{ANT}}$ found in the AASW layer. Considering the upper limits of $\frac{\partial C_{\text{ANT}}}{\partial t}$ in both layers (i.e., the change in $DIC^{BIO}$ is assumed to be due to circulation processes), the increase of $C_{\text{ANT}}$ in the AASW_{upw} layer represents no more than 69% of the increase in DIC (upper limit of $\frac{\partial C_{\text{ANT}}}{\partial t}$, Table 4) while the upper limit of $\frac{\partial C_{\text{ANT}}}{\partial t}$ for the AASW equals the increase of DIC. Thus, AASW_{upw} layer, at least ~30% of the increase in DIC (~0.18 μmol kg$^{-1}$ yr$^{-1}$) is still not explained and is most probably related to the upwelling of DIC-rich waters. The increase in non-anthropogenic DIC could be even higher, since we assume that the change in $DIC^{BIO}$ due to circulation does not affect DIC (see section 3.3).

In terms of the change in oxygen, Helm et al. (2011) found an average decrease in the concentration of O$_2$ between 100 and 1000 m from 1970 to 1992 of ~0.23 μmol l$^{-1}$ for the Southern Ocean (27% of the estimated global average change, -0.93 ± 0.23 μmol l$^{-1}$). Considering the volume of the first 1000 m of the water column of the Southern Ocean to be 19400 · 10$^9$ l (obtained using ETOP01 doi:10.7289/V5C8276M) and the volume of the first 1000 m of the SR03 section to be 2700 · 10$^9$ l, the decrease of O$_2$ found by Helm et al. (2011), if constant in time, would correspond to a decrease of ~1.7 μmol l$^{-1}$ yr$^{-1}$. We only found changes in oxygen within the surface water mass layers (STCW, AASW and AASW$_{upw}$) that approximately fill the first 300 m of the water column of the SR03. Then, the decrease of ~1.7 μmol l$^{-1}$ would correspond to an average change of O$_2$ of ~0.32 μmol kg$^{-1}$ yr$^{-1}$ for surface waters of the SR03. This means that values of ~0.20 μmol kg$^{-1}$ yr$^{-1}$ due to circulation processes can be expected in $\frac{\partial DIC^{BIO}}{\partial t}$ for surface waters, which is comparable to the average of our findings (Table 5), 0.32 ± 0.24 μmol kg$^{-1}$ yr$^{-1}$ and could indicate that the change in O$_2$ is related to circulation processes.

The variability of the SAMW and AAIW layers south of Tasmania has been related to variability in the northward Ekman transport that drives the northward movement of AASW (Rintoul and England, 2002; Sallée et al., 2006, 2012). A scenario of intensification of the upwelling near the Antarctic Divergence would lead to an increase in the northward Ekman transport, conditioning the properties of these water mass layers and particularly for SAMW, which is mostly form north of the SAF.

There is a significant freshening of the SAMW layer (-0.0026 ± 0.0001 psu yr$^{-1}$, not shown) between 1995 and 2011 that could be related to higher inputs of AASW into the SAMW layer and consistent with the increase in Ekman transport. Besides, an intensification of the winds due to the positive trend of the SAM favours the ventilation and thus the increase in $C_{\text{ANT}}$ uptake by both water mass layers (Matear and Lenton, 2008). Our results indicate that the change of DIC in the SAMW and AAIW layers is driven mostly by the uptake of atmospheric CO$_2$ ($\frac{\partial DIC}{\partial t} \approx \frac{\partial C_{\text{ANT}}}{\partial t}$, Table 4). The increase of $C_{\text{ANT}}$ in the SAMW layer is higher than that found for the upper ocean layers and closer to the expected from the increase of atmospheric CO$_2$ (~1 μmol kg$^{-1}$ yr$^{-1}$). The smaller increase of $C_{\text{ANT}}$ in the AAIW layer compared to the SAMW layer (Table 4) agrees with lower ventilation of the AAIW layer south of Tasmania (see section 2) due to the fact that this layer carries recently ventilated waters mixed
with older waters ventilated far out the SR03 section. The lack of measurable long-term changes in \( \text{DIC}^{\text{Bio}} \) and \( \text{DIC}^{\pi} \) in both AAIW and SAMW layers indicate that circulation and biological processes do not have a large effect on \( \frac{\partial \text{DIC}}{\partial t} \).

