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Influence of microbial community composition and metabolism on air-sea ΔpCO_2 variation off the western Antarctic Peninsula

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ABSTRACT: We studied CO₂ and O₂ dynamics in the western Antarctic Peninsula (WAP) waters in relation to (1) phytoplankton biomass, (2) microbial community primary production and respiration, and (3), for the first time, phytoplankton composition, during summer and fall in 3 consecutive years (2002, 2003 and 2004). The areal average of ΔpCO_2 (the difference between surface seawater and atmospheric partial pressure of CO_2) for the 3 yr was significantly negative (-20.04 \pm 44.3 µatm, p < 0.01) during the summer to fall period in the region, possibly indicating a CO₂ sink. In the southern WAP (i.e. south of Anvers Island), ΔpCO_2 was significantly negative (-43.60 \pm 39.06 µatm) during fall. In the northern WAP (north of Anvers Island), ΔpCO_2 values showed a more complex distribution during summer and fall (-4.96 ± 37.6 and 21.71 ± 22.39 µatm, respectively). Chlorophyll a (chl a) concentration averaged $1.03 \pm 0.25 \text{ µg l}^{-1}$ and was higher in the south of the peninsula. Phytoplankton composition influenced chl a concentration with higher and lower values for diatom- and phytoflagellate-dominated communities, respectively. A significant negative correlation existed between chl a and ΔpCO_2 . From incubation experiments performed in the northern WAP, respiration was low (averaging 5.1 mmol $O_2 m^{-3} d^{-1}$), and the net community production (NCP) correlated negatively with ΔpCO_2 and positively with $\%O_2$ saturation. However, despite the high NCP values measured, ΔpCO_2 was significantly positive in the northern WAP during the summer to fall period. Strong mixing and lower chl a concentration may explain this result. In contrast, ΔpCO_2 was significantly negative in the southern WAP, possibly because of high surface water chl a concentration.

KEY WORDS: CO_2 sinks \cdot CO_2 sources \cdot Phytoplankton biomass \cdot Phytoplankton composition \cdot Primary production \cdot Community respiration

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INTRODUCTION

The Southern Ocean contributes up to 15% of the global ocean primary production (Huntley et al. 1991) and has been considered among the most

important CO_2 sinks on Earth, being responsible for 24% (i.e. 0.39 Pg C yr⁻¹) of the global ocean CO_2 uptake (Takahashi et al. 2002). However, Takahashi et al. (2009) recently showed that the Southern Ocean CO_2 sink is smaller than previously thought

(0.05 Pg C yr⁻¹, i.e. 4% of the global ocean CO₂ uptake), in agreement with views developed by atmospheric scientists (Bousquet et al. 2000, Peylin et al. 2002). In addition, over the last decades, the Southern Ocean CO₂ sink seems to have decreased in capacity (Metzl 2009), probably because of global warming (Le Quéré et al. 2007, 2010). However, coastal and shelf waters, such as the western Antarctic Peninsula (WAP) and the Ross Sea, are among the most productive waters of the Southern Ocean (Ferreyra et al. 2004, Ducklow et al. 2006) and may still represent important CO₂ sinks (Agustí et al. 2004, Arrigo et al. 2008).

Within the Southern Ocean, the WAP has experienced some of the most rapid warming on Earth over the last 50 yr (Marshall et al. 2002, Vaughan et al. 2003, Turner et al. 2005). This warming has affected both air (Turner et al. 2005) and sea temperatures (Gille 2002, 2008, Meredith & King 2005, Whitehouse et al. 2008), and, along with increasing upwelling of warm Upper Circumpolar Deep Water onto the WAP shelf (Martinson et al. 2008), has triggered a change in sea-ice dynamics (i.e. a decrease in the extent and duration of sea-ice cover; Stammerjohn et al. 2008), all of which may have affected its potential as a CO_2 sink through physical, chemical, and biological processes.

Polar waters are considered to be rectified (1-way) annual CO_2 sinks because sea ice is present during winter when production is at a minimum, and absent during summer when production is at a maximum (Yager et al. 1995, Ducklow et al. 2007). This represents a rectification of the typical (ice-free, low latitude) seasonal cycle of air–sea CO_2 flux. Because of the observed changes in sea-ice dynamics (Stammerjohn et al. 2008), the rectified sink hypothesis for WAP waters is now being questioned (Wang et al. 2009). Indeed, if, in the future, sea ice is absent during low production periods, and therefore during times when the ocean acts as a CO_2 source, WAP waters may no longer be a rectified CO_2 sink.

Moreover, global change will continue to have an impact on marine organisms in the WAP, affecting the timing and magnitude of phytoplankton spring blooms (Montes-Hugo et al. 2009), the composition of communities at various trophic levels, such as phytoplankton (Moline et al. 2004, Montes-Hugo et al. 2009), zooplankton (Loeb et al. 1997, Atkinson et al. 2004), and top predators (Ducklow et al. 2007), and therefore the trophic interactions within the food web (Schofield et al. 2010). In turn, these effects may have an impact on the net community production (NCP, i.e. the rate of carbon fixation by photosynthesis relative to remineralization by respiration), on the transfer of carbon through the food web and on sedimentation processes. In polar ecosystems such as the WAP, air–sea CO_2 gas exchanges are driven by either physical (e.g. deep-water formation) or biological processes (e.g. primary production or respiration, Carrillo et al. 2004). Therefore, through its impact on microbial organisms, global change may also have an indirect impact on CO_2 dynamics.

The objective of the present study was to study the potential biological contribution to atmosphere– ocean CO_2 variability in the WAP. Three consecutive surveys were conducted on board the Argentinean icebreaker 'Almirante Irizar' during the austral summer and fall of 2002, 2003 and 2004. This was the first time air–sea CO_2 and O_2 exchanges were studied along the WAP (from the South Shetland Islands to Marguerite Bay) in relation to (1) phytoplankton biomass, (2) microbial community metabolism, and (3) phytoplankton composition.

MATERIALS AND METHODS

Study area

The WAP consists of 3 sub-regions (continental slope, shelf, and coastal regions, Fig. 1a; for a review of the WAP ecosystem, see Ducklow et al. 2007). These zones include the permanently open ocean zone, the sea-ice zone, and the coastal continental shelf zone (Tréquer & Jacques 1992, Smith et al. 1998). In the WAP, the coastal zone is the most productive, and the continental slope is the least productive (Ducklow et al. 2006). In addition, the dynamics of phytoplankton in the WAP are strongly subjected to the dynamics of sea ice (Garibotti et al. 2003), whose retreat brings water column stability (Garibotti et al. 2005b). For this study, we define the northern (southern) WAP as waters north (south) of Anvers Island (as defined by Montes-Hugo et al. 2009; Fig. 1a).

Sampling

Sampling was performed in the WAP on 9 occasions from January 2002 to April 2004 on board the Argentinean icebreaker 'Almirante Irizar' within the framework of the ARGAU (Programme de coopération entre la France et l'ARGentine pour l'étude de l'océan Atlantique AUstral) cooperative research program (see Table 1 and Fig. 1b for the

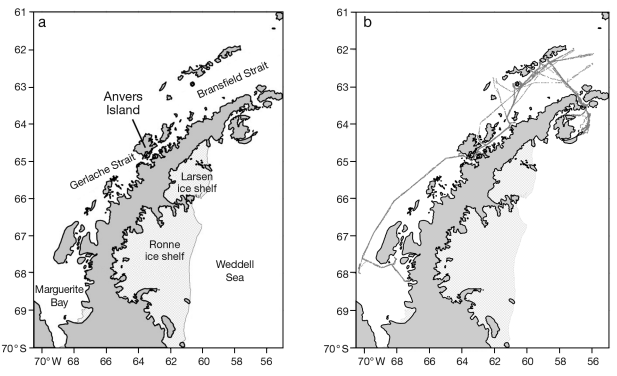


Fig. 1. (a) Western Antarctic Peninsula (WAP), and (b) ARGAU cruise itineraries (see 'Materials and methods: Sampling' for details)

dates and itineraries of the 9 cruises). The north of the peninsula was sampled both during the summer (before 21 March) and fall (after 21 March) while the south of the peninsula was only sampled during fall. The sampling methods of the ARGAU missions were described by Schloss et al. (2007) and Bianchi et al. (2005). Roughly, they consisted of a continuous acquisition and recording of meteorological, atmospheric, and seawater data. Incident photosynthetic available radiation (PAR) and the partial pressure of atmospheric CO_2 (p CO_2) were recorded continuously. The atmospheric pCO_2 was determined from an air intake placed on the bow of the ship. The air went through a flow-through equilibrator and an infrared analyzer (Siemens, Type Ultramat 5F) which was calibrated every 6 h with 3 gas standards con-

