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# Distribution of barium in the Weddell Gyre: Impact of circulation and biogeochemical processes

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### ABSTRACT

The Southern Ocean data base of dissolved barium (Ba<sub>d</sub>) has been augmented with two densely sampled sections across the Weddell Gyre sampled by the icebreaker FS Polarstern during February and March 2005. Bad was found to be relatively high in the surface layer as compared to the adjacent waters north (Antarctic Circumpolar Current) and east (Antarctic Zone of the Indian sector). Compared to the inflowing water into the Weddell Gyre and also to the surface water, the deep water is characterized by a significant  $Ba_d$ enrichment. Depletion of Bad in the surface layer is associated with well-known biogeochemical processes, but we speculate that also sea-ice formation leads to the extraction of Ba from solution via barite (BaSO<sub>4</sub>) precipitation. The particulate Ba settles down the water column and redissolves in deeper water where undersaturation of barite is prevalent. In the bottom layer, enrichment of  $Ba_d$  occurs, exhibited as a  $Ba_d$ maximum, which is caused by Ba efflux from the sediments. In recently formed Weddell Sea Bottom Water, though, a Bad minimum is observed, imposed by the shelf water component of bottom water, which has relatively low Ba<sub>d</sub> concentration. Like in other Southern Ocean regions, throughout the water column a strong correlation exists between Bad and dissolved silicate, although the relationship is different from that in the Antarctic regions to the east. In a tentative budget, the deep water is found to be supplied with  $Ba_d$ mainly by the surface layer, but also to some extent by the bottom layer. Weddell Sea Deep Water, enriched in Ba<sub>d</sub>, is transferred out of the Weddell Gyre into the ACC as Antarctic Bottom Water, making the gyre a source of Ba<sub>d</sub> to the deep and abyssal world oceans.

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#### 1. Introduction

Like other chemical species, the distribution of barium (Ba) in the oceans is controlled by water circulation. Consequently, barium has been utilized as a water mass tracer (e.g., Chan et al., 1977; Falkner et al., 1994). However, many studies have found that there is non-conservative behavior of Ba as well, namely, due to biologically associated uptake followed by dissolution (Chan et al., 1977; Jeandel et al. 1996). These conditions impose that Ba be categorized as a bio-intermediate element. Higher biomass and productivity in the upper mixed layer usually coincide with higher Ba concentrations in suspended matter (e.g., Cardinal et al., 2005; Jacquet et al., 2007a). Most of this upper mixed layer particulate, non-lithogenic Ba consists of Ba absorbed by or adsorbed on planktonic material (Cardinal et al., 2005; Collier and Edmond, 1984; Jacquet et al. 2007b, 2008), though

the presence of the mineral barium sulfate has been also demonstrated in living marine phytoplankton (Fresnel et al., 1979; Gayral and Fresnel, 1979) and in freshwater and marine benthic microbial organisms (e.g., Bertram and Cowen, 1997; Hopwood et al., 1997). Also, celestite ( $SrSO_4$ ) of acantharians appears to be significantly enriched in barium as compared to the surrounding seawater (Bernstein et al., 1998); these pelagic protozoans constitute a significant part of plankton in the Southern Ocean.

Underneath the surface mixed layer, it is well established that barite is the major carrier of Ba in suspended matter (Dehairs et al., 1980; Jacquet et al., 2007b; Stroobants et al., 1991), although the deeper parts of the ocean are generally undersaturated with respect to this mineral (Monnin et al., 1999; Rushdi et al., 2000). Barite formation occurs essentially in the mesopelagic zone (100–1000 m) within aggregates of decomposing organic detritus (Cardinal et al., 2005; Dehairs et al., 1980, 1990; Stroobants et al., 1991; van Beek et al., 2009). Experimental work by Ganeshram et al. (2003) showed that barite crystals form during the degradation of phytoplankton material. Mesopelagic particulate Ba-barite was observed to be

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correlated with the rate of oxygen consumption (Dehairs et al., 1997) and subsurface bacterial activity (Dehairs et al., 2008; Jacquet et al., 2008).

