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To cite this version:

L. Farías, L. Florez-Leiva, V. Besoain, G. Sarthou, C. Fernández. Dissolved greenhouse gases (nitrous oxide and methane) associated with the naturally iron-fertilized Kerguelen region (KEOPS 2 cruise) in the Southern Ocean. Biogeosciences, European Geosciences Union, 2015, 12 (6), pp.1925-1940. <10.5194/bg-12-1925-2015>. <hal-01207331>

HAL Id: hal-01207331
http://hal.upmc.fr/hal-01207331
Submitted on 30 Sep 2015

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Dissolved greenhouse gases (nitrous oxide and methane) associated with the naturally iron-fertilized Kerguelen region (KEOPS 2 cruise) in the Southern Ocean

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Received: 23 June 2014 – Published in Biogeosciences Discuss.: 21 August 2014
Revised: 9 February 2015 – Accepted: 23 February 2015 – Published: 24 March 2015

Abstract. The concentrations of greenhouse gases (GHGs), such as nitrous oxide (N2O) and methane (CH4), were measured in the Kerguelen Plateau region (KPR). The KPR is affected by an annual microalgal bloom caused by natural iron fertilization, and this may stimulate the microbes involved in GHG cycling. This study was carried out during the KEOPS 2 cruise during the austral spring of 2011. Oceanographic variables, including N2O and CH4, were sampled (from the surface to 500 m depth) in two transects along and across the KRP, the north–south (TNS) transect (46°–51°S, ∼72°E) and the east–west (TEW) transect (66°–75°E, ∼48.3°S), both associated with the presence of a plateau, polar front (PF) and other mesoscale features. The TEW presented N2O levels ranging from equilibrium (105 %) to slightly supersaturated (120 %) with respect to the atmosphere, whereas CH4 levels fluctuated dramatically, being highly supersaturated (120–970 %) in areas close to the coastal waters of the Kerguelen Islands and in the PF. The TNS showed a more homogeneous distribution for both gases, with N2O and CH4 levels ranging from 88 to 171 % and 45 to 666 % saturation, respectively. Surface CH4 peaked at southeastern stations of the KPR (A3 stations), where a phytoplankton bloom was observed. Both gases responded significantly, but in contrasting ways (CH4 accumulation and N2O depletion), to the patchy distribution of chlorophyll a. This seems to be associated to the supply of iron from various sources. Air–sea fluxes for N2O (from −10.5 to 8.65, mean 1.25 ± 4.04 µmol m−2 d−1) and for CH4 (from 0.32 to 38.1, mean 10.01 ± 9.97 µmol−2 d−1) indicated that the KPR is both a sink and a source for N2O, as well as a considerable and variable source of CH4. This appears to be associated with biological factors, as well as the transport of water masses enriched with Fe and CH4 from the coastal area of the Kerguelen Islands. These previously unreported results for the Southern Ocean suggest an intense microbial CH4 production in the study area.

1 Introduction

The increasing concentration of greenhouse gases (GHGs) in the troposphere, such as CO2, N2O and CH4, affect the Earth’s radiative balance. Additionally, the increasing concentration of ozone-depleting gases (such as chlorofluorocarbons and N2O) in the stratosphere is weakening the ozone shield, permitting higher levels of damaging ultraviolet radiation to reach the Earth’s surface. The relative potency of a
GHG is determined by its respective residence time in the atmosphere (Cicerone and Orelamland, 1988), and the extent of magnitude of emissions to the atmosphere, of which the input/contribution from the ocean is considered to be important (IPCC, 2013).

Although oceans are generally considered to be a net source of GHGs, for example N$_2$O and CH$_4$, to the atmosphere, the oceanic distribution of these GHGs and the amount exchanged via the air–sea interface is highly variable (Nevison et al., 1995; Holmes et al., 2000; Rhee et al., 2009). Thus, source and sink behaviours of GHGs have been observed on different spatial and temporal scales. In general terms, these behaviours depend on biological and physical processes that promote outgassing or sequestering mechanisms. Physical and biological features of the Southern Ocean suggest the existence of a potential for both the production and removal of CH$_4$ and N$_2$O (Rees et al., 1999; Tilbrook and Karl, 1994), although very little information on dissolved N$_2$O and CH$_4$ distributions is currently available for the region. The substantial spatial variation in regional gas exchange could be due to the increased gas solubility in low-temperature Subantarctic waters, combined with either the downwelling associated with intermediate and deep water formation in the northern and southern part, respectively, of the Antarctic Zone. Further variation may be caused by the upwelling of deep and intermediate waters in the southern part of the Antarctic Polar Frontal Zone (Park and Vivier, 2012).

In different regions of the Southern Ocean, the surface layer is permanently supersaturated with CH$_4$ (Bates et al., 1996; Tilbrook and Karl, 1994; Yoshida et al., 2011), but this is not the case for N$_2$O, which may occur in either under- or supersaturated conditions (Law and Ling, 2001; Rees et al., 1997; Chen et al., 2014). Aside from physical processes affecting GHG concentrations in the water column and their concomitant air–sea fluxes, the concentrations also depend on organic matter (OM) availability and oxygen levels, which determine whether aerobic or anaerobic respiration occurs (Codispoti et al., 2001; Reeburgh, 2007). The availability of dissolved iron (dFe) should also be included as a direct variable affecting the recycling of GHGs (through production and consumption), as several Fe-containing enzymes are required for GHG cycling, which are involved in the electron transfer chains in bacterial respiratory systems and nitrogen cycling (Arrieta et al., 2004; Kirchman et al., 2003; Morel and Price, 2003). Indirectly, dFe stimulates ocean productivity, which can enhance carbon and nitrogen export from the euphotic zone to the subsurface (Boyd and Ellwood, 2010), resulting in an increase in microbial activities and mediate GHG production via nitrification (Fuhrman and Capone, 1991).