Table 1. Dates of the 9 ARGAU cruises (see Fig. 1) in the western Antarctic Peninsula in 2002, 2003, and 2004

2002	2003	2004		
3–14 February 9–22 March 14–22 April	11–16 February 6–28 March 11–27 April 4–5 May	14–16 February 17 March–7 April		

taining 270.0, 361.0, and 489.9 ppm mole fraction of CO_2 . The atmospheric pCO_2 was further corrected for warming effects using temperature data obtained from high-accuracy sensors that were placed in the equilibrator and the air intake. In addition, a surface seawater intake (placed at a depth of 9 m on the ship's exterior) allowed the continuous measurement of surface seawater pCO_2 , temperature, salinity, and in vivo fluorescence. The surface seawater pCO₂ was obtained from the infrared analyzer described above and was corrected for atmospheric pressure variations, drift, and moisture effects as in Bianchi et al. (2005). The surface seawater pCO_2 was determined with a precision of 1 µatm (Metzl et al. 1995). The difference between atmospheric and surface seawater pCO_2 was later computed as ΔpCO_2 , which contributes to CO₂ fluxes and ultimately to CO₂ sinks (if $\Delta pCO_2 < 0$) and CO_2 sources (if $\Delta pCO_2 > 0$).

Discrete samples were collected every 3 h from the seawater intake to determine dissolved oxygen concentration (following the Winkler method and using an automatic potentiometer Mettler DL21 titrator; N = 137, 114, and 62 for the 2002, 2003, and 2004 missions, respectively). Discrete samples were also collected every 3 h to determine chlorophyll *a* (chl *a*) concentration (determined with a spectrophotometer; N = 129, 116, and 63 for 2002, 2003, and 2004,

respectively), nutrient concentrations (i.e. nitrite, nitrate, phosphate, and silicate; N = 132, 114, and 62 for 2002, 2003, and 2004, respectively), and microscopic counts (performed under an inverted microscope; N = 12 and 8 for 2002 and 2003, respectively) and to perform incubation experiments to determine the NCP, gross primary production (GPP), and community respiration (*R*) of the surface planktonic community (N = 14, 19, and 7 for 2002, 2003, and 2004, respectively).

For the analysis of nutrient concentrations, duplicate samples were filtered through Whatman GF/F filters and kept frozen at -20°C until analysis. Nitrate, nitrite, and silicate concentrations were determined with a Technicon II® automatic analyzer. Phosphate concentration was determined manually following the method of Strickland & Parsons (1972). For microscopic analyses, the biovolume of cells was calculated using the geometric shapes proposed by Hillebrand et al. (1999) and corrected to account for cell shrinkage caused by fixation of samples (Montagnes et al. 1994). The carbon content of cells was calculated with 2 different carbon-to-volume ratios for diatoms (which were composed of nano- and microphytoplankton) and for phytoflagellates (an arbitrary group composed of both flagellates and dinoflagellates and including both pico- and nanophytoplankton) according to Menden-Deuer & Lessard (2000). Incubations were run in replicates with 3 clear and 2 to 3 dark bottles incubated between 6 and 24 h to measure GPP and NCP as well as *R*.

Samples for incubation were collected at a depth of 9 m using the ship's intake. The same water was further circulated through the incubation system to keep samples at in situ water temperature. In 2002 and 2003, the irradiance in the incubation system simulated PAR at the sampling depth, corresponding to $\sim 50\%$ of incident PAR as measured by a PUV-500 profiler spectroradiometer (Biospherical Instruments) in the water column before the incubation experiments. In 2004, incubation bottles were exposed to natural light and were therefore wrapped with neutral filters to reach ~50% of the sea-surface irradiance, corresponding to the irradiance measured at the sampling depth. The initial and final oxygen concentrations in the clear and dark bottles were determined as described above using the Winkler method. NCP was calculated as the difference between the initial and final oxygen concentration in the clear bottles. R was calculated as the difference between the initial and final oxygen concentration in the dark bottles. GPP was calculated as R + NCP. NCP and R were computed as daily estimates by multiplying hourly estimates by 24 following the method of Agustí et al. (2004). Daily GPP estimates were computed as GPP (d^{-1}) = GPP (h^{-1}) × DL, where DL is day length (h). Finally, the GPP:*R* ratio (i.e. the metabolic balance) was used to discriminate between autotrophic (GPP:*R* > 1) and heterotrophic communities (GPP:*R* < 1). This method had a strong analytical precision with an average coefficient of variation of 0.29% between replicates.

%O2 saturation was calculated following García & Gordon (1992) and using the solubility coefficients of Benson & Krause (1984) and the salinity and temperature data measured from the continuous sampling system. Areal averages were used to describe data averages over the sampling areas. Data were log transformed to meet the assumptions of normality and homoscedasticity, and model II linear regression analyses were performed to describe relationships between variables. In addition, 1-sample t-tests were used to determine whether $\Delta pCO2$ and $\%O_2$ saturation were significantly different from 0 and 100%, respectively, for the years 2002, 2003, and 2004 and for all 3 cruises pooled together. Finally, ANOVAs were run to determine differences between seasons and between the northern and southern WAP for all variables. Data are represented as averages \pm SE.

RESULTS

Surface water temperature and salinity

Sea surface temperature over the whole WAP showed very little interannual variability, with an areal average of 0.33 ± 0.97 °C over all 3 years (Table 2). As expected, sea surface temperature decreased significantly from February to May (r = -0.24, p < 0.01; data not shown). Moreover, during fall, the southern peninsula (with an average of -0.18 ± 0.77 °C over all 3 years) was significantly colder (p < 0.01) than the northern peninsula (with an average of 0.28 ± 0.83 °C over all 3 years; Table 2 and Fig. 2a).

The areal average of surface water salinity over the whole WAP was 33.63 ± 0.41 over all 3 years (Table 2). Surface water salinity in the northern WAP did not change significantly from summer to fall, with respective areal averages of 34.02 ± 0.21 and $34.04 \pm$ 0.25 (Table 2). However, during fall, surface water salinity was significantly higher (p < 0.01) in the north than in the south (with an areal average of 33.28 ± 0.26 in the south, Fig. 2b).

Year	—— Whole WAP ——		—— Summer north ——		——— Fall north ———		——— Fall south ———	
	Temp.	Salinity	Temp.	Salinity	Temp.	Salinity	Temp.	Salinity
2002	-0.04 ± 1.46	33.65 ± 0.39	1.22 ± 1.03	33.99 ± 0.24	0.33 ± 0.6	34.15 ± 0.19	-0.74 ± 0.4	33.31 ± 0.16
2003	0.36 ± 0.77	33.61 ± 0.39	1.02 ± 1.08	34.08 ± 0.15	0.16 ± 0.75	33.92 ± 0.27	0.3 ± 0.59	33.34 ± 0.36
2004	0.47 ± 0.64	33.64 ± 0.33	0.66 ± 0.85	34.03 ± 0.15	0.17 ± 0.85	33.97 ± 0.23	0.46 ± 0.33	33.35 ± 0.27
2002, 2003,	0.33 ± 0.97	33.63 ± 0.41	1.07 ± 1.1	34.02 ± 0.21	0.28 ± 0.83	34.04 ± 0.25	-0.18 ± 0.77	33.28 ± 0.26
and 2004								

Table 2. Areal average (± SE) of sea surface temperature (°C) and salinity (see 'Materials and methods: Sampling' for details). WAP: western Antarctic Peninsula

$\%O_2$ saturation and ΔpCO_2

Over all years, $\%O_2$ saturation ranged from 76.66 to 148.17%, with an areal average of 93.98 ± 2.1% (Table 3, Fig. 2c). In 2002, the WAP was supersaturated with respect to oxygen in the north in summer (average of 101.07 ± 2.85%) and undersaturated in both the north (93.56 ± 3.51%) and south (93.26 ± 1.4%) during fall (Table 3). In 2003 and 2004, WAP

waters were mostly undersaturated with respect to oxygen, except for a few locations in the northern WAP in summer. The $\%O_2$ saturation in the north was significantly higher during summer than during fall (p < 0.01, Table 3). In addition, $\%O_2$ saturation was significantly higher in the south than in the north during fall (p < 0.01).