Because of the link of barium and barite to ocean biological productivity, and the fact that barite is well preserved in sediments due to its low solubility, barium and barite have been used as a paleoproxy, i.e., a paleo recorder of export production (e.g., Dymond et al., 1992; Paytan and Griffith, 2007). Since dissolved Ba in the oceans is the main constraint for the chemistry of barite, it is essential that the large-scale Ba distribution and the barium cycle in the world oceans be known. The present study explores the distribution of dissolved Ba (and for comparison that of silicate) in the water column of the Weddell Gyre, an area for which there is a dearth of data on Ba, despite its importance in terms of formation of ventilated deep and bottom water, upwelling and sea-ice formation. Earlier data for the Weddell region were obtained during the GEOSECS project, but they consist of just a few full-depth profiles, plus some surface and bottom water data (Chan et al., 1977; Östlund et al., 1987). Expedition ANT-XXII/3 in 2005 offered the possibility to investigate Ba along the Prime Meridian and at a section across the Weddell basin (Fig. 1), thereby significantly expanding the Southern Ocean data set, which thus far consists of sections along 30°E (WOCE I6; unpublished results of Dehairs et al.), 145°E (WOCE SR3; Jacquet et al., 2004), a section in the Crozet-Kerguelen Basin (Jacquet et al., 2005) and a series of profiles in the Indian sector (INDIGO 1 and 3; Jeandel et al., 1996).

#### 2. Material and methods

Data were collected (Fig. 1) during FS Polarstern cruise ANT-XXII/3 from Cape Town, South Africa to Punta Arenas, Chile, 22 January to 6 April 2005 (Fahrbach, 2006). Water was sampled with the rosette sampler coupled to the CTD instrument (conductivity temperature depth; SBE911plus) throughout the water column, with a bias to the upper 1000 m. Temperature and salinity were measured with a precision of 0.001 °C and 0.002, respectively (Fahrbach, 2006).

#### 2.1. Dissolved barium

Volumes of 15 ml of unfiltered seawater were sampled in polypropylene vials (Nalgene), which were rinsed three times with the sample seawater. Samples were acidified with 15 µl HCl (Merck Suprapur) and stored at room temperature until analysis in the home laboratory. No filtration of the seawater was attempted based on the well documented knowledge that dissolved Ba represents, in general, a very large fraction (>99%) of total Ba. In productive surface mixed



**Fig. 1.** Map of the Weddell Sea and environs with station locations during cruise ANT-XXII/3. Arrows denote the schematic surface circulation in the region.

layers this may be different, but also here particulate Ba is usually <3%of total Ba (e.g., see Jacquet et al., 2007a; 2008). Dissolved Ba was measured using an isotope dilution method described below; in fact, the dissolved Ba measurements presented here represent soluble Ba, consisting of dissolved Ba plus a very small fraction (generally <1% of total Ba) that is generated from the particulate Ba pool as a result of the acidification. For the sake of simplicity, we prefer to use the term dissolved Ba (Ba<sub>d</sub>) in this paper. Sample preparation is as follows: 1 g of seawater is spiked with 0.7 g of <sup>135</sup>Ba-spike solution, yielding a <sup>138</sup>Ba/<sup>135</sup>Ba ratio between 0.7 and 1 to minimize propagated error (Klinkenberg et al., 1996; Webster, 1960). Subsequently, the sample is diluted with acidified (nitric acid) Milli-Q grade water to a final weight of 30 g. Blanks consist of acidified (with nitric acid) Milli-Q water. Quantities of sample, spike and dilution water were accurately assessed by weighing. Isotope ratios were measured with sector-field inductively-coupled plasma mass spectrometry (SF-ICP-MS; Element 2 Thermo Finnigan). Reproducibility of our method is  $\pm 1.5\%$  (RSD) as tested on repeat preparations of reference solutions. Average Ba values obtained for reference waters SLRS-3 and an in-house standard (a Mediterranean Sea standard prepared by C. Jeandel) were  $13.48 \pm$  $0.21 \ \mu g \ l^{-1} (1\sigma)$  with RSD of 1.55% and  $10.49 \pm 0.29 \ \mu g \ l^{-1} (1\sigma)$  with RSD of 2.75%, respectively, which is in good agreement with the certified values (SLRS-3:  $13.4 \pm 0.6 \ \mu g \ l^{-1}$  and OMP:  $10.4 \pm 0.2 \ \mu g \ l^{-1}$ ). Overall precision (including sampling precision) based on 6 dissolved Ba profiles sampled in a hydrographically stable environment is  $\pm 0.3 \,\mu g \, l^{-1}$  (1 $\sigma$ ), with an RSD of 5% (Dehairs et al., 2008; Jacquet, 2007); for further details we refer to the latter two studies. Bad concentrations are expressed in nmol  $kg^{-1}$ .