Deep to bottom layers of the section show significant trends for DIC that are not explained by the increase of \( \text{C}_{\text{ANT}} \). These trends are most likely due to the advection of old and DIC-rich waters. Concretely for deep waters (UCDW and LCDW), the trends could result from an intensification of upwelling at high latitudes being offset by enhanced transport of old and CO\(_2\)-rich waters to replace the upwelled waters, since the increase of DIC follows the upwelling path of the UCDW and LCDW layers (Fig. 4). We separated the UCDW layer into two latitudinal sectors: north and south of the SAF (Fig. 4). The increase of DIC in the UCDW layer north of the SAF is 0.44 ± 0.04 μmol kg\(^{-1}\) yr\(^{-1}\) while south of the SAF is smaller at 0.26 ± 0.04 μmol kg\(^{-1}\) yr\(^{-1}\) (not shown), consistent with a greater supply of waters from the north at depth. A decrease of O\(_2\) in the UCDW to the north of the SAF occurs mostly in the upper to middle parts of the UCDW layer (Fig. 4), and this is not observed south of the SAF. The decrease of O\(_2\) is also in agreement with the arrival of waters from Indian-Pacific origin since these waters provide the characteristic oxygen minimum zone that defines UCDW (Callahan, 1972; Talley 2013). Another feature that agrees with the hypothesis of upwelling intensification is the shoaling of the ASD following the path of upwelling of the UCDW layer. This feature was also described by Bostock et al. (2013) in an oceanic climatology of \( \Omega_{\text{A}} \) and could be due to the naturally lower buffer capacity of the UCDW layer (low value of TA/DIC ≈ 1.043) with respect to upper layers (TA/DIC ≈ 1.06 in the AASW layer). However, the greatest shoaling of the ASD in the UCDW layers compared to the AAIW layer (Table 6) is consistent with the upwelling of UCDW, as both water masses have similar TA/DIC ratios (TA/DIC ≈ 1.043 for the UCDW layer and TA/DIC ≈ 1.042 for the AAIW layer). Furthermore, the increase of SiO\(_4\) found in deep-bottom layers (Table 4) could also indicate the arrival of old waters to the section that are progressively enriched in SiO\(_4\) (e.g., Callahan, 1972).

Statistically significant decreases in pH\(_T\) with time were observed in all water mass layers (Table 4), with the greatest change in surface water masses, coinciding with the greatest DIC changes. The decrease of pH\(_T\) in the STCW and SAMW layers is related to the increase in the uptake of \( \text{C}_{\text{ANT}} \), while for the AASW and AASW\(_{\text{upw}}\) layers the pH\(_T\) change appears to be linked to the upwelling of DIC-rich waters at high latitudes. At deep layers, the tongue of water of pH\(_T\) ≈ 7.9 off the shelf is reduced in 2011 compared to 1995 (Fig. 2e,f), which is consistent with the advection of DIC-rich waters in the section due to the enhanced upwelling. The different rates of pH\(_T\) change in the water masses is in part related to the buffering capacity of the waters. AASW layer has lower temperature than the STCW layer (~2.3 °C for the AASW layer compared to ~11.0 °C for the STCW layer, mean values for the period 1995-2011) and lower buffer capacity than the STCW (TA/DIC ≈ 1.058 for the AASW layer versus TA/DIC ≈ 1.095 for STCW). For similar increases in DIC (Table 4) the decrease of pH\(_T\) in the AASW is expected to be higher than in the STCW layer.

In terms of carbon, previous studies concluded that the intensification of the upwelling (as a consequence of the SAM variability) caused a reduction in the uptake of CO\(_2\) by the Southern Ocean between 1980s and 2000s due to the outgassing of CO\(_2\) near the Antarctic Divergence (Le Quéré et al, 2007; Lovenduski et al., 2008). Landschützer et al. (2015) showed that the
efficiency of the Southern Ocean CO$_2$ sink declined through the 1990’s, and the trend reversed from about 2002, although the reversal in the sink efficiency was not zonally uniform. The results from Landschützer et al. (2015) are consistent with a carbon sink influenced by the upwelling of DIC-rich waters at high latitudes and superimposed on this is the near surface response to atmospheric forcing that modifies the sink efficiency and could mask longer term trends in the upwelling of DIC-rich waters at high latitudes. A comparison of our results with those of Landschützer et al. (2015) is problematic as their data is restricted to surface waters and our analysis is on long-term trends in water mass properties below 50m depth. Both data sets do show continued uptake of CO$_2$ throughout the period of study and indicate the importance of the circulation in influencing the regional carbon sink, which has also been established by recent model results (DeVries et al., 2007).