N$_2$O is mainly produced during the first step of nitrification (the aerobic oxidation of NH$_4^+$ to NO$_2^-$), as well as during partial denitrification (the anaerobic reduction of NO$_3^-$/NO$_2^-$ to N$_2$O). N$_2$O can also be consumed by complete denitrification via dissimilatory reduction to N$_2$ (Codispoti and Cristiensen, 1985) or assimilatory N$_2$O reduction to NH$_4^+$ (Farias et al., 2013). CH$_4$, on the other hand, is mainly formed by methanogens during anaerobic OM degradation (Wuebbles and Hayhoe, 2002) or by methylotrophs during CH$_4$ formation derived from transformations of methyl compounds such as methylphosphonate (MPn; Karl et al., 2008), dimethylsuloniopropionate (DMSP; Damm et al., 2010) and dimethylsulfide (DMS; Florez-Leiva et al., 2013). In addition, CH$_4$ can be consumed (oxidized) via aerobic methanotrophy (Hanson and Hanson, 1996). Since Southern Ocean waters are well oxygenated, the dominant mechanisms of production of the GHGs in surface waters are nitrification in the case of N$_2$O (Nevison et al., 2003) and methanogenesis in suspended particles (Scranton and Brewer, 1977) and/or methylotrophy (Sun et al., 2011) for the case of CH$_4$.

The main objective of the present study is to describe for the first time the N$_2$O and CH$_4$ contents in the Southern Ocean under the influence of natural fertilization during the spring phytoplankton bloom in the Kerguelen Plateau region (KPR). The determination of the role of the Southern Ocean in CH$_4$ and N$_2$O air–sea exchange may be critical in understanding the factors that influence GHG cycling. This includes dFe which comes from different sources within the KPR, inducing mesotrophic conditions associated with the coastal waters of the Kerguelen Islands and of the central Kerguelen Plateau, an area within the central plateau area of the KPR that demonstrates an annually recurrent phytoplanktonic bloom in the (Blain et al., 2008; Chever et al., 2010), and with the Antarctic PF and other mesoscale structures (Mongin et al., 2008; Lasbleiz et al., 2014).

2 Methods

2.1 Study area

Samples were collected within the KPR (Fig. 1) during the KEOPS 2 cruise at stations along the north–south transect (TNS; 46°–51° S) and the east–west transect (TEW; 66°–75° E). The cruise took place from 11 October to 11 December 2011 onboard the research vessel (RV) Marion Dufresne. Some of the stations were located in the naturally Fe-enriched polar front (PF) zone, in the coastal shell waters of the Kerguelen Islands and within the southeastern KPR bloom (including the A3 Stations from the previous KEOPS 1 cruise). Station R-2, located east of the Kerguelen Plateau, was considered to be typical of high-nutrient, low-chlorophyll (HNLC) conditions (see Table 1). The positions of the stations were selected according to a strategy based on real-time ocean colour and altimetry satellite images (d’Ovidio et al., 2015).
2.2 Sampling

Continuous vertical profiles of temperature, salinity, dissolved O₂, fluorescence and photosynthetically active radiation (PAR) were obtained using a conductivity—temperature—depth (CTD) sensor. Seawater samples were collected using a Sea-Bird SBE 911plus CTD unit mounted on a 24-bottle (12 L each) rosette. Water samples for gases (N₂O, CH₄), nutrients and pigments (sampled in this consecutive order) were obtained from nine depths distributed between the surface and 500 m depth. Water samples for CH₄ (triplicate) and N₂O (triplicate) analyses were taken in 20 mL glass vials and poisoned with HgCl₂ (0.1 mL of saturated HgCl₂ solution per vial). Subsequently, the vials were sealed with a butyl-rubber septum and an aluminium cap, avoiding bubble formation, and stored in darkness at room temperature until laboratory analysis. Syringes of 50 mL were directly connected to the spigot of the Niskin bottles to take nutrient samples (NO₃, PO₄, SiO₂) at each sampled depth. Duplicate samples were collected and drawn through a 0.45 µm Up-tidisc adapted to the syringe, and then immediately analysed using an autoanalyser (more details in Blain et al., 2015). Total chlorophyll a (Chl a) samples in triplicate were filtered into a 25 mm glass-fibre filter (GF/F), and then immediately frozen (−20°C). Samples were kept until later analysis by high-performance liquid chromatography (HPLC) (more details in Lasbleiz et al., 2014).

Figure 1. Map showing the location of biogeochemical stations sampled during the KEOPS 2 cruise. Bathymetric topography is shown in the main oceanographic region. The orange line delimits the position of the polar front. The transects are indicated.
2.3 Chemical analysis

N$_2$O and CH$_4$ were analysed by generating a 5 mL ultra-pure helium headspace in a vial using a gastight syringe and then equilibrating the gas and liquid phases at 40°C within the vial. Following this the gases within the vial were quantified through the use of a gas chromatograph determined by headspace equilibration (5 mL helium headspace and 15 mL of seawater) at 40°C. Finally quantification via chromatography was carried out. N$_2$O was analysed in a Varian 3380 gas chromatograph using an electron capture detector (ECD) at 350°C and connected to an autosampler device. CH$_4$ was analysed in a Shimadzu 17A gas chromatograph using a flame ionization detector (FID) at 250°C through a capillary column GS-Q at an oven temperature of 30°C. A calibration curve was made with four concentrations for N$_2$O (0.1, 0.32, 0.5 and 1 ppm, by Matheson standards) and four concentrations for CH$_4$ (0.5, 1.78, 2 and 10 ppm, by Matheson standards). Both detectors linearly responded to these concentration ranges. The analytical error for the N$_2$O and CH$_4$ analyses was less than 3 and 5%, respectively. The ECD and FID linearly responded to these concentration ranges, and the analytical error for the N$_2$O measurements for this study was about 3%. The uncertainty of the measurements was calculated from the standard deviation of the triplicate measurements by depth. Samples with a variation coefficient higher than 10% were not taken into account for the gas database.