The atmospheric pCO_2 averaged 372.2 \pm 0.7 μatm from 2002 to 2004, while the sea pCO_2 was much

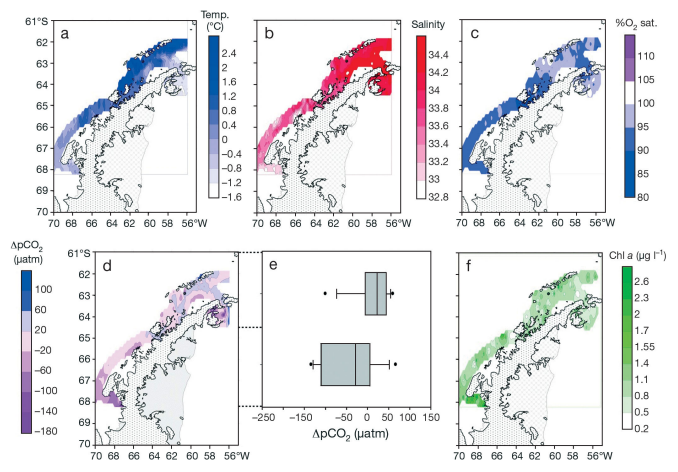


Fig. 2. (a) Sea surface temperatures (°C), (b) sea surface salinity, (c) O_2 saturation, and (d) ΔpCO_2 along the western Antarctic Peninsula (WAP) in 2002, 2003, and 2004 (combined). (e) Box plots describing the ΔpCO_2 data in the northern and southern WAP. The box plots indicate the 5th and 95th percentile (dots), the lower and upper quartiles, and the median. (f) Chl *a* along the WAP in 2002, 2003, and 2004 (combined). To avoid distortion, chl *a* values > 3 µg l⁻¹ were not used in (f)

ARGAU	Whole WAP			Summer north			
cruise	Sat O_2	ΔpCO_2	Chl a	Sat O ₂	ΔpCO_2	Chl a	
2002	96.19 ± 3.37	-27.13 ± 42.31	0.78 ± 0.28	101.07 ± 2.85	-27.88 ± 39.59	0.91 ± 0.27	
2003	91.48 ± 2.44	0.01 ± 39.20	1.08 ± 0.25	93.78 ± 3.37	18.62 ± 19.51	0.88 ± 0.4	
2004	93.33 ± 1.26	-14.03 ± 41.81	1.65 ± 0.32	95.06 ± 2.56	19.59 ± 32.07	1.41 ± 0.21	
2002, 2003, and 2004	93.98 ± 2.10	-20.04 ± 44.30	1.03 ± 0.25	98.39 ± 2.35	-4.96 ± 37.6	0.98 ± 0.24	
ARGAU		— Fall north —			— Fall south —		
cruise	Sat O ₂	ΔpCO_2	Chl a	Sat O ₂	ΔpCO_2	Chl a	
2002	93.56 ± 3.51	23.31 ± 23.17	0.42 ± 0.2	93.26 ± 1.40	-49.26 ± 30.27	1.01 ± 0.38	
2003	89.27 ± 1.66	35.59 ± 16.22	0.94 ± 0.17	94.74 ± 0.98	-31.30 ± 37.97	1.39 ± 0.48	
2004	91.20 ± 1.18	17.96 ± 18.33	1.57 ± 0.56	93.73 ± 0.55	-42.93 ± 45.8	1.79 ± 0.16	
2002, 2003, and 2004	90.99 ± 1.99	21.71 ± 22.39	0.85 ± 0.22	94.02 ± 0.54	-43.60 ± 39.06	1.31 ± 0.32	

Table 3. Areal average (\pm SE) of ΔpCO_2 (µatm), $\%O_2$ saturation (sat O_2), and chl *a* (µg l⁻¹) (see 'Materials and methods: Sampling' for details). WAP: western Antarctic Peninsula

more variable and ranged from 176.3 to 503.2 µatm. Over the 3 yr, the areal average of ΔpCO_2 in the WAP was negative (-20.04 ± 44.3 µatm, contributing to a CO_2 sink) during the summer to fall period (p < 0.01, Table 3). However, ΔpCO_2 showed important variations among years, and although variation in the data is high, it could have led the WAP to be a significant CO_2 sink in 2002 and in 2004 (areal averages of -27.13 ± 42.31 and -14.03 ± 41.81 µatm, respectively, p < 0.01) while a balanced CO_2 budget was observed in 2003 (areal average of 0.01 ± 39.2 µatm).

For all years, a south to north gradient in ΔpCO_2 seemed to exist as shown in Fig. 2d and the box plots in Fig. 2e. During summer, ΔpCO_2 was negative in the north (p < 0.01), contributing to an atmospheric CO_2 sink in 2002 only (with areal averages of $-27.88 \pm$ 39.59, 18.62 ± 19.51 , and 19.59 ± 32.07 µatm for 2002, 2003, and 2004, respectively). In addition, in the north, ΔpCO_2 was significantly lower during summer (with an average of $-4.96 \pm 37.6 \mu atm$) than during fall $(21.71 \pm 22.39 \mu atm, p < 0.01; Table 3)$. In fact, during fall, ΔpCO_2 was significantly positive in the north, which could have led to a CO₂ source for the atmosphere during all years (p < 0.01). The opposite situation was evident in the south during fall over all years (with a significantly negative ΔpCO_2 average of -43.6 \pm 39.06 µatm, p < 0.01). Therefore, ΔpCO_2 in the south could have led to a significant sink for atmospheric CO₂, while the role of the northern peninsula was more complex and depended on the season (Fig. 2d).

Chl a and phytoplankton groups

Surface water chl *a* areal average was $1.03 \pm 0.25 \ \mu g \ l^{-1}$ for all years (Table 3). Small interannual differences in chl *a* were observed, with areal aver-

ages of 0.78 ± 0.28, 1.08 ± 0.25, and 1.65 ± 0.32 µg l⁻¹ for 2002, 2003, and 2004, respectively (Table 3 and Fig. 2f). In the north, chl *a* was insignificantly higher during summer (average of 0.98 ± 0.24 µg l⁻¹) than during fall (average of 0.85 ± 0.22 µg l⁻¹, p = 0.31). In addition, with an average of 1.31 ± 0.32 µg l⁻¹, chl *a* was significantly higher in the south during fall than in the north during either summer or fall (p < 0.01).

The highest chl *a* concentrations measured in 2002 (between 17.5 and 22.3 μ g l⁻¹) were found near the coast (at Deception Island, in the Gerlache Strait, and in the north of the peninsula near the Weddell Sea) on 4, 8, and 10 February. In 2003, the highest chl *a* concentrations (between 18.3 and 26.4 μ g l⁻¹) were measured near the coast in Marguerite Bay on 14 and 15 April. In 2004, the highest chl *a* concentrations (between 12 and 29.4 μ g l⁻¹) were measured near the coast (at Deception Island and in the northern WAP near the Weddell Sea) on 15 February and 19 and 21 March.