Our results also agree with the conclusions from different model simulations done by Matear and Lenton (2008), who established that intense wind regimes (associated to a positive phase in the SAM) favour the uptake of C$_{ANT}$ and ventilation of the SAMW and AAIW layers. These authors highlighted the complex response of the uptake of CO$_2$ by the Southern Ocean due to the diverse forcing acting on upper layers, which can be also seen in our results (e.g., differences in the biogeochemical changes in the AASW and AASW$_{upw}$ layers). Matear and Lenton (2008) also noticed the complex relationship between the upwelling and subduction areas of the Southern Ocean, with the same drivers acting in opposite direction for the changes in non-anthropogenic DIC with respect to the changes in C$_{ANT}$ uptake.

6 Sensitivity of the results to underlying assumptions.

This section considers the sensitivity of assumptions used to calculate temporal changes in C$_{ANT}$, including errors associated with the assumption of steady state in the oceanic circulation and remineralization processes, and the sensitivity to stoichiometric ratios for the biological processes.

6.1 Comparison of C$_{ANT}$ changes using other methods.

We compared the changes of C$_{ANT}$ obtained in our study with the results from two regression-based methodologies (Table 7); the extended multiple linear regression (eMLR) method (Friis et al., 2005) and the two-regression method (Thacker, 2012). These methods use repeat hydrodynamic sections to quantify the temporal change in C$_{ANT}$.

The eMLR method (Friis et al., 2005) estimates the change in C$_{ANT}$ between two repeats of a hydrodynamic section by establishing MLRs for each section and relating the observed DIC for each observation to a set of other measured oceanic variables:

$$DIC(t) = a_0(t) + a_{1(t)}P_{1(t)} + \cdots + a_{n(t)}P_{n(t)}$$

(6)

where $a_{x(t)}$ are the coefficients of the fit between DIC and the n observed variables ($P_1,\ldots,P_n$) chosen for the fit, all measured at the time (t) of the survey.

Taking the difference between DIC at two times, t1 and t2, gives an equation for the change in C$_{ANT}$ over the time period between the two hydrographic surveys ($\Delta C_{ANT}$):
\[ \Delta C_{\text{ANT}} = a_{0(t2)} - a_{0(t1)} + (a_{1(t2)} - a_{0(t1)})P_{1(t2)} + \cdots + (a_{n(t2)} - a_{0(t1)})P_{n(t2)} \] (7)

The two-regression method was introduced by Thacker (2012) as an improvement in regression-based methods. The region of study is first divided into sub-regions since the empirical relationships between DIC and other environmental variables vary spatially (Thacker, 2012). MLRs are investigated between DIC and other measured variables (predictors) using a stepwise technique. The procedure is applied for each sub-region using all data from the repeat surveys within the period to be investigated, resulting in an optimal MLR for each sub-region (similar to Eq. 6). A linear regression with time is established for the residuals (observed DIC - predicted DIC) of the regional fits, which directly gives \( \Delta C_{\text{ANT}} \) averaged over the space-time in each sub-region. The purpose of the first MLR is to remove the natural variability of DIC, leaving the anthropogenic signal and noise (random variability) in the residuals and the second MLR is used to separate the anthropogenic signal from the noise.

We applied both methodologies within the different water mass layers separated by \( \gamma^n \) used as sub-regions. The predictor variables of \( \theta \), \( S \), \( \sigma_0 \), nitrate (NO\(_3\)), SiO\(_4\) and AOU were used for the MLR procedures. The three methodologies estimate similar rates of increase in \( C_{\text{ANT}} \) for most water mass layers (Table 7). In the STCW layer, the value of \( \Delta C_{\text{ANT}} \) (eMLR) is higher than our maximum estimate of \( \frac{\partial C_{\text{ANT}}}{\partial t} \) and the value obtained from the two-regression method. The eMLR method is less suitable for the upper layers of the ocean subject to high seasonal to interannual variability, such as the STCW layer, resulting in large residuals that bias the regression (Friis et al., 2005). For the AAIW layer, the increase of \( C_{\text{ANT}} \) estimated by the two-regression method is half the increase that is established by our method and the eMLR method. The lower value of the trend estimated by the two-regression method is due to the fact that the two-regression method uses stepwise MLR. This means that the two-regression method only considers those predictors that give the best fit while the eMLR method is forced to consider all the predictors in the fit. This is also the cause of the low RMSE obtained with the two-regression method compared to both our trends and those obtained by the eMLR method. For deep to bottom layers, the two-regression method estimates a small increase of \( C_{\text{ANT}} \) similar to the one found in this study for the AABW layer (Table 7) that can be considered negligible given the resolution of the back-calculation method (± 6 \( \mu \)mol kg\(^{-1}\)). The eMLR method finds increases of \( C_{\text{ANT}} \) (with relatively high uncertainties) higher than the two-regression method that over the 16-year period also give values of \( C_{\text{ANT}} \) change lower than the resolution of our back-calculation method (although close to it for the LCDW layer, ~ 5 \( \mu \)mol kg\(^{-1}\) for the 16-year period).