More details regarding the analysis of both gases can be found in Farias et al. (2009). Nutrients were immediately analysed onboard using standard automated colorimetric methods (Tréguer and LeCorre, 1975) using the continuous flow autoanalysers (Skalar). The precision and detection limit of the method was ±30 and 110 nM for PO$_4$$^{3-}$, ±2 and 10 ppm, by Matheson standards). Both detectors linearly responded to these concentration ranges. The analytical error for the N$_2$O and CH$_4$ analyses was less than 3 and 5%, respectively. The ECD and FID linearly responded to these concentration ranges, and the analytical error for the N$_2$O measurements for this study was about 3%. The uncertainty of the measurements was calculated from the standard deviation of the triplicate measurements by depth. Samples with a variation coefficient higher than 10% were not taken into account for the gas database.

2.4 Data analysis

To interpret the vertical variation of N$_2$O and CH$_4$, and assess how biogeochemical processes may affect their concentrations, the water column was divided into two layers according to density gradient: (1) well mixed and (2) sub-surface from the base of the mixed layer (ML) to 500 m (arbitrary depth used only for comparison proposes). Nutrient inventories for TChl $a$, N$_2$O and CH$_4$ were calculated by numerical integration of data with linear interpolation at intervals of 1 m, based on at least 4–6 sampled depths per layer. Saturation percentages of gases were calculated from the measured CH$_4$ and N$_2$O concentrations and those estimated to be in equilibrium with the current gas concentrations in the atmosphere register (NOAA/ESRL programme http://www.esrl.noaa.gov/gmd/hats/combined/N2O.html) based on in situ temperature and salinity records according to the solubility parameterization of CH$_4$ (Wiesen-burg and Guinasso, 1979) and N$_2$O (Weiss and Price, 1980). GHG flux through the air–sea interface was determined using the following equation, modified by Wanninkhof (1992):

$$F = kw(T^2, \text{salinity}) \cdot (C_w - C_a),$$

where kw is the transfer velocity from the ML to the atmosphere, as a function of wind speed, temperature and salinity in the ML according to parameterization; $C_w$ is the mean gas concentration in the mixed layer; and $C_a$ is the gas concentration in the mixed layer expected to be in equilibrium with the atmosphere. Since gas transfer velocity is related to wind speed, this was calculated according to the well-known exchange models of Liss and Merlivat (1986) (hereafter referred to as LM86) and Wanninkhof (1992) (hereafter referred to as W92), based on the dependence of the transfer velocity on wind speed. Wind speed and direction were obtained from the ship’s meteorological station. Wind speed was estimated as a moving 7-day average prior to the sampling period in order to smooth out short-term fluctuations and highlight longer-term trends. The mixed layer depth was calculated using a potential density-based criterion, defining the mixed layer depth (ML) as the shallowest depth at which density increased by 0.02 kg m$^{-3}$ from the sea surface value. Pearson product-moment correlations (rs) were determined for GHGs, and TChl $a$ and nutrient inventories were estimated for both the ML and for the whole water column from the surface to 500 m depth. The threshold value for statistical significance was set to $p < 0.05$. A principal component analysis (PCA) using the empirical orthogonal function (EOF; Emery and Thomson, 1997) was performed to find the co-variability patterns of a number of stations located in spatial gradients in terms of nutrients, gases (O$_2$, N$_2$O, and CH$_4$), TChl $a$ and $d$Fe. This analysis excluded the stations from the TNS as no measurements were recorded (Quéroué et al., 2015). PCA was done with all biogeochemical variables measured in the ML and with those variables obtained in the water column from the surface to a depth of 500 m in order to detect differences in the vertical structure.

3 Results

3.1 Oceanographic conditions

Oceanographic characteristics of the sample stations during the KEOPS 2 cruise are shown in Table 1. Two transects, carried out almost synoptically across and along the KPR (survey region – Fig. 1), were undertaken to establish the position of the main mesoscale structures as fronts (Fig. 2). The polar front (PF) crosses the KPR and denotes certain physical structures (i.e. convergence processes) visible throughout temperature and salinity (Park and Vivier, 2012).

Regarding the TEW (66 to 75°E, along 47°S), vertical cross sections of temperature and salinity along with a temperature–salinity ($T$–$S$) diagram are illustrated in Fig. 2a,
Temperature and salinity varied between 2.41 and 3.30 °C and between 33.60 and 34.67, respectively. A weak structure with colder and fresher surface waters was registered in the PF, which crossed these transects twice, at \( \sim 71^\circ \text{E} \) (TEW-3, TEW-4) and at \( \sim 73.5^\circ \text{E} \) (TEW-7, TEW-8). Middle stations (TEW-4, TEW-5 and E) are located in an area with a complex recirculatory system. This is a superficial section inundated by mixed Antarctic Surface Water (AASW) and coinciding with an area of a PF northward inflexion (Fig. 1). The presence of Subantarctic Mode Water (SAMW) was observed east of 73.5° E (TEW-7, TEW-8; Fig. 2f). In addition, a marked variability in subsurface water was observed, attributed to the mixing of water masses; this was particularly strong in TEW within the PF, revealing a vertical mixing process produced by convergence, particularly evident at TEW-7 (Fig. 2e). Regarding the TNS (46°–51° S, along \( \sim 72^\circ \text{E} \)), Fig. 2b, d and f show vertical cross sections of temperature, salinity and a \( T-S \) diagram, respectively. Temperature and salinity fluctuated from 1.67 to 4.17 °C and from 33.67 to 34.68, respectively, and a gradual decrease in temperature and increase in salinity were observed in the surface layer from north to south (Fig. 2b, d). A water parcel of a relatively cold water mass was observed to be spreading northward in subsurface waters. This is noted as an expression of the PF, which marks the location where the AASW moves northward, descends rapidly, and sinks below 200 m depth (Fig. 2b). These distributions coincided with the expected water mass distribution, this being the case for the northern (TNS-1, TNS-2) and southern (A3, TNS-10) stations, mainly occupied from the surface to 250 m by the SAMW and the AASW, respectively (Fig. 2f).