Among the samples analyzed by microscopy, diatoms were the dominant group at the 4 stations with the highest chl a concentrations measured (between 8.04 and 22.33 μ g l⁻¹). Measurements were made at these 4 stations in February 2002 and February 2003, and were found in the northern WAP (i.e. in the Bransfield Strait and near the Weddell Sea). In fact, there was a significant positive relationship between chl a and the carbon biomass of diatoms (r = 0.88, p < 0.01, Fig. 3a). In contrast, no significant relationship was observed between chl a and the carbon biomass of phytoflagellates (Fig. 3b). Therefore, the composition of the phytoplankton community influences the chl a concentration found in surface waters of the WAP. Of the 20 stations for which phytoplankton composition was analyzed, 12 were dominated by diatoms and the other 8 by phytoflagellates. Therefore, and consistent with the literature (e.g. Garibotti et al. 2003), both

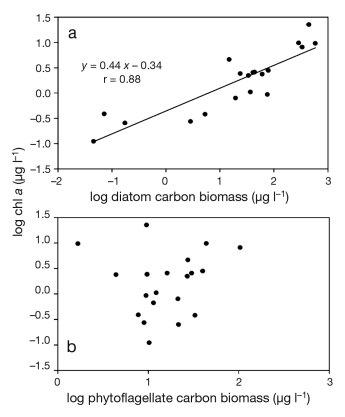


Fig. 3. Chl a and carbon biomass of (a) diatoms and (b) phytoflagellates (all log-transformed). A significant (p < 0.01) linear regression was found for chl a and diatom carbon biomass

phytoplankton groups were important in WAP waters and probably contributed to the observed spatial and temporal differences in the accumulation of chl *a*.

A significant negative correlation existed between chl *a* and ΔpCO_2 (r = -0.66, p < 0.01, Fig. 4a) when all the stations were pooled. Moreover, this relationship held true for stations dominated by diatoms (i.e. defined by Schloss et al. 2007 as stations with more than 50% of the phytoplankton biomass represented by diatoms; r = -0.68, p < 0.01; Fig. 4b) but not for stations dominated by phytoflagellates (i.e. with more than 50% of the phytoplankton biomass represented by phytoflagellates; Fig. 4c).

A positive relationship existed between chl *a* and $%O_2$ saturation (r = 0.6, p < 0.01, Fig. 4d). The correlation coefficient was greater when considering only stations dominated by diatoms (r = 0.81, p < 0.01, Fig. 4e), and there was a lack of correlation between chl *a* and $%O_2$ saturation for stations dominated by phytoflagellates (Fig. 4f). Therefore, phytoplankton composition plays a crucial role in the relationship between chl *a* and CO_2 and O_2 dynamics in the WAP.

Community metabolism

R was neither correlated with temperature (p = 0.94) nor with chl *a* (p = 0.95). The average daily *R* for all studied years was 5.2 mmol $O_2 \text{ m}^{-3} \text{ d}^{-1}$. Two of

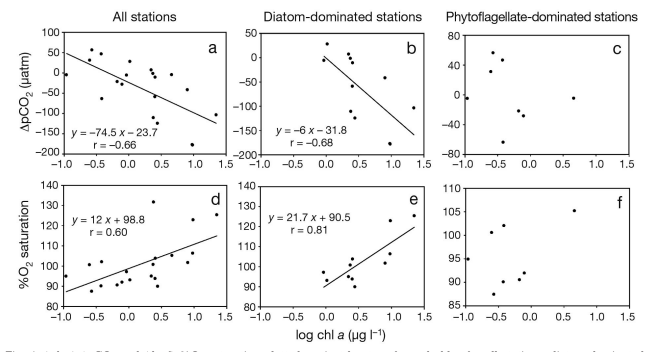


Fig. 4. (a,b,c) ΔpCO_2 and (d,e,f) $\%O_2$ saturation plotted against log-transformed chl *a* for all stations, diatom-dominated stations and phytoflagellate-dominated stations. Significant (p < 0.01) linear regressions are also plotted

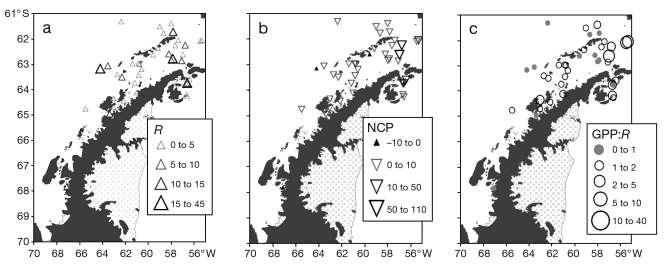


Fig. 5. (a) Community respiration (R, mmol O₂ m⁻³ d⁻¹), (b) net community production (NCP, mmol O₂ m⁻³ d⁻¹), and (c) gross primary production:respiration ratio (GPP:R, dimensionless) measured in incubation experiments in 2002, 2003, and 2004 (combined)

the highest *R* values (i.e. 22.8 and 42.1 mmol $O_2 m^{-3} d^{-1}$) were observed in 2002 and were found offshore (on the continental slope, Fig. 5a). Another high *R* value (16.8 mmol $O_2 m^{-3} d^{-1}$) was measured in 2004 in the north, in the Bransfield Strait. High *R* values were rare and were mostly found in the stations located relatively far offshore.

NCP ranged from -3.5 to 103.9 mmol O₂ m⁻³ d⁻¹. High NCP values (between 15.2 and 103.9 mmol O₂ m⁻³ d⁻¹) were generally found at the northern tip of the peninsula (in the Bransfield Strait and towards the Weddell Sea, Fig. 5b). Negative NCP values were rare and were found in Marguerite Bay, Deception Island, and on the WAP continental slope. NCP was higher than *R* for 70% of the stations (28/40 stations), showing that, for the majority of stations, plankton communities were potentially net autotrophic. Finally, NCP was significantly correlated with chl *a* (r = 0.61, p < 0.01, Fig. 6a) and GPP (r = 0.86, p < 0.01, Fig. 6b).

As described above, 70% of the stations had a GPP:R > 1, although this ratio showed a high interannual variability. In 2002 and in 2004, only 57% of the stations had a GPP:R > 1 (12/21 stations for 2002 and 2004), while in 2003, 84% of the stations had a GPP:R > 1 (16/19 stations). No seasonal differences were observed, with 68 and 72% of the stations with a GPP:R > 1 for summer and fall, respectively (15/22 and 13/18 stations, respectively). It also seems that most of the stations with GPP:R > 1 were found close to the coast (Fig. 5c). Finally, there was no difference in nutrient concentration between autotrophic (with averages of 24.4 ± 3.6, 1.8 ± 0.3, and 67.9 ± 9.3 µM for nitrate + nitrite, phosphate, and silicate, respectively)

and heterotrophic communities (with averages of 21.6 ± 7.8, 1.6 ± 0.5, and 63.4 ± 17.2 µM for nitrate + nitrite, phosphate, and silicate, respectively). However, chl *a* was usually lower in heterotrophic (average of 1 ± 0.7 µg l⁻¹) than in autotrophic communities (average of 2.2 ± 2.2 µg l⁻¹). Only 1 station, where the community was net heterotrophic (GPP:*R* = 0.33), exhibited a high chl *a* concentration (18.6 µg l⁻¹) together with low nitrate + nitrite and phosphate concentrations (3.1 and 0.4 µM, respectively).

Except for 1 station in Marguerite Bay in fall 2002 (not included in the analysis), NCP was negatively correlated with ΔpCO_2 (r = -0.42, p < 0.01, Fig. 6c) and positively correlated with $\%O_2$ saturation (r = 0.59, p < 0.01, Fig. 6d). In contrast, *R* was neither correlated with ΔpCO_2 nor with $\%O_2$ saturation (data not shown). Finally, for the 14 incubation stations for which phytoplankton species biomass was determined, GPP and the GPP:R ratio were significantly correlated with diatom biomass (r = 0.82, p < 0.01 and r = 0.68, p = 0.01 for GPP and GPP:R, respectively; Fig. 7 a,b) but not with phytoflagellate biomass (data not shown).