### 6.2 Circulation and biological processes at steady state.

The back-calculation method assumes the circulation of the ocean and the biological processes are in steady state. The contribution of non-linear mixing is unknown and some of the changes in DIC in the water mass layers could be erroneously included in the estimates of \( C_{\text{ANT}} \) rather than as a non-anthropogenic change in DIC. The non-steady state of the circulation in our analysis is included to some extent through the changes in \( C\text{DIS}^\pi \) (Eqs. (3, 4), which is solved by the OMP analysis, which is subjected to the limitations of quantifying the mixing mostly through thermohaline changes in the water masses (section A.2 of the Appendix).
For biological processes, remineralization rates are usually considered to be in steady state (Sarmiento et al., 1992). Climate change has been suggested as potentially driving changes in carbon fixation and export that can influence the uptake of CO₂ by the oceans (Falkowski et al., 1998). Pahlow and Riebesell, (2000) first suggested that decadal changes in remineralization rates occurred in the deep waters of the Northern Hemisphere, although this is still a matter of debate (e.g., Li and Peng, 2002; Najjar, 2009).

Metzl et al. (1999) and Shadwick et al. (2015) observed that the uptake of CO₂ over the sub-Antarctic zone (SAZ, between the SAF and the STF) in summer is mostly controlled by biological processes. If a change in remineralization rates has occurred, i.e. the changes in $DIC^{BIO}$ are due to biological effects, a change in nutrient concentrations of the water masses would be expected. The detection of long-term trends of nutrients at upper layers of the ocean can be masked by short time scale physical processes such as changes in the mixed layer depth, mesoscale activity and advection (Sambrotto and Mace, 2000; Rintoul and Trull, 2001; Sallée et al., 2010). We did not find a measurable trend in nutrient concentrations for any of the layers in the period 1995-2011, except for an increase of SiO₄ over the Antarctic Divergence (Table 4) that is most likely due to the upwelling of SiO₄-rich deep waters. We cannot confidently assign the changes of $DIC^{BIO}$ and its impact on $\frac{\partial C_{ANT}}{\partial t}$ in upper layers (Table 5) to any particular process and, instead we provide a range of values for $\frac{\partial C_{ANT}}{\partial t}$ (Table 4). For the scenario of intensified upwelling, the increase in low-O₂ and DIC-rich waters would increase the content of DIC in subsurface waters, leading (at least for the AASW and AASW$_{upw}$ layers) to values of $\frac{\partial C_{ANT}}{\partial t}$ closer to the lower limit of the range (i.e., $\frac{\partial DIC}{\partial t} >> \frac{\partial C_{ANT}}{\partial t}$, Table 4).

In deep layers of the section, the increase of DIC is not explained by the long-term change of any of the terms in Eq. (1), which is other implication of considering the circulation in steady state. The differences found in the increase of DIC in the UCDW layer north and south of the SAF (section 4.3, Fig. 4) add consistency to the idea of the advection of older waters to the section. Considering these differences we can assign a change in DIC of at least $\sim$0.20 ± 0.02 μmol kg$^{-1}$ yr$^{-1}$ (lower rate of increase of DIC found in deep-bottom layers, Table 4) due to the upwelling intensification. Since the AASW and AASW$_{upw}$ layers are the most affected by the upwelling we can correct the values of $\frac{\partial C_{ANT}}{\partial t}$ in these layers for this effect (trends with ** in Table 4).

### 6.3 Changes in the rates of export of particulate organic carbon and silicate from surface layers.

We assume that the export of particulate organic carbon (POC) from upper ocean layers (STCW, AASW and AASW$_{upw}$) and remineralization in the water column was constant between 1995 and 2011. The high-latitudes are considered important in terms of POC export, mostly because these areas are dominated by large phytoplankton, in particular diatoms (Buesseler, 1998; Sambrotto and Mace, 2000), and rapid carbon export to deep waters from phytoplankton blooms is possible (DiTullio et al., 2000; Lourey and Trull, 2001). The POC exported is remineralized to DIC below the mixed layer (Wassman et al., 1990; Asper and Smith, 1999; Trull et al., 2001a; Fripiat et al., 2015).
Our results show that the increase of DIC in mode and intermediate waters is fully explained by the uptake of atmospheric CO₂, which could indicate that there was no detectable change in the rate of export of POC over the 1995-2011 period. Estimates of POC export in the SAZ (~3800 m), the SAF (~3100 m) and the PFZ (~1500 m) using moored sediment traps near the section (Trull et al., 2001b) were 0.5, 0.8 and 1.0 g C m⁻² yr⁻¹, respectively. If all this POC is fully remineralized in the UCDW layer (with a mean thickness of 2000m), we obtain a range of 0.02 – 0.04 μmol kg⁻¹ yr⁻¹ for the maximum increase of DIC due to the export of POC. This increase is close to the uncertainty of the total DIC increase estimated for the UCDW layer, which means that in order to generate an increase in DIC similar to that found in the UCDW layer, the rate of POC export should be ~10 times higher than the observed rates. This change should be certainly noticeable in \(\frac{\partial \text{DIC}}{\partial t}\) in surface waters but most probably in deep waters as well, which we do not see.