### 3.2 Biogeochemical variables

Figure 3 shows vertical cross sections along the TEW of biogeochemical variables including nutrients (only NO_3\(^-\) and PO_4^{3-} \), TChl \( a \), O\(_2\) and GHGs. The surface layer continuously showed NO_3\(^-\) concentrations, fluctuating from 22 to 27 \( \mu \text{mol} \text{L}^{-1} \) (typical condition of the AASW). However, a relative depletion of NO_3\(^-\) was observed at the stations lo-
Vertical cross section of (a) nitrate (µmol L\(^{-1}\)), (b) phosphate (µmol L\(^{-1}\)); (c) chlorophyll \(a\) (µg L\(^{-1}\)), (d) dissolved oxygen (µmol L\(^{-1}\)), (e) nitrous oxide (nmol L\(^{-1}\)) and (f) methane (nmol L\(^{-1}\)) for zonal transect between 69 and 75°E. Arrows indicate the position of the PF crossing this transect.

cated to the north and east of the PF (Fig. 3a). \(PO_4^{3-}\) presented the same pattern as \(NO_3^-\), and the N:P ratio of dissolved nutrients averaged around 14.5, with the exception of some values of 13.2 from stations located close to the PF (Fig. 3b). \(TChl \ a\) fluctuated from 0.005 to 4.69 µg L\(^{-1}\) and peaked at TEW-1 and TEW-2 (both located in a coastal area 10 and 75 km away from Hillsborough Bay coast) as well as TEW-7 and TEW-8 (to the north of the PF). \(TChl \ a\) showed a relative decrease at stations located in the central section (Fig. 3c). \(O_2\) concentration varied from 320 µM (in surface water) to 185 µM (at 500 m depth), consistently maintaining supersaturation conditions (Fig. 3d).

\(N_2O\) fluctuated from 14.0 to 25.4 nmol L\(^{-1}\) (equivalent to a range of 102–182.2 % saturation, Fig. 3e). Superficially, in the western and central section (70.5–73° E), the \(N_2O\) concentration was close to equilibrium with the atmosphere in surface waters, and in surface waters at sites where the PF crosses the transect, i.e. TEW-4 and TEW-7 (Fig. 3e) \(N_2O\) was slightly undersaturated (around 90 %). \(N_2O\) levels increased slightly attaining around 120 % saturation towards the subsurface water. \(CH_4\) ranged from 1.4 to 31.35 nmol L\(^{-1}\), equivalent to a saturation range of 43–969 %. In contrast to \(N_2O\), surface waters were always supersaturated in \(CH_4\), demonstrating the greatest increase in gas levels (up to 970 %) in coastal waters close to the Kerguelen Islands, a relative decrease (< 200 % saturation) in the central section (between 71° and 73.5° S, or TEW-4, TEW-5 and E-2), and a strong increase (up to 778 %) at TEW-7. Remarkably, \(CH_4\) concentrations in subsurface waters were low compared to the surface waters (Fig. 3f).

Vertical cross sections of biogeochemical variables along the TNS are shown in Fig. 4. \(NO_3^-\) and \(PO_4^{3-}\) gradually increased from north to south from 24 to 30 µmol L\(^{-1}\) and from 1.5 to 2 µmol L\(^{-1}\), respectively (Fig. 4a, b). This spatial trend coincided with the expected transition of water mass dominance and its mixing between the SAMW and the AASW (Fig. 2f). \(TChl \ a\) ranged from 0.005 to 2.391 µg L\(^{-1}\) and peaked in the southernmost stations (TNS-8, TNS-9 and A3-2; Fig. 4b) and coincided with a slight increase in nutrients. There a deep Fe-enriched and lithogenic silica reservoir seemed to influence the area (Lasbleiz et al., 2014; Quéroué et al., 2015). \(O_2\) distribution was similar to that observed in the TET.

\(N_2O\) concentrations ranged from 12.37 to 23.8 nmol L\(^{-1}\), equivalent to 88.5 to 171 % saturation. \(N_2O\) levels close to equilibrium or undersaturation were often observed in surface waters, except at TNS-08 (Fig. 4e). \(CH_4\) varied from 1.47 to 21.88 nM, or 45 to 666 % saturation, and peaked in southern stations (Fig. 4f). Notably, high levels of \(TChl \ a\) were associated with high concentrations of \(CH_4\) in this tran-
sect. Southern stations had extremely low N$_2$O concentrations (less than 6.9 nM or 70 % saturation), such as A3-1, which is located in an area of relatively high dFe availability and within a phytoplankton bloom.