DISCUSSION

Over the 3 consecutive years studied (2002, 2003, and 2004) ΔpCO_2 was significantly negative during the summer to fall period (areal average of $-20.46 \pm$ 44.3 µatm), which could have led to a significant CO_2 sink in the WAP. However, during summer, ΔpCO_2 was significantly negative in the north of the peninsula in 2002 only, while during fall, it was significantly positive in the north and significantly negative

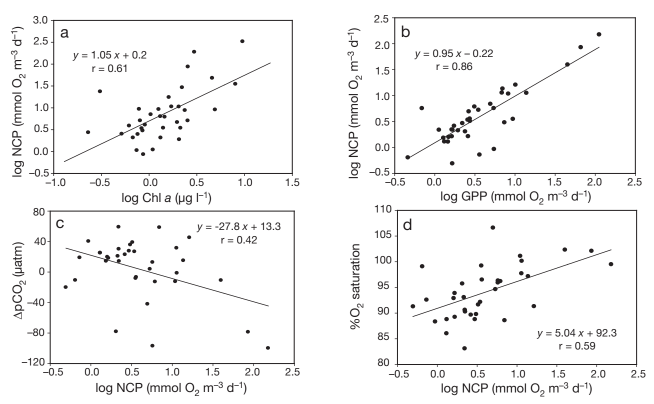
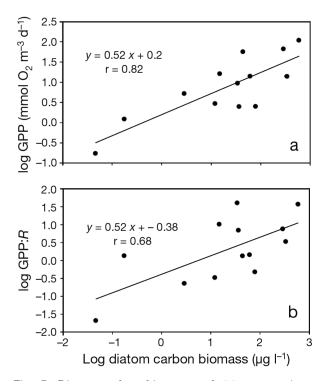


Fig. 6. Net community production (NCP) and (a) chl a and (b) gross primary production (GPP) (all log-transformed). (c) ΔpCO_2 and (d) O_2 saturation and NCP (log-transformed). Significant (p < 0.01) linear regressions are also plotted



ings are coherent with the literature, which suggests that the southern coastal WAP is a CO₂ sink during summer (Carrillo et al. 2004). These results are also consistent with the results of Carrillo & Karl (1999) and Álvarez et al. (2002) which suggest that CO_2 sinks and sources show a complex distribution in the northern WAP, but they are not entirely consistent with the suggestion of Agustí et al. (2004) that the northern WAP is a net CO₂ sink during summer because it is mainly autotrophic. The differences observed between the north and the south are reported here for the first time, and some of the possible mechanisms controlling ΔpCO_{2} , such as community composition, community metabolism (i.e. primary production and respiration), and sea surface temperature, are discussed below.

in the south, possibly leading to, respectively, CO₂

sources and sinks (Fig. 2d and Table 3). These find-

Influence of community composition on the WAP CO₂ dynamics

Fig. 7. Diatom carbon biomass and (a) gross primary production (GPP) and (b) GPP:respiration (R) ratio (all log-transformed). Significant (p < 0.01 and p = 0.01 for panels a and b, respectively) linear regressions are also plotted

A significant negative correlation existed between chl *a*, as a proxy of primary producers, and ΔpCO_2 , and a positive relationship existed between chl *a* and $%O_2$ saturation (Fig. 4), suggesting that the dynamics of these 2 gases in the WAP were influenced by primary producers. Moreover, and consistent with the findings of Schloss et al. (2007) at the Patagonian shelf off Argentina, the 2 relationships between chl a and ΔpCO_2 and $\%O_2$ saturation only held true for stations dominated by diatoms. Chl a concentration was itself influenced by the composition of the phytoplankton community, with high and low chl a concentrations found for diatom- and phytoflagellatedominated communities, respectively. In fact, in the majority of samples analyzed, diatom cells were significantly larger than phytoflagellates. This is consistent with previous findings in the literature (e.g. Montes-Hugo et al. 2008, Olguín & Alder 2011) which showed that, in the WAP, high and low chl a concentration are usually associated with the presence of large and small cells, respectively. One may therefore extrapolate and argue that large- and small-cell-dominated phytoplankton communities may lead to negative and positive ΔpCO_2 and therefore to CO_2 sinks and sources, respectively.

It has been hypothesized that the community composition of the WAP is shifting from diatom- to cryptophyte-dominated waters because of regional warming (Moline et al. 2004) and in relation to temporal changes in sea-ice dynamics (Garibotti et al. 2005a) which were observed in the WAP during the last decades (Stammerjohn et al. 2008). More specifically, Montes-Hugo et al. (2008) showed that the phytoplankton shifts from large to small cells that have taken place in the northern WAP in the last decade were probably related to changes in sea-ice anomalies. In another study, Montes-Hugo et al. (2009) showed that, because of regional warming, chl a concentration has decreased by 12% in the whole WAP during the summer (December to February) over the last 30 yr. In contrast, Montes-Hugo et al. (2008) found higher chl a and larger cells in the southern part of the peninsula. Moreover, and contrasting with observations in the north, Montes-Hugo et al. (2009) observed increases in surface chl a in the southern WAP. These authors hypothesized that the WAP was undergoing a southward displacement of species of all trophic levels, from phytoplankton to top predators, with likely greater contributions of diatoms and large cells in the south and the opposite trend in the north. This is consistent with our observations of higher chl *a* concentrations in the southern WAP (Table 3), and can potentially explain the lower ΔpCO_2 values observed in the south in this study.

Other than a direct impact on total phytoplankton biomass, a shift in phytoplankton composition and

cell size might also have other implications on carbon fluxes. Indeed, small cells, such as cryptophytes or other phytoflagellates, are believed to be responsible for less carbon export than diatoms. Large diatoms are the base of multivore and herbivore food webs which are responsible for a high export of carbon from the euphotic zone to deep waters through direct sedimentation of phytoplankton cells or the sinking of zooplankton fecal pellets (Legendre & Rassoulzadegan 1995, Armbrust 2009). In contrast, small phytoplankton cells are the base of microbial food webs for which most of the carbon is expected to be consumed within the euphotic layer (Legendre & Rassoulzadegan 1996). For example, several authors measured higher sinking rates for waters dominated by large diatoms than for waters dominated by small phytoplankton (Serret et al. 2001, Anadón et al. 2002).

Community metabolism and CO₂ dynamics

First, it should be considered that most of the incubation experiments presented here (95%) were performed in the northern WAP (north of Anvers Island, Fig. 5). The average daily respiration rate for all studied years was low (present summer to fall data of 5.2 mmol $O_2 m^{-3} d^{-1}$) which is consistent with average respiration rates of 2.65 and 6.95 mmol $O_2 \text{ m}^{-3}$ d^{-1} reported by Agustí et al. (2004) for the Bransfield and Gerlache Straits, respectively. High respiration rates were rare and were mostly found far from the coast and mostly during summer. In contrast, the net community production was high (ranging from -3.5 to 103.9 mmol $O_2 m^{-3} d^{-1}$) over all studied years when compared to previously reported estimates of NCP in the WAP, with NCP ranging from -6.29 to 35.4 mmol $O_2 \text{ m}^{-3} \text{ d}^{-1}$ in the study of Aqustí et al. (2004). In addition, NCP was correlated to chl a and to GPP (Fig. 6a,b), showing that NCP in the WAP was mainly driven by primary production rather than by respiration. This is consistent with the findings of other studies which described an uncoupling between phytoplankton production and heterotrophic respiration in Antarctic waters (Morán et al. 2002, Agustí et al. 2004, Pearce et al. 2007).