#### 2.2. Silicate

Nutrients were measured on board with a Technicon TRAACS 800 rapid flow autoanalyzer (see also Fahrbach, 2006) by the Royal Netherlands Institute for Sea Research (NIOZ, Texel). Accuracy was determined using stock standards diluted in low-nutrient sea water, as prepared in the home lab by weighing. The precision for silicate was estimated to be 0.66  $\mu$ mol l<sup>-1</sup> at two calibration stations, where all 24 rosette bottles were fired at the same depth. Part of the surface layer nutrient data of this cruise have also been reported by Hoppema et al. (2007). Silicate (or silicic acid) concentrations are expressed in  $\mu$ mol kg<sup>-1</sup>.

#### 2.3. BaSO<sub>4</sub> saturation index calculation

The barite saturation index (SI) is defined as the ratio of the ionic product of barium sulfate (Q) to the solubility product of barite ( $K_{sp}$ ):

$$SI = \frac{Q}{K_{\rm sp}} = \frac{m_{\rm Ba(aq)} \cdot m_{\rm SO_4(aq)} \cdot \gamma_{\rm BaSO_4(aq)}^2}{K_{\rm sp}} \tag{1}$$

in which m is the measured concentration (molality) and  $\gamma$  the activity coefficient of aqueous barium sulfate. Monnin (1999) has developed a model of the Na–K–Ca–Mg–Sr–Ba–Cl–SO<sub>4</sub>–H<sub>2</sub>O system that allows the calculation of the solubility and saturation indices of some minerals, including barite, as a function of the solution composition, temperature (up to 200 °C) and pressure (up to 1 kbar). It has been used to investigate the saturation state of the ocean with respect to pure barite (Monnin et al., 1999) and Sr-substituted barites (Monnin and Cividini, 2006). Alternative models restricted to the case of seawater, such as the one proposed by Rushdi et al. (2000), have been shown to give similar results when applied to the GEOSECS Ba data. The criterion for equilibrium retained in these studies is a saturation index between 0.9 and 1.1, which has been inferred from an evaluation of the mean accuracy of the solubility calculations during model development (Monnin, 1999). The measured quantities

used are dissolved barium molarity, salinity, pressure and potential temperature. The concentrations of K, Ca, Mg, Ba, Sr, SO<sub>4</sub> and Cl are calculated from the measured salinity and the composition of standard seawater of salinity 35 using:

$$c_i(S) = c_i(35) \times \frac{S}{35} \tag{2}$$

with  $c_i(S)$  the concentration (molarity) of the *i*th element for the sample with salinity *S*. The Na concentration is calculated from the electroneutrality condition. Molarities are converted to molalities using the VOlumetric\_PrOperties (VOPO) code (Monnin, 1994). The thermodynamic model then calculates the stoichiometric BaSO4(aq) activity coefficient, the barite solubility product and the barite saturation index at the given temperature and pressure.