PCA was performed including dFe and GHG data obtained from the TEW, as shown in Fig. 5. The results did not change when O$_2$ was removed from the analysis, indicating that O$_2$ availability does not modify the percentage of the variance. When the data set used in the PCA is restricted to the ML (Fig. 5a), stations located on the TEW were grouped into three sets, clearly separating stations located in eastern (north of the PF, TEW-7), western, and coastal areas (TEW-1, TEW-2), and within the central section (TEW-4, TEW-5, E2). The variability among stations can be predominantly explained by the first component, accounting for 75.7 % of the variance. Figure 5 suggests possible interpretations of the relationships between the variables with their respective weights assigned to each of them (illustrated with an eigenvector). The figure shows a close relationship between N$_2$O, nutrients, CH$_4$, dFe and TChl a. PCA using data from the entire water column (Fig. 5b) reproduced a similar grouping to those performed with data from surface waters (Fig. 5a).

3.3 Vertical distribution of gases and other variables at selected stations

Figure 6 shows typical profiles of oceanographic and biogeochemical variables (including gases). Stations were separated a priori according to biogeochemical (PCA for the case of the TEW; Fig. 5) and oceanographic criteria (T–S diagram, Fig. 2e, f). Selected stations included those of A3, with a recurrent annual bloom (historical station sampled in KEOPS 1; Blain et al., 2007) and moderate dFe levels (∼0.18 nmol L$^{-1}$). Some stations displayed evidence of an active uptake of dFe; TEW-7 had one of the highest dFe (∼0.40 nmol L$^{-1}$) and TChl a levels, and also showed evidence of rapid dFe uptake (Fourquez et al., 2014). For comparative purposes, we include the most northern station of the TNS (TNS-01), R-2 (in the HNLC area), and a coastal station close to the Kerguelen Islands, which had the highest dFe levels (up to 3.82 nmol L$^{-1}$). Vertical distribution of N$_2$O and CH$_4$ clearly varied, while elevated CH$_4$ concentrations were generally observed superficially and in the ML base, and concentrations decreased with increased depth, whereas N$_2$O concentrations gradually increased with depth. Gas contents also differed between stations and were observed to correlate in a similar way with TChl a and dFe levels.

The stations located at the extremities of the TEW (i.e. TEW-1 and TEW-7) had the highest CH$_4$ levels (Fig. 6),
while N$_2$O levels were relatively low. Conversely, TNS-1 and A3-2, located respectively in the extreme north and south of the TNS, presented relatively low levels of CH$_4$ compared to the TEW. Station R-2, which is located in the HNLC area, had the lowest N$_2$O and CH$_4$ content, and both gases were homogeneously distributed with depth (Fig. 6). This is consistent with TChl $a$ levels of less than 0.5 µg L$^{-1}$.

### 3.4 Nutrient, TChl $a$, dFe, and GHG inventories and air–sea GHG exchanges

Table 2 shows the inventories of NO$_3^-$, PO$_4^{3-}$ and GHGs in the ML and the water column from the surface to 500 m; mean GHG concentrations in the ML, wind speed, and air–sea GHG fluxes are also included. ML depths varied widely from 16 m (at the station near the Kerguelen Islands) to 181 m. The TChl $a$ pool, estimated on the basis of the photic layer, fluctuated from 8.77 to 75.45 mg m$^{-2}$. Levels were notably elevated at A3-2 (up to 5-fold greater), in comparison to the more oligotrophic stations such as R-2. Surface NO$_3^-$ and PO$_4^{3-}$ inventories did not show significant differences among stations and varied between 1.56 and 16.03 and between 0.13 and 1.07 mol m$^{-2}$, respectively. Minimal values were registered at stations TEW-7 and TNS-1, both located north of the PF.

N$_2$O pools varied from 0.201 to 2.55 mmol m$^{-2}$ and from 1.12 to 10.05 in the ML and the whole water column, respectively. Minimum values were registered in the ML at stations within the PF and also to the north. These surface pools did not significantly correlate with TChl $a$, but correlated strongly and negatively with nutrients (rs: 0.91 $p < 0.001$ for NO$_3^-$; rs: 0.92, $p < 0.001$ for PO$_4^{3-}$). CH$_4$ inventories fluctuated between 0.19 and 3.31 mmol m$^{-2}$ for the ML, and 1.06 and 7.44 mmol m$^{-2}$ for the whole water columns. Once again, inventories in the ML were 2- and 5-fold higher at TEW-7 and A3-2, respectively, than at R-2. CH$_4$ inventories were 4- and 7-fold higher at TEW-7 and A3-2, respectively, than at R-2. The comparison between the CH$_4$ inventories (standardized by the thickness of the layer) obtained from the ML and from the entire water column indicates that the maximum values came from the ML’s base, notably in the PF (Table 2). CH$_4$ pools correlated positively with TChl $a$ pools (rs = 0.69; $p < 0.05$) but did not show any correlation with NO$_3^-$ and PO$_4^{3-}$. Thus, minimum values for both nutrients were found when TChl $a$ was higher.