As a consequence, the northern WAP waters were mainly autotrophic, and the GPP:*R* ratio exceeded 1 at 70% of the stations, very similar to the results of Agustí et al. (2004), who found that 73% of the stations they analyzed in the northern WAP during the summer were autotrophic. In addition, there was no strong difference in the proportion of stations with a

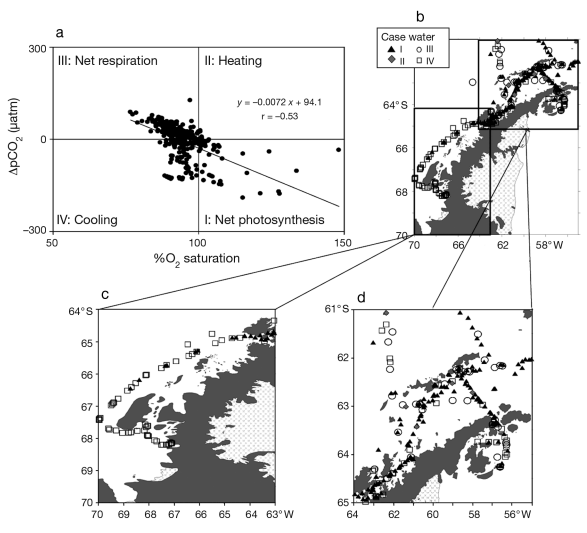


Fig. 8. (a) ΔpCO₂ and %O₂ saturation. The plot is divided into 4 quadrants. The significance of these quadrants is explained in the 'Discussion'. (b) Geographical distribution of Case I, II, III, and IV waters described in (a). (c) Geographical distribution of Case I, II, III, and IV waters within the southern portion of the western Antarctic Peninsula (WAP) and (d) geographical distribution of Case I, II, III, and IV waters within the northern WAP

GPP:R > 1 between summer and fall (with 68 and 72% of the stations, respectively). Because the northern WAP is mainly autotrophic, and because respiration seems to be uncoupled to primary production (i.e. *R* was not correlated with chl *a*, NCP, or GPP), one would expect the northern WAP to behave as a significant CO₂ sink. Indeed, NCP correlated negatively with ΔpCO_2 and positively with the %O₂ saturation (Fig. 6c,d), while R was neither correlated to ΔpCO_2 nor to the %O₂ saturation. According to the linear regression between ΔpCO_2 and NCP presented in Fig. 6c, the minimum NCP value required to yield negative ΔpCO_2 was 3.02 mmol O_2 m⁻³ d⁻¹. This value was achieved in 36% and 11% of the incubation experiments performed during the summer and fall, respectively. This is consistent with the lower and higher ΔpCO_2 values observed in the northern WAP during the summer and fall, respectively (Table 3). However, this is not consistent with the proportion of autotrophic stations sampled in the northern WAP (70%). Therefore, NCP, and more particularly primary production, seems to play a significant role in CO_2 and O_2 dynamics in the WAP, although autotrophy does not necessarily lead to negative ΔpCO_2 values in the northern WAP.

Spatial variability of CO_2 and O_2 dynamics in the WAP

In order to explain the observed discrepancies between community metabolism and the distribution of negative and positive ΔpCO_2 values and the possible mechanisms responsible for their distribution, we

studied the distribution of both ΔpCO_2 and $\%O_2$ saturation (Fig. 8). Fig. 8a shows a scatter diagram of ΔpCO_2 as a function of $\%O_2$ saturation. A significant negative correlation existed between these 2 variables (r = -0.53, p < 0.01), confirming the hypothesis of Agustí et al. (2004) and Álvarez et al. (2002) that photosynthesis and respiration, rather than other physical processes, are the driving processes in O_2 and CO₂ dynamics in the WAP. This figure was inspired and modified from the work of Carrillo et al. (2004), who studied the relationship between $%O_2$ saturation and the surface seawater fugacity of CO₂ (which is similar to pCO_2) relative to the atmosphere (fCO_2) in the southern WAP waters. In Fig. 8a, the graph was divided into 4 quadrants: Quadrant I (simultaneous positive ΔpCO_2 and O_2 undersaturation) implies respiration as the process controlling seawater pCO_2 and $\%O_2$ undersaturation, Quadrant II (simultaneous positive ΔpCO_2 and O_2 supersaturation) implies heating as the process controlling seawater pCO₂ and %O₂ supersaturation; Quadrant III (simultaneous negative ΔpCO_2 and O_2 supersaturation) implies photosynthesis as the main process controlling seawater pCO_2 and $\%O_2$ supersaturation; and Quadrant IV (simultaneous negative ΔpCO_2 and O₂ undersaturation) implies cooling as a the process controlling seawater pCO₂ and %O₂ undersaturation.

From this figure, a clear demarcation appeared between the north and the south (i.e. north and south of Anvers Island; Fig. 8b), with southern waters mainly represented by case IV waters and northern waters represented by case I, III, and IV waters. Indeed, ΔpCO_2 was negative in 91% of the stations in the southern peninsula, consistent with the findings of Carrillo et al. (2004), who showed that most of the coastal southern WAP was represented by case III and case IV waters (i.e. with negative ΔpCO_2). However, and although ΔpCO_2 was negative in 91% of the stations in the south, the $%O_2$ saturation was above 100% in only 1.5% of the stations (Fig. 8a). Because primary production plays a significant role in CO₂ dynamics in WAP waters, one would expect southern WAP waters to be supersaturated with respect to O₂. However, as described previously, surface water was particularly cold in the south, a fact that may have increased oxygen solubility and would explain the oxygen undersaturation found in these waters. The %O₂ saturation was positively correlated with sea surface temperature (r = 0.45, p < 0.01, data not shown). Moreover, the relatively high chl a concentration found in the south (Fig. 2f, Table 3) may indicate that primary production was strong even though incubation experiments were not performed in this part of the peninsula. Therefore, even though most of the southern WAP waters were undersaturated with respect to O_2 , primary production may still have been the main driver for the significant negative ΔpCO_2 values observed.

North of Anvers Island, only 35% of the stations showed negative ΔpCO_2 values (Fig. 8d). Of these, 52% were supersaturated with respect to O_2 (quadrant III) and the other 48 % were undersaturated with respect to O₂ (quadrant IV). There was no clear distribution pattern of any of the 4 quadrant types in the northern WAP (Fig. 8d). Moreover, ΔpCO_2 was mainly positive in the northern WAP as shown by the dominance (65%) of quadrant I waters, possibly leading to a significant CO_2 source in the northern WAP. Therefore, the role of the northern WAP with regards to CO₂ was much more complex than and not as strong as the southern WAP waters, consistent with the results presented in Table 3 and with the results of Carrillo & Karl (1999) and Álvarez et al. (2002), who described a complex distribution of CO₂ sinks and sources in the northern WAP.

Fig. 9 describes the possible influence of community metabolism (represented by the GPP:*R* ratio) on the distribution of ΔpCO_2 and $\%O_2$ saturation in the northern WAP, which will ultimately contribute to the distribution of CO₂ sinks and sources. Unexpectedly, there was no clear distribution pattern in GPP:*R* from *in vitro* measurements regarding ΔpCO_2 and $\%O_2$ saturation. In fact, autotrophic and heterotrophic

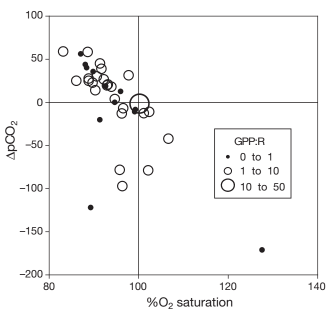


Fig. 9. Distribution of the gross primary production:respiration (GPP:*R*) ratio in relation to ΔpCO_2 and $\%O_2$ saturation

communities (with GPP:*R* ratios > 1 and < 1, respectively) were both found in Quadrants I, III, and IV with no clear pattern. Therefore, the community metabolism balance, represented by the GPP:*R* ratio, cannot explain by itself the distribution of negative ΔpCO_2 values in the northern WAP and suggests that the relationship between community metabolism and ΔpCO_2 and $\%O_2$ saturation in the WAP is more complex than previously thought (Fig. 9).

In the WAP, mixing is believed to play a major role in primary production (Mitchell & Holm-Hansen 1991). In fact, surface water salinities measured in the northern WAP were much higher than in the south (Fig. 2b and Table 2), suggesting that mixing was more important in the north, therefore limiting primary production and the drawdown of CO₂ from the atmosphere. Schloss et al. (2002) showed that King George Island phytoplankton communities were potentially very productive, although they rarely met the adequate growth conditions in situ. In a mesocosm experiment, it took 12 d before the exponential growth of phytoplankton started and finally reached levels of chl *a* as high as $36 \ \mu g \ l^{-1}$ with complete nutrient exhaustion (Schloss et al. 2002), showing the strong primary production potential of WAP waters. Mixing is caused by winds which can be very intense in the WAP (Smith et al. 2008). Moreover, associated with regional warming, there has been a 15 to 20% increase in the strength of westerly winds in the WAP since the 1960s (Orr et al. 2004), which may have limited the biological uptake of CO₂ (Le Quéré et al. 2010). This strengthening of winds over the last decades in the WAP may explain the complex distribution of ΔpCO_2 observed in the north in this study.