#### 3. Hydrographic background

The region of investigation is situated in the Antarctic Zone, i.e., south of the Polar Front. The boundary between the Antarctic Circumpolar Current (ACC) and the Weddell Gyre is found at about 56°S on the Prime Meridian (Klatt et al., 2005). Thus the majority of data stem from the Weddell region. North of 56°S, the potential temperature maximum of the Upper Circumpolar Deep Water (UCDW) can be distinguished, centered at 500-600 m (Fig. 2). Circumpolar Deep Water (CDW), characterized by a maximum in the potential temperature  $(\theta)$ , is transferred into the Weddell Gyre near its eastern end. We observed the temperature maximum at 64-69°S at 150-400 m depth reaching 0.8-1 °C (Fig. 2); this coincides essentially with the westward flowing southern limb of the Weddell Gyre. Note that the  $\theta$  maximum layer is interrupted near 66°S by a lower  $\theta$  maximum around the undersea mountain Maud Rise. At the northern part (56-63.5°S) of the Prime Meridian section, constituting the northern limb of the gyre with generally eastward flow, a  $\theta$ maximum was observed as well, but it reaches only about 0.5 °C (Fig. 2). This is CDW that has been modified by mixing with waters above and below during its course through the Weddell Gyre. The  $\theta$ maximum layer is separated from the base of the surface layer (identified by a temperature minimum) by a sharp pycnocline. In the upper part of the surface layer, temperatures were often >1.0 °C, but in the coastal region it was significantly colder. These are summer temperatures; in winter the entire surface layer is at, or close to, the freezing point of about -1.85 °C. Note that the surface layer is deeper in the coastal zone than elsewhere in the basin. The Weddell Gyre is a divergent feature with upwelling of deep water, mainly more or less modified CDW; the surface layer is eventually generated from this upwelled water. In the deep Weddell basin,  $\theta$  decreases monotonically from the  $\theta$  maximum to a minimum at the sea floor. Weddell Sea Bottom Water (WSBW) is present along the section, being defined by  $\theta < -0.7$  °C. Between the lower boundary of the CDW at 0 °C and the WSBW, the most voluminous water mass of the Weddell Gyre is found, the Weddell Sea Deep Water (WSDW). It is replenished by upward mixing of WSBW with CDW, by local water mass formation processes (Orsi et al., 1993), but also by advective transport from the east (Meredith et al. 2000; Hoppema et al., 2001). In the deep Weddell basin,  $\theta$  decreases monotonically from the  $\theta$  maximum to a minimum at the sea floor.

#### 4. Results

The most conspicuous feature in the Bad distribution is the difference between the upper 500-1000 m and the deep basin (Fig. 3). Concentrations below about 500–1000 m mainly range between 90 and 105 nmol kg $^{-1}$ , which appears to be the background level of Bad of the deep Weddell-Enderby basin. At the Prime Meridian, Ba<sub>d</sub> concentrations in the deep basin (>1000 m) appear to be highest in the central part (Fig. 3A), reaching up to 107 nmol kg $^{-1}$ between 2000 and 3000 m at 61.5°S. Over the seamount Maud Rise (65-66°S) and over the northernmost part of the Weddell Gyre (56-58°S), deep and bottom water Ba<sub>d</sub> concentrations are the lowest. Further south, between Maud Rise and the Antarctic margin (67-68°S), Bad in the bottom layer is again slightly higher (up to 100 nmol kg<sup>-1</sup>). In the deep Weddell Sea (Fig. 3B; below 1000 m) the Ba<sub>d</sub> concentration range is 84-104 nmol kg<sup>-1</sup>, with lowest values in the northwestern part (in the vicinity of the peninsula margin) and highest in the southeastern part of the Weddell transect.