The observed increase of SiO₄ in deep and bottom layers of the ocean is consistent with the transport of SiO₄-rich older waters to the section. For UCDW, the increase of SiO₄ north of the SAF is higher than at southern latitudes (0.31 ± 0.08 μmol kg⁻¹ yr⁻¹ respect to 0.19 ± 0.04 μmol kg⁻¹ yr⁻¹, not shown). Nelson et al., (1995) observed a lower dissolution rate of diatom dominated SiO₄ exported in high-latitudes regions compared to lower latitudes. Nelson et al. (1995) estimated a mean silica production rate of 0.7 – 1.2 mol Si m⁻² yr⁻¹ for regions over diatomaceous sediments and concluded that 15 – 25 % of the silica produced in the upper ocean accumulates in the seabed. Of the silica produced in the mixed layer of the upper ocean layers at least 50% is believed to dissolve in the upper 100 m of the water column (e.g. Nelson et al., 1991; DeMaster et al., 1992). For deep waters, the production rates of Nelson et al. (1995) could result in a mean increase of SiO₄ of 0.08 – 0.14 μmol kg⁻¹ yr⁻¹ in the water column (~3400 m, mean depth). The maximum value of this increase could explain the trends of SiO₄ found for the AABW layer (Table 4), but 32-48 % of the increase of SiO₄ in deep bottom layers is not explained by the remineralisation of exported silica and is most likely the result of the advection of older SiO₄-rich waters to the section.

6.4 Stoichiometric ratios for biological processes.

The back-calculation method and the OMP analysis assume constant stoichiometric ratios for remineralization. The theoretical Redfield ratios (Redfield, 1934; 1958) are usually considered as a mean for the whole ocean, although they can vary from the theoretical value due to changes in phytoplankton species composition, the food-web structure and nutrient availability (Martiny et al., 2013).

We carried out a sensitivity analysis on the Redfield ratios following Álvarez et al., (2014), to obtain values of stoichiometric ratios for the section. A battery of OMP analyses were done with varying values of \(R_N\) between 9 and 10 in increments of 0.2, \(R_P\) between 120 and 145 in increments of 5, and \(R_{Si}\) between 0 and 8 in increments of 2. For each variation in the stoichiometric ratios, an OMP analysis was made for each section in order to determine best-fit R values and if there were differences in time for the stoichiometric ratios along the sections. The smallest residuals (differences between the nutrients measured and the estimated by the OMP analysis) were obtained for \(R_N = 9\) and \(R_P = 125\). The residuals of SiO₄ did not change significantly for
any value of $R_{Sl}$ and we consider SiO$_4$ as a conservative variable. These results indicate $\frac{R_P}{R_N} = 13.8$, in agreement with values obtained for the region ($\frac{R_P}{R_N} \in [8-15]$, Lourey and Trull, 2001).

### 7 Conclusions.

The results of our analysis south of Tasmania over the 1995-2011 period support a scenario of intensification of upwelling in the vicinity of the Antarctic Divergence due to an increase in the westerly winds at high latitudes most probably linked to the variability of the SAM. The intensification of the upwelling favours the advection of older waters to deep-bottom layers of the section where we found net increase in DIC over the 16-year period. The enhanced upwelling causes the eventual entrainment of low-O$_2$ and DIC-rich waters into upper layers, explaining the trends of decreasing O$_2$ and increasing non-anthropogenic DIC found in surface waters close to the Antarctic Divergence. This scenario also implies the intensification of the convergence north of the SAF, implying a more efficient ventilation of the SAMW and AAIW layers and thus an efficient uptake of atmospheric CO$_2$ by these layers. The enhanced upwelling lowers the uptake of C$_{ANT}$ in the AASW layer but the effect of ventilation more than compensates that of the upwelling allowing the increase in C$_{ANT}$ in this layer. The atmospheric warming reduces the dissolution of CO$_2$ in upper layers north of the PF, presenting increases in DIC lower than expected from the atmospheric CO$_2$ increase.