Average hourly wind velocity during the cruise was 10.53 ± 5.52 m s$^{-1}$, occasionally falling below 0.31 m s$^{-1}$ or rising above 29.1 m s$^{-1}$. The ML depth did not show any significant relationship to wind speed (rs: 0.20 $p = 0.41$) or the water mass structure (Table 1 and Fig. 2) but seems to be related to the complex mesoscale circulation observed in the KPR (Park et al., 2014; Zhou et al., 2014). N$_2$O fluxes, estimated by LM86, fluctuated between ~9.69 and 10.02 µmol m$^{-2}$ d$^{-1}$ (mean: 1.25 ± 4.04 µmol m$^{-2}$ d$^{-1}$), while those estimated by W92 varied from ~18.69 to 20.2 µmol m$^{-2}$ d$^{-1}$ (mean: 2.41 ± 7.88). Substantial differences were observed between the cubic (LM86) and the quadratic parameterizations (W92) during periods of high wind speeds, such as those measured during the TNS (21–23 October 2011, mean value of 12.08 m s$^{-1}$), compared to those registered during the TEW (31 October–2 November, mean value of 5.61 m s$^{-1}$). The W92 parameterization
Figure 6. Vertical distribution of biogeochemical variables from selected stations. Different biogeochemical regimes are defined as HNLC area (R), northern and southern area of the polar front (TNS-1 and A3-2) and close to the polar front (TEW-3 and TEW-7).
showed an increase in calculated fluxes by approximately a factor of 2 at high wind speeds, while at low wind speeds the difference between LM86 and W92 was up to a factor of ~1.6 higher (see Table 2). CH$_4$ fluxes varied from 0.21 to 38.1 µmol m$^{-2}$ d$^{-1}$ (mean: 10.01 ± 9.97) and from 0.32 to 70.24 µmol m$^{-2}$ d$^{-1}$ (mean: 21.27 ± 21.07) when LM86 and W92 were employed, respectively. At times the study area acted as a source of very high CH$_4$ effluxes into the atmosphere, particularly at stations TNS-9 and A3-2, where emissions were around 3 times higher compared to those calculated for R-2. There are important differences between the two parameterizations, although the same trend was obtained among stations (Table 2).

### 4 Discussion

Iron fertilization in the KPR influences phytoplankton growth and primary production (PP) and other microbial activities (Cavagna et al., 2014; Christaki et al., 2014), as well as relative CH$_4$ accumulation (Fig. 3f and 4f) and some N$_2$O depletion (Fig. 3e). The gas distribution pattern clearly matched those patterns in Quéréau et al. (2014) for TChl a and PCA grouped stations using dFe. The segregation of stations includes the coastal area (TEW-1, TEW-2), the PF (TEW-7) and the central plateau region (A3-2). In the case of KEOPS 2, phytoplanktonic blooms were mainly represented by a microplanktonic community (Lasbeiz et al., 2014), as those observed in the north polar front (TEW-7) and the central part of the KPR (A3) stations displayed high rates of iron uptake (Quéréau et al., 2014). These previously indicated areas demonstrated variable but high particulate Fe of biogenic origin (Van der Merwe et al., 2015). This confirms an increased biological uptake, which in turn determines a rapid dFe turnover. The observed gas distribution patterns raise questions as to how the complex circulation and some mesoscale structures support relatively high TChl a accumulation and microbial activities in comparison to surrounding waters, and particularly whether there are some fertilization mechanisms (including the addition of Fe and nutrients) that promote GHG cycling and its associated microbial activities.

#### 4.1 N$_2$O cycling

Fuhrman and Capone (1991) pointed out that the stimulation of ocean productivity through the addition of Fe enhances nitrogen export from the euphotic zone to the subsurface layer and may result in enhanced N$_2$O formation via the stimulation of nitrification. This stimulation may occur through the activation of metallo-proteins that are involved in the various stages of ammonium and nitrite oxidation, as ammonia-oxidizing nitrifiers oxidize NH$_4^+$ and NH$_2$OH to NO$_2^-$ using iron-containing ammonia monoxygenase (AMO) and hydroxylamine oxidoreductase (HAO), respectively (Morel et al., 2003). Since N$_2$O is a powerful greenhouse gas, 300 times more radiative than CO$_2$ per molecule, Fe addition could counteract the climatic benefits of atmospheric CO$_2$ drawdown (Jain et al., 2000). The link between Fe fertilization and enhanced N$_2$O formation via nitrification was supported by Law and Ling (2001), who found a small but significant N$_2$O accumulation in the pycnocline during the Southern Ocean Iron Enrichment Experiment (SOIREE) at
The process is stimulated by enhanced Fe supply. N (Gonzalez et al., 2014), suggesting that N fixation (TEW-7) also had the highest surface N fixation more commonly than originally expected. Coincidently,dez et al., 2011), suggesting that N fixation levels of Fe, high levels of particulate N (within the PF) (with moderate salinity from 34 to 10) would be required to produce the observed undersaturation, neither of which was observed during the sampling (Table 1) or expected during this season (Park et al., 2014). Additionally, if the two water masses were mixed proportionally, as they are, the resulting process cannot produce undersaturation in regard to the original N2O levels and the signature temperature and salinity. Recently, Chen et al. (2014) reported that surface waters of the Indian sector of the Southern Ocean were undersaturated in N2O, suggesting a N2O influx. This phenomenon in the surface water may result from the intrusion of freshwater from ice melt and the northeastward transport of the AASW. However, in the KPR, N2O undersaturation seems to be located in an area of high particle concentration under the influence of the SAMW (northern the PF).

Thus, a preliminary analysis indicates that biological processes are responsible for the N2O undersaturation and the concomitant influx from the atmosphere. In contrast, subsurface waters have higher N2O concentrations (saturations from 120 to 180 %) than surface waters, which indicate a net accumulation. In this case the most likely process responsible for N2O accumulation is aerobic ammonium oxidation (Codispoti et al., 2001), but no significant difference was noted at the stations with the highest TChl a levels, indicating that N2O production by nitrification was not substantially stimulated at those stations.

4.2 CH4 cycling

There have been few studies on CH4 distribution and production in the Southern Ocean (Lamontagne et al., 1973; Tilbrook and Karl, 1994; Heeschen et al., 2004). Surface water in the Southern Ocean has been reported to be undersaturated or
lightly saturated with respect to atmospheric CH$_4$, as a result of the entrainment of CH$_4$-depleted deep water into surface water and the seasonal ice covers acting as a barrier against gas exchange (Yoshida et al., 2011). Regarding the effect of iron addition on CH$_4$ cycling, Wingenter et al. (2004) found low levels of CH$_4$ production (less than 1%) during artificial Southern Ocean Fe enrichment experiments (SOFex). Simulated large-scale Southern Ocean Fe fertilization (OIF) also resulted in anoxic conditions which may favour anaerobic methanogenesis (Oschlies et al., 2010).