The deep waters (i.e. below 200 m in the central part of the Weddell section and below 600 m toward the western and the eastern margins of the basin) are separated from an upper layer which has concentrations <90 nmol kg<sup>-1</sup>, largely coinciding with the surface mixed layer (Fig. 3B). Since surface waters of the Weddell Gyre are eventually formed from upwelled deep water, the lower Bad contents in surface waters must be the result of non-conservative processes which cause a partial depletion of Ba<sub>d</sub>. Between 58° and 65°S at the Prime Meridian, the Ba<sub>d</sub> isolines in the upper 500–1000 m exhibit an upward hump, typical of the divergent Weddell Gyre with westward flow in the south and eastward flow in the north, which is also seen in the distributions of other variables like  $\theta$  (Fig. 2) and silicate (see below). As a consequence, the vertical extent of the Bad depleted upper layer is zonally variable, being larger near the northern and southern boundaries of the section. North of 56°S, i.e. outside the Weddell Gyre, the Ba<sub>d</sub> depletion zone appears to overlap with the  $\theta$ 



Fig. 2. Section at the Prime Meridian cutting the Weddell Gyre and the southernmost part of the Antarctic Circumpolar Current, contoured for potential temperature (°C) at cruise ANT-XXII/3 in 2005.



**Fig. 3.** Sections contoured for dissolved barium (nmol  $kg^{-1}$ ) during cruise ANT-XXII/3 in 2005, A) at the Prime Meridian cutting the Weddell Gyre and the southernmost part of the Antarctic Circumpolar Current; and B) across the Weddell Sea from Kapp Norvegia (right) to Joinville Island at the tip of the Antarctic Peninsula (left).

maximum core of the UCDW (Fig. 2). As UCDW waters were reported to contain less Ba<sub>d</sub> than the deep waters of the Antarctic Zone (Jacquet et al., 2004; Jeandel et al., 1996), this hints that the low Bad in the upper water column of the Weddell Gyre could, at least partly, result from advection of CDW into the Weddell Basin via the westward flowing southern limb of the gyre. It should be appreciated that the front between the ACC and the Weddell Gyre at 56°S also constitutes a divide between the near-surface water with lower Bad to the north and waters with higher Ba<sub>d</sub> to the south. In the coastal current at the southern end of the Prime Meridian, the mixed layer is deep because of convection, mixing and downwelling (see also  $\theta$  distribution; Fig. 2), which also leads to a deeper penetration of low-Bad waters. In addition, the Bad concentration in the upper few hundred meters of the coastal current appears to be lower (about 77 nmol kg<sup>-1</sup>) than north of it along the section, where it reaches about 85 nmol kg $^{-1}$ . In the area between 64°S and 66.5°S the Ba<sub>d</sub> depletion reaches greater depths than north and south of it. Here, Maud Rise exerts influence on the entire water column: as circulation-topography interaction induces a Taylor column over the rise, the mixed layer extends deeper over the crest of Maud Rise than over its flanks.

Along the Weddell Sea section, the near-surface layer exhibits similar  $Ba_d$  concentrations (Fig. 3B) as observed at the Prime Meridian. At both ends of the section, the low- $Ba_d$  layer also extends deeper than in the central part. At the continental shelf break of the Antarctic Peninsula, the low- $Ba_d$  surface layer is linked to a plume of  $Ba_d$  depleted water hugging the continental slope down to about 4000 m depth. This plume consists of Weddell Sea Bottom Water, which has very recently been formed at the southern and western shelves of the Weddell Sea and which has not yet reached the bottom of the central basin (Fahrbach et al., 2001; Gordon et al., 1993). Since WSBW is a mixing product of CDW and shelf water, it carries the low-Ba<sub>d</sub> properties of the latter water mass. In the WSBW at the continental slope, the shelf water component is particularly large; further along its way through the Weddell Sea, mixing with adjacent deep waters will reduce the overall contribution of the shelf water component to the WSBW, thus also increasing the Ba<sub>d</sub> concentration.