Our results rely on a limited number of sections spread every 3-7 years and can only provide a long term (decadal) average view of changes in water masses. More surface observations and repeat deep ocean sections are needed to help resolve interannual changes in the Southern Ocean carbon sink and to determine the main drivers and feedback to the carbon-climate system. The effort to maintain hydrographic sections with CO$_2$ system measurements would also benefit from additional direct measurements of more variables of the carbon system, e.g. pH, which has not been measured on the SR03 section.

### Data availability

The section data are available through the Global Ocean Data Analysis Project (http://cdiac.ornl.gov/oceans/GLODAPv2; Key et al., 2015; Olsen et al., 2016). The original data for the different cruises were corrected following the QC recommendations in GLODAPv2.

### Acknowledgements

The SR03 section was sampled as part of the World Ocean Circulation Experiment/CO$_2$ Survey (WOCE, http://woceatlas.ucsd.edu), and more recently the Global Ocean Ship-Based Hydrographic Investigation Program (GO-SHIPS, http://www.go-ship.org). Carbon system parameters contribute to the International Ocean Carbon Coordination Project of the United Nations Intergovernmental Oceanographic Commission (IOCCP, http://www.ioccp.org). Support for measurements on
the section were provided to S. R. and B. T. by the Antarctic Climate and Ecosystems Cooperative Research Centre (ACE CRC) and the Australian Climate Change Science Program. Logistic support for the section and ship time on the RSV Aurora Australis was provided by the Australian Antarctic Division. The many scientific staff involved in the hydrographic sections and the officers and crew of the ship were critical to obtaining good quality data. We especially want to thank the work by Kate Berry and Mark Pretty for high quality DIC and TA data, and for the hydrochemistry teams and CTD watches of multiple cruises and especially Mark Rosenberg and Rebecca Cowley. The first author of this paper is a postdoctoral fellow supported by the ACE-CRC Project R2.1: Carbon Uptake and Chemical Change.

References.


Redfield, A.: The biological control of chemical factors in the environment, Am. Sci., 46, 205–221, 1958.


Uppstroom, L. R.: The boron/chloronity ratio of deep-sea water from the Pacific Ocean, Deep-Sea Research, 21, 161-162, 1974


Fig. 1. Transects of the 4 summer repeats of the G0-SHIP hydrodynamic line SR03 in the Southern Ocean south of Tasmania for the period 1995 – 2011 and main hydrography features of the region. ZC = Zeehan Current. EAC = East Australian Current. Black arrows indicate the flow of the Antarctic Circumpolar Current (ACC), with STF = Subtropical Front; SAF = Sub-Antarctic Front, PF = Polar Front and SACCF = Southern ACC Front.
Fig. 2. Distribution of DIC (a and b), $C_{\text{ANT, BC}}$ (c and d) and pH$_T$ (e and f) in the SR03 section south of Tasmania for the years 1995 (a, c, e) and 2011 (b, d, f). Red lines in plots (a) and (b), pink lines in plots (c) and (d) and green lines in plots (e) and (f) indicate the neutral surfaces that define the different water masses ($\gamma^o$). The position of the fronts (coloured triangles and grey dotted lines) in each of the cruises is also shown.
Fig. 3. Aragonite Saturation Depth ($\Omega_{\text{Ar}} = 1$) for the 1995 and 2011 cruises. (a) Distribution of ASD (blue dotted and solid lines) and $\gamma^n$ (green and yellow lines) in latitude in the SR03 section. (b) section zoomed for the first 1500 m of the water column. The position of the fronts (triangles) in 1995 (yellow) and 2011 (green) is also shown. The blue palette in the background indicates the distribution of $\Omega_{\text{Ar}}$ for the 2011-cruise.
Fig. 4. Distribution of (a and b) dissolved inorganic carbon (DIC) and (c and d) dissolved oxygen (O$_2$) in the 1995 and 2011 cruises for deep-bottom layers of the section. The location of the Sub-Antarctic Front (SAF) is also shown as well as the neutral surfaces ($\gamma^n$, white lines) limiting the UCDW, LCDW and AABW layers.
C Tables

<table>
<thead>
<tr>
<th>EXPOCODE</th>
<th>Dates</th>
<th>REF.</th>
</tr>
</thead>
<tbody>
<tr>
<td>09AR19941213</td>
<td>20/12/1994</td>
<td>1995</td>
</tr>
<tr>
<td>09AR20011029</td>
<td>29/10/2001</td>
<td>2001</td>
</tr>
<tr>
<td>09AR20080322</td>
<td>23/03/2008</td>
<td>2008</td>
</tr>
<tr>
<td>09AR20110124</td>
<td>4/01/2011</td>
<td>2011</td>
</tr>
</tbody>
</table>

Table 1. Repeats of the GO-SHIP SR03 line on board the Aurora Australis from 1995 to 2011 with the expocode from GLODAPv2 database.