In addition, nutrient limitation may have explained the discrepancies found between the community metabolism balance (i.e. GPP: R ratio) and the distribution of ΔpCO_2 and $\%O_2$ saturation in the WAP. For example, 1 station sampled for an in vitro experiment showed a high chl a concentration (18.6 μ g l⁻¹) together with low nitrate + nitrite and phosphate concentrations (3.1 and 0.4 μ M, respectively) and low ΔpCO_2 (-171 µatm). However, this community had a low NCP (-3.5 mmol $O_2 \text{ m}^{-3} \text{ d}^{-1}$) and a fairly high R value (8.8 mmol $O_2 \text{ m}^{-3} \text{ d}^{-1}$) together with a GPP:*R* of 0.33, representative of a heterotrophic community. It is probable that this station was sampled after a phytoplankton bloom took place, which consumed nitrite, nitrate, and phosphate present in the water column. The high respiration rate measured probably reflects the consumption of organic matter by heterotrophs (i.e. bacteria and micro- and mesozooplankton). However, this particular scenario was only found once, and nutrient concentrations were high in the remaining incubation experiments, suggesting that physical factors (e.g. water column mixing) limited *in situ* primary production rather than nutrient concentrations.

As pointed out by Schloss et al. (2007), metabolic activities such as respiration and primary production rely on different time and spatial scales than CO2 and O₂ exchanges between the atmosphere and the ocean. Indeed, gas exchanges between the ocean and the atmosphere are processes occurring over weeks, whereas incubation experiments reflect short-term metabolic activities. In addition, recent studies have shown that incubation experiments performed in bottles may underestimate NCP and GPP when compared to non-incubation methods (e.g. Quay et al. 2010), although the authors did not conclude which method gives the more realistic results. This possible limitation needs to be considered for the present study, and simultaneous incubation and non-incubation estimations should be done simultaneously in order to solve this question. Therefore, more studies are needed to determine the relationship between CO₂ and O₂ exchanges between the atmosphere and the ocean and the community metabolism in WAP waters.

Finally, although the role of the WAP and its community composition in CO₂ and O₂ dynamics was determined, these results only concerned summer and fall. However, due to the increasing disappearance of sea ice along with regional warming (Stammerjohn et al. 2008), the seasonal rectification hypothesis of Yager et al. (1995) may require reconsideration. Some recent studies performed during winter to spring in the WAP (e.g. Wang et al. 2009) additionally point towards a weak CO₂ source during winter but a moderate sink during spring. Ideally, as described by Montes-Hugo et al. (2010), the study of CO₂ dynamics in relation to sea-ice dynamics over periods of several years is needed to be able to adequately assess the role of WAP waters in CO₂ dynamics.

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LITERATURE CITED

- Agustí S, Satta MP, Mura MP (2004) Summer community respiration and pelagic metabolism in upper surface Antarctic waters. Aquat Microb Ecol 35:197–205
- Álvarez M, Ríos AF, Rosón G (2002) Spatio-temporal variability of air-sea fluxes of carbon dioxide and oxygen in the Bransfield and Gerlache Straits during Austral summer 1995–96. Deep-Sea Res II 49:643–662
- Anadón R, Alvarez-Marqués F, Fernández E, Varela M, Zapata M, Gasol JM, Vaqué D (2002) Vertical biogenic particle flux during Austral summer in the Antarctic Peninsula area. Deep-Sea Res II 49:883–901
- Armbrust EV (2009) The life of diatoms in the world's oceans. Nature 459:185–192
- Arrigo KR, van Dijken G, Long M (2008) Coastal Southern Ocean: a strong anthropogenic CO₂ sink. Geophys Res Lett 35:L21602 doi:10.1029/2088GL035624
- Atkinson A, Siegel V, Pakhomov E, Rothery P (2004) Longterm decline in krill stock and increase in salps within the Southern Ocean. Nature 432:100–103
- Benson BB, Krause DJ (1984) The concentration and isotopic fractionation of oxygen dissolved in freshwater and seawater in equilibrium with the atmosphere. Limnol Oceanogr 29:620–632
- Bianchi AA, Bianucci L, Piola AR, Pino DR, Schloss I, Poisson A, Balestrini CF (2005) Vertical stratification and air-sea CO₂ fluxes in the Patagonian shelf. J Geophys Res 110: C07003 doi:10.1029/2004JC002488
- Bousquet P, Peylin P, Ciais P, Le Quere C, Friedlingstein P, Tans PP (2000) Regional changes in carbon dioxide fluxes of land and oceans since 1980. Science 290: 1342–1347
- Carrillo CJ, Karl DM (1999) Dissolved inorganic carbon pool dynamics in northern Gerlache Strait, Antarctica. J Geophys Res 104:15873–15884
- Carrillo CJ, Smith RC, Karl DM (2004) Processes regulating oxygen and carbon dioxide in surface waters west of the Antarctic Peninsula. Mar Chem 84:161–179
- Ducklow HW, Fraser W, Karl DM, Quetin LB and others (2006) Water-column processes in the West Antarctic Peninsula and the Ross Sea: interannual variations and foodweb structure. Deep-Sea Res II 53:834–852
- Ducklow HW, Baker K, Martinson DG, Quetin LB and others (2007) Marine pelagic ecosystems: the West Antarctic Peninsula. Philos Trans R Soc Lond B Biol Sci 362:67–94
- Ferreyra G, Schloss I, Demers S (2004) Rôle de la glace saisonnière dans la dynamique de l'écosystème marin de l'Antarctique: impact potentiel du changement climatique global. Vertigo 5:1–11
- García HE, Gordon LI (1992) Oxygen solubility in seawater: better fitting equations. Limnol Oceanogr 37:1307–1312
- Garibotti IA, Vernet M, Ferrario ME, Smith RC, Ross RM, Quetin LB (2003) Phytoplankton spatial distribution pat-

terns along the western Antarctic Peninsula (Southern Ocean). Mar Ecol Prog Ser 261:21–39