#### 5. Discussion

The concentrations of Bad are generally higher in Weddell Gyre surface waters compared to the Antarctic Zone in the Indian sector of the Southern Ocean (Jacquet et al., 2005; Jeandel et al., 1996). Such zonal variation is mainly caused by varying upwelling activity of Baenriched deep waters, although zonal differences in phytoplankton activity probably contribute as well. The latter usually is highest in the Subantarctic zone and Polar Frontal Zone to the north and decreases southwards into the Antarctic Zone, where it increases again in the vicinity of the ice edge (Jochem et al., 1995; Moore et al., 1999; Savoye et al., 2004). Our Weddell data were collected during summer, at the height of the growth season, and thus we expect the impact of biological uptake on Ba<sub>d</sub> to have been at its climax. The smaller Ba<sub>d</sub> depletions in Weddell surface waters could thus reflect lower production in the Weddell region compared to elsewhere. However, it is likely that this smaller apparent Bad depletion is in part due to the intensity of upwelling in the divergent Weddell Gyre transferring Baenriched waters into the surface layers. Additionally, deep water that

is being upwelled into the surface layer appears to have a higher  $Ba_d$  concentration in the Weddell Gyre than in the Antarctic Zone of the Indian sector (*cf.* Jacquet et al., 2004; Jeandel et al., 1996).

#### 5.1. Correlation between barium and silicate

The distribution of Ba<sub>d</sub>, both lateral and vertical, is known to show similarities with that of silicate in several ocean provinces (e.g., Chan et al., 1977; Jacquet et al., 2007a; Jeandel et al., 1996). We explore the occurrence of such a similarity for the Weddell Gyre with the corresponding silicate sections along the Prime Meridian and across the Weddell Sea (Fig. 4).

The vertical profile of silicate in the Weddell Gyre is typical for that of the major nutrients, i.e., relatively low concentrations in the surface layer, a maximum at intermediate depth, and a decrease towards the sea floor. Frequently, a silicate maximum is observed in the bottom layer (Edmond et al., 1979; Hoppema et al., 1998). Our data demonstrate a high agreement between the distributions of silicate and Ba<sub>d</sub> in the Weddell Gyre (Figs. 3 and 4). Even the lateral variation of the extent of the surface layer appears to be congruent. This condition is largely due to hydrographic processes such as upwelling and convective overturning of the surface layer.

Newly formed WSBW at the continental slope off Joinville Island can be recognized by its low silicate and Ba<sub>d</sub> concentrations (see left side of the Weddell section in Figs. 3B and 4B). It is intriguing that even the wide, but not very pronounced, silicate minimum at 3000– 3500 m emerging from the lower continental slope off Kapp Norvegia (see right side of the Weddell section in Figs. 3B and 4B) appears to correspond to a Ba<sub>d</sub> minimum. This core of ventilated water, characterized by a CFC (chlorofluorocarbon) maximum (Hoppema et al., 2001; Meredith et al., 2000) and thus a large surface water component, originates east of the Weddell Gyre. Just like WSBW from the Weddell Sea, it is characterized by relatively low concentrations of silicate and  $Ba_d$ , although the signal has been attenuated during transport from the region of origin.

The deep silicate maximum (Fig. 4), which is slightly visible in plots of silicate versus salinity (Fig. 5) is caused by dissolution of biogenic silica at depth (Rutgers van der Loeff and Van Bennekom, 1989). There is no concomitant coherent Bad maximum – the variability of Bad in the deep Weddell Gyre appears too large although the Bad data from the Weddell section seem to point to a weak feature (Fig. 5B). Differences between the deep distributions of silicate and Ba<sub>d</sub> should not be surprising, because the saturation state of ocean waters with respect to silica/opal and BaSO<sub>4</sub> is different, as are the dissolution kinetics of these minerals. The water column in the Weddell Gyre is oversaturated with respect to barite in the upper 1500 m with barely any regional variability (Fig. 6). This indicates that no dissolution of BaSO<sub>4</sub> should occur in the upper 1500 m of the water column. Based on the saturation state, no large Bad depth gradients are expected. Silicate shows a much larger surface to depth gradient since the entire oceanic water column is undersaturated with respect to opal (Brzezinski et al., 2003; Hurd, 1972).