<table>
<thead>
<tr>
<th>Layer</th>
<th>Definition</th>
<th>Complete Name</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>STCW</td>
<td>$\gamma^a &lt; 26.85 \ &amp; \ S \geq 34.3$</td>
<td>Subtropical Central Water</td>
<td>Rintoul (1998)</td>
</tr>
<tr>
<td>AASW</td>
<td>$\gamma^a \leq 27.7 \ &amp; \ S &lt; 34.3$</td>
<td>Antarctic Surface Water</td>
<td>Rintoul (1998); Williams et al. (2015)</td>
</tr>
<tr>
<td>AASW$_{upr}$</td>
<td>$\gamma^a \geq 27.7 \ &amp; \ depth \leq 300m$</td>
<td>Antarctic Surface Water</td>
<td>------------------------------------------------</td>
</tr>
<tr>
<td>SAMW</td>
<td>$26.85 \leq \gamma^a &lt; 27.2 \ &amp; \ S \geq 34.3$</td>
<td>Sub-Antarctic Mode Water</td>
<td>Rintoul (1998); Rintoul and Bullister (1999)</td>
</tr>
<tr>
<td>AAIW</td>
<td>$27.2 \leq \gamma^a &lt; 27.7 \ &amp; \ S \geq 34.3$</td>
<td>Antarctic Intermediate Water</td>
<td>Rintoul and Bullister (1999)</td>
</tr>
<tr>
<td>UCDW</td>
<td>$27.7 \leq \gamma^a &lt; 28.18 \ &amp; \ depth &gt; 300m$</td>
<td>Upper Circumpolar Deep water</td>
<td>Williams et al. (2015)</td>
</tr>
<tr>
<td>LCDW</td>
<td>$28.18 \leq \gamma^a &lt; 28.25$</td>
<td>Lower Circumpolar Deep Water</td>
<td>Lacarra et al. (2011); Williams et al. (2015)</td>
</tr>
<tr>
<td>AABW</td>
<td>$\gamma^a \geq 28.25$</td>
<td>Antarctic Bottom Water</td>
<td>Rintoul and Bullister (1999); Williams et al. (2015)</td>
</tr>
</tbody>
</table>

Table 2. Definition of the water mass layers between neutral surfaces ($\gamma^a$) and references used to accordingly decide the limits of the layers. We also consider limits in salinity, depth and position of the ACC fronts (see Table 3) to differentiate better some of the layers.
Table 3. Location of the ACC fronts south of Tasmania for each of the cruises following the definitions of Sokolov and Rintoul (2002) for hydrographic data. * The range in the location of the SAF for the 2011-cruise could be related to a diversification of the PF or SAF into different jets (Sokolov and Rintoul, 2009) but also to crossing the meander of the front twice.

<table>
<thead>
<tr>
<th>Cruise</th>
<th>STF</th>
<th>SAF</th>
<th>PF</th>
<th>SACC</th>
</tr>
</thead>
<tbody>
<tr>
<td>1995</td>
<td>~45.4’S</td>
<td>~51.8’S</td>
<td>~55.2’S</td>
<td>~63.8’S</td>
</tr>
<tr>
<td>2001</td>
<td>~46.4’S</td>
<td>~50.1’S</td>
<td>~54.5’S</td>
<td>~63.7’S</td>
</tr>
<tr>
<td>2008</td>
<td>~46.4’S</td>
<td>~52.2’S</td>
<td>~54.6’S</td>
<td>~63.5’S</td>
</tr>
<tr>
<td>2011</td>
<td>~45’S</td>
<td>~52.16-50.5’S *</td>
<td>~54.5’S</td>
<td>~63.4’S</td>
</tr>
</tbody>
</table>

Table 4. Trends in the water mass layers for the period 1995-2011 of dissolved inorganic carbon (\(\partial\text{DIC}/\partial t\)), anthropogenic carbon (\(\partial\text{CANT}/\partial t\)), silicate (\(\partial\text{SiO}_4/\partial t\)) total pH (\(\partial\text{pHT}/\partial t\)), aragonite saturation (\(\partial\Omega_{Ar}/\partial t\)) and % of change in the aragonite saturation (% \(\partial\Omega_{Ar}/\partial t\)). RMSE = root mean square error. * Trends in C\text{ANT} for the AASW and AASW\text{upw} layers considering an approximate value for the increase in DIC due to the advection of old deep waters to the section (see section 4.4.1 in the text). ** The value of \(\partial\text{CANT}/\partial t\) in the AABW layer is considered negligible because it falls below the accuracy of the back-calculation method.
Table 5. Trends of anthropogenic carbon from estimates of the back-calculation method and of the terms DIC^{BIO} and DIC^{π} (see section 3.2 of the text). RMSE = root mean square error. Bottom: schematic of the estimation of the upper and lower limits of $\frac{\partial C_{ANT}}{\partial t}$ (see section 3.3).
Table 6. Approximated rates of $C_{\text{ANT}}$ storage based on the trends from Table 4.