- Garibotti IA, Vernet M, Ferrario ME (2005a) Annually recurrent phytoplanktonic assemblages during summer in the seasonal ice zone west of the Antarctic Peninsula (Southern Ocean). Deep-Sea Res I 52:1823–1841
- Garibotti IA, Vernet M, Smith RC, Ferrario ME (2005b) Interannual variability in the distribution of the phytoplankton standing stock across the seasonal sea-ice zone west of the Antarctic Peninsula. J Plankton Res 27: 825–843
- Gille ST (2002) Warming of the Southern Ocean since the 1950s. Science 295:1275–1277
- Gille ST (2008) Decadal-scale temperature trends in the Southern Hemisphere ocean. J Clim 21:4749–4765
- Hillebrand H, Dürselen CD, Kirschtel D, Pollingher U, Zohary T (1999) Biovolume calculation for pelagic and benthic microalgae. J Phycol 35:403–424
- Huntley ME, Lopez MDG, Karl DM (1991) Top predators in the Southern Ocean: a major leak in the biological carbon pump. Science 253:64–66
- Le Quéré C, Rödenbeck C, Buitenhuis ET, Conway TJ and others (2007) Saturation of the Southern Ocean CO_2 sink due to recent climate change. Science 316: 1735–1738
- Le Quéré C, Takahashi T, Buitenhuis ET, Rödenbeck C, Sutherland SC (2010) Impact of climate change and variability on the global oceanic sink of CO₂. Global Biogeochem Cycles 24:GB4007 doi:10.1029/2009GB003599
- Legendre L, Rassoulzadegan F (1995) Plankton and nutrient dynamics in marine waters. Ophelia 41:153–172
- Legendre L, Rassoulzadegan F (1996) Food-web mediated export of biogenic carbon in oceans: hydrodynamical control. Mar Ecol Prog Ser 145:179–193
- Loeb V, Siegel V, Holm-Hansen O, Hewitt R, Fraserk W, Trivelpiecek W, Trivelpiecek S (1997) Effects of sea-ice extent and krill or salp dominance on the Antarctic food web. Nature 387:897–900
- Marshall GJ, Lagun V, Lachlan-Cope TA (2002) Changes in Antarctic Peninsula tropospheric temperatures from 1956 to 1999: a synthesis of observations and reanalysis data. Int J Climatol 22:291–310
- Martinson DG, Stammerjohn SE, Iannuzzi RA, Smith RC, Vernet M (2008) Western Antarctic Peninsula physical oceanography and spatio-temporal variability. Deep-Sea Res II 55:1964–1987
- Menden-Deuer S, Lessard EJ (2000) Carbon to volume relationships for dinoflagellates, diatoms, and other protist plankton. Limnol Oceanogr 45:569–579
- Meredith MP, King JC (2005) Rapid climate change in the ocean west of the Antarctic Peninsula during the second half of the 20th century. Geophys Res Lett 32:L19604 doi:10.1029/2005GL024042
- Metzl N (2009) Decadal increase of oceanic carbon dioxide in Southern Indian Ocean surface waters (1991–2007). Deep-Sea Res II 56:607–619
- Metzl N, Poisson A, Louanchi F, Brunet C, Schauer B, Bres B (1995) Spatio-temporal distributions of air-sea fluxes of CO_2 in the Indian and Antarctic oceans: a first step. Tellus Ser B 47:56–69
- Mitchell BG, Holm-Hansen O (1991) Bio-optical properties of Antarctic Peninsula waters: differentiation from temperate ocean models. Deep-Sea Res I 38:981–1007
- Moline MA, Claustre H, Frazer TK, Schofield O, Vernet M (2004) Alteration of the food web along the Antarctic

Peninsula in response to a regional warming trend. Glob Change Biol 10:1973–1980

- Montagnes DJS, Berges JA, Harrison PJ, Taylor FJR (1994) Estimating carbon, nitrogen, protein, and chlorophyll a from volume in marine phytoplankton. Limnol Oceanogr 39:1044–1060
- Montes-Hugo MA, Vernet M, Martinson D, Smith R, Iannuzzi R (2008) Variability on phytoplankton size structure in the western Antarctic Peninsula (1997–2006). Deep-Sea Res II 55:2106–2117
- Montes-Hugo M, Doney SC, Ducklow HW, Fraser W, Martinson D, Stammerjohn SE, Schofield O (2009) Recent changes in phytoplankton communities associated with rapid regional climate change along the western Antarctic Peninsula. Science 323:1470–1473
- Montes-Hugo M, Sweeney C, Doney SC, Ducklow H and others (2010) Seasonal forcing of summer dissolved inorganic carbon and chlorophyll *a* on the western shelf of the Antarctic Peninsula. J Geophys Res 115:C03024
- Morán XAG, Estrada M, Gasol JM, Pedrós-Alió C (2002) Dissolved primary production and the strength of phytoplankton-bacterioplankton coupling in contrasting marine regions. Microb Ecol 44:217–223
- Olguín HF, Alder AV (2011) Species composition and biogeography of diatoms in antarctic and subantarctic (Argentine shelf) waters (37–76°S). Deep Sea Res II 58: 139–152
- Orr A, Cresswell D, Marshall GJ, Hunt JCR, Sommeria J, Wang CG, Light M (2004) A 'low-level' explanation for the recent large warming trend over the western Antarctic Peninsula involving blocked winds and changes in zonal circulation. Geophys Res Lett 31:L06204
- Pearce I, Davidson AT, Bell EM, Wright S (2007) Seasonal changes in the concentration and metabolic activity of bacteria and viruses at an Antarctic coastal site. Aquat Microb Ecol 47:11–23
- Peylin P, Baker D, Sarmiento J, Ciais P, Bousquet P (2002) Influence of transport uncertainty on annual mean and seasonal inversions of atmospheric CO_2 data. J Geophys Res 107:4385 doi:10.1029/2001JD000857
- Quay PD, Peacock C, Bjrkman K, Karl DM (2010) Measuring primary production rates in the ocean: enigmatic results between incubation and non-incubation methods at Station ALOHA. Global Biogeochem Cycles 24:GB3014 doi:10.1029/2009GB003665
- Schloss IR, Ferreyra GA, Ruiz-Pino D (2002) Phytoplankton biomass in Antarctic shelf zones: a conceptual model based on Potter Cove, King George Island. J Mar Syst 36: 129–143
- Schloss IR, Gustavo AF, Ferrario ME, Almandoz GO and others (2007) Role of plankton communities in sea-air differences in pCO_2 in the SW Atlantic Ocean. Mar Ecol Prog Ser 332:93–106

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- Schofield O, Ducklow HW, Martinson DG, Meredith MP, Moline MA, Fraser WR (2010) How do polar marine ecosystems respond to rapid climate change? Science 328:1520–1523
- Serret P, Fernández E, Anadón R, Varela M (2001) Trophic control of biogenic carbon export in Bransfield and Gerlache Straits, Antarctica. J Plankton Res 23:1345–1360
- Smith RC, Baker KS, Vernet M (1998) Seasonal and interannual variability of phytoplankton biomass west of the Antarctic Peninsula. J Mar Syst 17:229–243
- Smith RC, Martinson DG, Stammerjohn SE, Iannuzzi RA, Ireson K (2008) Bellingshausen and western Antarctic Peninsula region: pigment biomass and sea-ice spatial/ temporal distributions and interannual variability. Deep-Sea Res II 55:1949–1963
- Stammerjohn SE, Martinson DG, Smith RC, Iannuzzi RA (2008) Sea ice in the western Antarctic Peninsula region: spatio-temporal variability from ecological and climate change perspectives. Deep-Sea Res II 55:2041–2058
- Strickland JDH, Parsons TR (1972) A practical handbook of sea-water analysis. Bull 167. Fisheries Research Board of Canada, Ottawa, ON
- Takahashi T, Sutherland SC, Sweeney C, Poisson A and others (2002) Global sea-air CO_2 flux based on climatological surface ocean pCO_2 , and seasonal biological and temperature effects. Deep-Sea Res II 49:1601–1622
- Takahashi T, Sutherland SC, Wanninkhof R, Sweeney C and others (2009) Climatological mean and decadal change in surface ocean pCO₂, and net sea–air CO₂ flux over the global oceans. Deep-Sea Res II 56:554–577
- Tréguer P, Jacques G (1992) Dynamics of nutrient and phytoplankton and cycles of carbon, nitrogen and silicon in the Southern Ocean: a review. Polar Biol 12:149–162
- Turner J, Colwell SR, Marshall GJ, Lachlan-Cope TA and others (2005) Antarctic climate change during the last 50 years. Int J Climatol 25:279–294
- Vaughan DV, Marshall GJ, Connolley WM, Parkinson C and others (2003) Recent rapid regional climate warming on the Antarctic Peninsula. Clim Change 60:243–274
- Wang X, Yang GP, López D, Ferreyra G, Lemarchand K, Xie H (2009) Late austral autumn to spring evolutions of water-column dissolved inorganic and organic carbon in the Scholaert Channel, West Antarctic. Antarct Sci 22: 145–156
- Whitehouse MJ, Meredith MP, Rothery P, Atkinson A, Ward P, Korb RE (2008) Rapid warming of the ocean around South Georgia, Southern Ocean, during the 20th century: forcings, characteristics and implications for lower trophic levels. Deep-Sea Res I 55:1218–1228
- Yager PL, Wallace DWR, Johnson KM, Smith WO Jr, Minnett PJ, Deming JW (1995) The Northeast Water Polynya as an atmospheric CO₂ sink: a seasonal rectification hypothesis. J Geophys Res 100:4389–4398

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