However, our results show that surface and subsurface waters are supersaturated in CH$_4$ with a 4-fold enrichment in CH$_4$ with respect to the control area (Fig. 3e); this was associated with areas with elevated TChl $a$ levels and iron uptake by microbial communities (Fourquez et al., 2014). Results showed a marked spatial differences in CH$_4$ content measured in the TEW and TNS (Student’s $t$: 3.21 $p < 0.001$) (Fig. 3f and 4f), and that surface CH$_4$ accumulation generally coincided with areas of relatively higher dFe levels, which in turn favours primary production (PP). Likewise, the CH$_4$ accumulation at pycnoclines (Fig. 6) indicates that most CH$_4$ came from accumulated particles sinking from the surface water, as commonly observed by Holmes et al. (2000) in different marine systems. PCA, which included the measurement of dFe, revealed a close relationship between CH$_4$ accumulation and Fe availability and clearly grouped in areas with different biogeochemical characteristics. The fact that the western and eastern sections showed high Fe levels (Quéroué et al., 2014) relative to the central section of the TEW, and that these sections had high CH$_4$ levels, suggests that Fe stimulates CH$_4$ production. A similar situation occurs at the A3 stations with high TChl $a$ levels and PP rates, as shown by Cavagna et al. (2014). For example, A3-2 and TEW-7 (maximum TChl $a$) had the highest integrated primary production rates (up to 3380 mg m$^{-2}$ d$^{-1}$) and the lowest C export level of around 2–3 % (Cavagna et al., 2014); this suggests an intense level of PP supported by regenerated N sources. The situation observed at stations TEW-7 and A3-2 contrasted with that observed at R-2 with with the lowest rate of regenerated production (with a PP rate of around 135 mg m$^{-3}$ d$^{-1}$ and an exported C rate of around 25 % of PP).

Two hypotheses exist for CH$_4$ production in surface waters. The first is that production only occurs in association with anoxic particles (Karl and Tilbrook, 1994), produced for the most part by grazing zooplankton, as methanogenic bacteria were considered to be present in an anaerobic microenvironment in organic particles (pellets) or in the guts of zooplankton (Allredge and Cohen, 1987; Karl and Tilbrook, 1994). The other hypothesis formulated more recently is that phytoplankton blooms should favour zooplankton grazing process and/or stimulate bacterioplankton activity as CH$_4$ is generated via the degradation of organic methyl compounds by bacteria (Karl et al., 2008).

Increased grazing of microbes by microzooplankton, as observed by Christaki et al. (2014), may contribute to particle recycling (rich in organic carbon and DMSP) and increase the potential for methanogenesis (Weller et al., 2013). Yoshida et al. (2011) found that high CH$_4$ production in the Southern Ocean probably resulted from the grazing processes of Antarctic krill and/or from zooplankton that feed on phytoplankton and the subsequent microbial methanogenesis. This agrees with the findings for sites enriched with iron and biomass that exhibit high carbon fluxes at 100 m depth, dominated by large faecal pellets rather than phytodetrital aggregates (Laurenceau-Cornet et al., 2015).

Conversely, aerobic CH$_4$ production in the water column could be associated with heterotrophic activities. Christaki et al. (2014) showed that the highest bacterial production rates (up to 110 mg C m$^{-2}$ d$^{-1}$) and the greatest abundance of heterotrophic bacteria were associated with stations where the phytoplankton bloom was developed (TEW-7 and A3-2). Recent evidence indicates that methylotrophs are candidates for mediated CH$_4$ generation using methylated compounds such as DMSP and DMS (Florez-Leiva et al., 2013; Weller et al., 2013). Among these heterotrophic microorganisms DMS degradation can be ascribed to methylotrophic bacteria (Vissher et al., 1994) that derive energy from the conversion of methyl into other products, as well as using S as a source of methionine biosynthesis (Kiene et al., 1999). Current studies of natural and cultivated SAR11 alphaproteobacteria (strain Ca. P. ubique HTCC1062; Sun et al., 2011) indicate that these microorganisms, among the most abundant heterotrophic bacteria in surface waters, possess genes that are encoded for oxidation pathways of a variety of C1 compounds and that they have the capacity for demethylation and C1 oxidation but do not incorporate C1 compounds as biomass. This suggests that there is a close relationship between phytoplankton, the only producers of DMSP (Yoch, 2002), and microbial communities which may be recycling DMS. Phyto- and bacterioplankton relationships control DMS turnover, which could result in several mechanisms of DMSP/DMS degradation (Simó et al., 2002; Viá-Costa et al., 2006) and produce CH$_4$ (Damm et al., 2010; Florez-Leiva et al., 2013; Weller et al., 2013). These publications show that phytoplankton species composition and biomass in different bloom phases, as well as eddy dynamics, were important determinants of CH$_4$ saturation and emission.