Regression of  $Ba_d$  against silicate shows a high correlation for the combined Prime Meridian and Weddell sections ( $r^2 = 0.81$ ; Fig 7). The  $Ba_d$ -silicate regression for the Weddell region is slightly different from that for a meridional section south of the Polar Front along 30°E (Fig. 7: previously unpublished data by F. Dehairs and C. Jeandel of the CIVA-1 cruise 1993, WOCE 16 line). For a given silicate concentration,



**Fig. 4.** Sections contoured for dissolved silicate ( $\mu$ mol kg<sup>-1</sup>) during cruise ANT-XXII/3 in 2005, A) at the Prime Meridian cutting the Weddell Gyre and the southernmost part of the Antarctic Circumpolar Current; and B) across the Weddell Sea from Kapp Norvegia (right) to Joinville Island at the tip of the Antarctic Peninsula (left).



Fig. 5. Plots of silicate (µmol kg<sup>-1</sup>) vs. salinity (A, C) and dissolved Ba (nmol kg<sup>-1</sup>) vs. salinity (B, D) for the Weddell and the Prime Meridian sections to investigate property maxima in the deep water.

the Ba<sub>d</sub> concentrations in the Weddell system exceed those at 30°E by about 5 nmol kg<sup>-1</sup> – note that the 30°E section is quite near the eastern rim of the Weddell Gyre. Also, the slope of the Ba<sub>d</sub> versus silicate regression is larger at 30°E, whereas the Ba<sub>d</sub> intercept at zero silicate is smaller (differences are significant at p<0.005). In the Weddell region, the higher Ba<sub>d</sub> level highlights the decoupling between barium and silicate and points to a mechanism for retaining Ba<sub>d</sub> within the gyre system relative to silicate. A likely mechanism is Ba translocation to particles in surface waters and Ba efflux from the sediments as discussed below. However, a smaller biological Ba to Si uptake ratio in the upper waters of the Weddell Sea, as compared to the ACC, could also lead to the observed differences in Ba vs. silicate regressions between both regions.

## 5.2. Barium depletion in the surface layer

Depletions of nutrients or  $TCO_2$  in the surface layer have been used frequently in the Weddell region to determine the net community production from the onset of the growth season until the time of measurement (e.g., Hoppema et al., 2007). We are using an approximate depletion here to obtain the seasonal changes of the Ba<sub>d</sub> concentration between the winter and the time of sampling. Usually, depletions are computed from the vertical integration of the concentrations above the remnant Winter Water, which is identified by its temperature minimum, and is assumed to reflect the initial prebloom conditions. Since sampling resolution in the upper layers was insufficient to warrant a reliable integration over the surface layer, we simply consider the difference in  $Ba_d$  between the temperature minimum layer and the shallowest sample (at about 20 m). This difference in  $Ba_d$  concentrations is proportional to, and an approximate measure of, the depletion.

At the Prime Meridian, the  $Ba_d$  difference between the temperature minimum and surface waters is generally in the range 0– 6 nmol kg<sup>-1</sup>, with an average of  $3.6 \pm 3.9$  nmol kg<sup>-1</sup> (Fig. 8A). A slight trend of higher values towards the centre of the transect may be discerned. At the Weddell section, the  $Ba_d$  difference exhibits a range of -2 to 6 nmol kg<sup>-1</sup>, with few deviations to higher and lower values (Fig. 8B). The Ba difference is independent of temperature within the temperature minimum (Fig. 8C and D). At the Weddell section, the temperature minimum remains close to the freezing point of seawater of -1.85 °C and it therefore appears that the observed  $Ba_d$  depletions are real and not affected by erosion of the temperature minimum layer.