<table>
<thead>
<tr>
<th>Layers</th>
<th>Thickness (m)</th>
<th>$C_{\text{ANT}}$ storage rates (mol m$^{-2}$ yr$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>STCW</td>
<td>230</td>
<td>[0.17 ± 0.06 - 0.22 ± 0.08]</td>
</tr>
<tr>
<td>AASW</td>
<td>280</td>
<td>[0.10 ± 0.04 - 0.24 ± 0.07]</td>
</tr>
<tr>
<td>AASW$_{upw}$</td>
<td>275</td>
<td>[0 - 0.11 ± 0.04]</td>
</tr>
<tr>
<td>SAMW</td>
<td>450</td>
<td>0.43 ± 0.14</td>
</tr>
<tr>
<td>AAIW</td>
<td>900</td>
<td>0.39 ± 0.14</td>
</tr>
<tr>
<td>SR03 section</td>
<td>---</td>
<td>0.30 ± 0.24</td>
</tr>
</tbody>
</table>

Table 7. Comparison between $\frac{\partial C_{\text{ANT}}}{\partial t}$ from the present study and the values of $\Delta C_{\text{ANT}}$ from the two-regression and eMLR methods.

<table>
<thead>
<tr>
<th>Layer</th>
<th>$\frac{\partial C_{\text{ANT}}}{\partial t}$ (µmol kg$^{-1}$ yr$^{-1}$) 1995-2011</th>
<th>$\Delta C_{\text{ANT}}$ (Two-regression) (µmol kg$^{-1}$ yr$^{-1}$) 1995-2011</th>
<th>$\Delta C_{\text{ANT}}$ (eMLR) (µmol kg$^{-1}$ yr$^{-1}$) 1995-2011</th>
</tr>
</thead>
<tbody>
<tr>
<td>STCW</td>
<td>[0.71 ± 0.08 - 0.93 ± 0.08]</td>
<td>0.72 ± 0.04, RMSE=5</td>
<td>1.21 ± 0.09, RMSE=3</td>
</tr>
<tr>
<td>AASW</td>
<td>[0.35 ± 0.14 - 0.85 ± 0.21]</td>
<td>0.61 ± 0.04, RMSE=6</td>
<td>0.67 ± 0.10, RMSE=10</td>
</tr>
<tr>
<td>AASW$_{upw}$</td>
<td>[0 - 0.41 ± 0.16]</td>
<td>0.20 ± 0.03, RMSE=3</td>
<td>0.30 ± 0.04, RMSE=3</td>
</tr>
<tr>
<td>SAMW</td>
<td>0.92 ± 0.09, RMSE=10</td>
<td>0.94 ± 0.05, RMSE=5</td>
<td>0.96 ± 0.09, RMSE=10</td>
</tr>
<tr>
<td>AAIW</td>
<td>0.42 ± 0.06, RMSE=10</td>
<td>0.28 ± 0.02, RMSE=4</td>
<td>0.43 ± 0.02, RMSE=7</td>
</tr>
<tr>
<td>UCDW</td>
<td>-----</td>
<td>0.09 ± 0.01, RMSE=2</td>
<td>0.15 ± 0.09, RMSE=3</td>
</tr>
<tr>
<td>LCDW</td>
<td>-----</td>
<td>0.08 ± 0.01, RMSE=2</td>
<td>0.28 ± 0.15, RMSE=3</td>
</tr>
<tr>
<td>AABW</td>
<td>** 0.07 ± 0.01, RMSE=2</td>
<td>0.09 ± 0.01, RMSE=1</td>
<td>0.18 ± 0.08, RMSE=2</td>
</tr>
</tbody>
</table>

Table 7. Comparison between $\frac{\partial C_{\text{ANT}}}{\partial t}$ from the present study and the values of $\Delta C_{\text{ANT}}$ from the two-regression and eMLR methods.

RMSE= root mean square error. ** The value of $\frac{\partial C_{\text{ANT}}}{\partial t}$ in the AABW layer is considered negligible because it falls below the accuracy of the back-calculation method.