With regard to the distribution, vertical profiles of the gas indicate that most CH$_4$ is being formed at the surface and pycnoclines (at the base of the ML) and consumed at subsurface and intermediary depths (Fig. 6). Thus, CH$_4$ distribution appears to be controlled largely by biological mechanisms rather than by mixing, contrary to what has been reported by Heeschen et al. (2004). In general, surface waters of the Southern Ocean were undersaturated with respect to atmospheric CH$_4$ as the result of the entrainment of CH$_4$-depleted deep water to the surface and from seasonal ice cover acting as a barrier for gas exchange. We observed CH$_4$ undersatura-
tion, fluctuating between 40 and 90%, at most sampled stations at depths of > 200 m. It is unlikely that this undersaturation results from the entrainment of CH<sub>4</sub>-depleted waters that have high levels of gas solubility; more likely, it results from a biological consumption, as it is more likely that a biological mechanism is involved. The only known process able to consume CH<sub>4</sub> is methanotrophy, and the fact that subsurface waters were depleted of CH<sub>4</sub> suggests that CH<sub>4</sub> consumption is higher than production or that no production occurs in subsurface waters. Interestingly, although CH<sub>4</sub> microbial oxidation occurs throughout the water column and is recognized as an important process that reduces CH<sub>4</sub> emissions (Reeburgh et al., 2007; Rehder et al., 1999), there have been few investigations on microbial communities mediating aerobic CH<sub>4</sub> oxidation. There do exist a few measurements of aerobic CH<sub>4</sub> oxidation in marine environments, as well as measurements taken from open systems under oligotrophic regimes (Tilbrook and Karl, 1994, Holmes et al., 2000) which report lower levels of oxidation than in the oxic–anoxic interface (Sansone and Martens, 1978; Reeburgh et al., 1991).

4.3 CH<sub>4</sub> and N<sub>2</sub>O emission in the Southern Ocean

Highly dynamic gas exchanges were registered in the KPR, with source and sink scenarios for N<sub>2</sub>O and only a source scenario for CH<sub>4</sub>. Since the mean wind speed did not exceed 14 m s<sup>-1</sup>, LM86 and W92 parameterizations represent the more conservative overestimation estimates of gas exchange in the area (Frost and Upstill-Goddard, 2002). The gas inventories in the ML reflect the effect of gas transport mainly via turbulent mixing and advection, which can be accelerated by the action of wind as well as by the microbial activity in surface waters. The ML depth did not correlate with wind speed (rs: 0.31, p < 0.05) and this aids in estimation of the gas content in the ML and whether it is a result of in situ production or consumption. CH<sub>4</sub> fluxes were higher at stations located at the PF and A3, where phytoplanktonic blooms were observed (see Table 2), but the tendency was the reverse for N<sub>2</sub>O, with an influx into the aforementioned stations. CH<sub>4</sub> emission rates during this study were higher than previously measured (Table 2), with a range of 0.1 to 3.0 µmol m<sup>-2</sup> d<sup>-1</sup> for the Pacific Ocean (Bates et al., 1996; Holmes et al., 2000; Sansone et al., 2001) and 0.5 to 9.7 µmol m<sup>-2</sup> d<sup>-1</sup> for the Atlantic Ocean (Oudot et al., 2002; Forster et al., 2009). In the South Pacific ocean (10°–64°S, 140° E), crossing the PF, Yoshida et al. (2011) reported CH<sub>4</sub> fluxes ranging from 2.4 to 4.9 µmol m<sup>-2</sup> d<sup>-1</sup>.

In the case of N<sub>2</sub>O, the estimates in this study were in the expected range for the oligotrophic open ocean (Nevison et al., 1995). N<sub>2</sub>O undersaturation and the concomitant influx were estimated, although this situation has not yet been well described for the Southern Ocean. N<sub>2</sub>O sinks can occasionally be observed (Butler et al., 1989; Law and Ling, 2001), which can be explained by probable N<sub>2</sub>O assimilation by N-fixing microorganisms. This process may be responsible for the estimated N<sub>2</sub>O influx.

5 Implications

The dynamics of both gases differ substantially both spatially and vertically (surface to 500 m depth), indicating that different mechanisms are being activated to produce an active gas during recycling. Our findings also show that, in areas of active fertilization and biogenic particle accumulation, CH<sub>4</sub> accumulates while N<sub>2</sub>O becomes depleted. This study suggests that the Antarctic Polar Zone plays a significant role in surface CH<sub>4</sub> production and subsequent air–sea gas exchange. These results did not agree with some previous studies of artificial fertilization experiments in the Southern Ocean, although only a few studies of this nature exist. This indicates that the turnover and evolution of microbial communities in mesoscale structures are fundamental for the development of substrates and conditions for CH<sub>4</sub> regeneration. Surface N<sub>2</sub>O production via nitrification does not seem to respond to spatial patterns of natural iron fertilization, but N<sub>2</sub>O consumption appears to be intensified in areas with higher dFe supply. However, in subsurface water, N<sub>2</sub>O accumulation seems to take place via nitrification.

Acknowledgements. We would like to thank the captain and crew of the RV Marion Dufresne. We recognize the support of both project leaders (Stephane Blain and Bernard Queguiner). This work was supported through the French Research programme of INSU-CNRS LEFE-CYBER (Les enveloppes fluides et l’Environnement – Cycles biogéochimiques, environnement et ressources), the French ANR (Agence Nationale de la Recherche, SIMI-6 programme, ANR-10-BLAN-0614), the French CNES (Centre National d’Etudes Spatiales) and the French Polar Institute IPEV (Institut Polaire Paul-Emile Victor). We are also grateful to Louise Oriol and Stephane Blain for nutrient data and Marine Lasbleiz for the HPLC analysis of chlorophyll measurements. The altimeter and colour/temperature products for the Kerguelen area were produced by Ssalto/Duacs and CLS with support from CNES. We also recognize all our colleagues who contributed to KEOPS 2. C. Fernández and L. Farias were supported by the Proyecto Ecos-Conicyt C09B02 and the International Associated Laboratory MORFUN. C. Fernández received partial support through Fondap no. 15110027. L. Farias performed the analysis of samples obtained in the KEOPS 2 cruise with FONDECYT no. 1120719. This is a contribution to the FONDAP-CONICYT programme (no. 15110009).

Edited by: S. Blain
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