As a bio-intermediate element, Ba becomes depleted in the surface layer with respect to the deep water underneath it (e.g., Chan et al., 1977; Jacquet et al. 2007a). Our data show that surface water Ba depletions reaching up to 8 and 12 nmol kg<sup>-1</sup> (for the Prime Meridian and Weddell section, respectively; Fig. 8) are generated within a few months after the end of winter. There are several processes that may be responsible for this: first, active or passive biologically associated uptake of Ba; and second, abiotic precipitation of Ba as the mineral barite (BaSO<sub>4</sub>). As to the first cause, there are only very few phytoplankton



Fig. 6. Barite saturation state SI (ion product of Ba and sulfate divided by the solubility of barite) for the sections along the Prime Meridian (A) and across the Weddell Sea (B). If SI>1 the water is supersaturated with barite.

species known to actively take up Ba (e.g., Gayral and Fresnel, 1979). However, high particulate Ba concentrations have been reported in Southern Ocean surface waters (Cardinal et al., 2005; Jacquet et al., 2007a), usually associated with elevated biomass. This Ba is generally not present as barite (Dehairs et al., 1980; Stroobants et al., 1991), but is adsorbed onto biogenic matter (diatom tests; Sternberg et al., 2005) or is incorporated into the skeletal matrix, e.g., in acantharians (Bernstein et al., 1998; Jacquet et al., 2007b). As to the second cause, our results for Weddell Gyre surface waters point to a barite supersaturation in the upper water column with a maximum saturation index of 1.4 (Fig. 6). Together with the Black Sea (Falkner et al., 1993), the Weddell Sea appears to be one of the few regions in the world's ocean where BaSO<sub>4</sub> supersaturation prevails. Note that when barite saturation is reached, equilibrium should be the rule, as expected for salts whose kinetics of dissolution/precipitation are generally believed to be fast. Nevertheless, large BaSO<sub>4</sub> supersaturations have also been documented for the sedimentary environment (e.g., Aloisi et al., 2004).

In the seasonally ice-covered Weddell Gyre, a third cause of depletion should be considered, namely, barite precipitation during sea-ice formation. In an already oversaturated surface ocean, salinity and solute concentrations in brine channels of the ice may reach very high values (e.g., Anderson and Jones, 1985), eventually leading to supersaturation of barite, which will result in inorganic precipitation. The large biomass of sea-ice algae, visible as green or brown ice at the base of ice floes, may further enhance Ba depletion as a result of Ba adsorption on algal cells and may also result in barite formation due to organic matter degradation within the sea-ice (Carson, 2008). During the EPOS-2 cruise in the Weddell Scotia Confluence in 1988, high particulate Ba concentrations were measured in brown ice (up to 8 nM Ba, i.e., about 40 times more than the usually observed surface water particulate Ba concentrations), correlating with high POC contents (F. Dehairs, unpublished results). Carson (2008) reports dissolved Ba concentrations in sea-ice brine ranging widely from surface water values to as low as 6 nM, suggesting that significant Ba translocation from the dissolved to the particulate phase is likely in sea-ice brines. On the other hand, it must be mentioned that in laboratory experiments on freezing sea water no evidence for barite formation was found (Taylor et al., 2003; K.K. Falkner, personal communication). In the Arctic, strong depletions of Ba were found in the surface layer associated with the ice edge (Falkner et al., 1994),



**Fig. 7.** Dissolved Ba (nmol kg<sup>-1</sup>) plotted against silicate (µmol kg<sup>-1</sup>) for data from the Prime Meridian and Weddell sections combined. Also shown is the regression for a section at 30°E (1993, CIVA1 cruise, WOCE I6 line), where only stations south of the Polar Front were included.

suggesting that processes within the sea-ice play a role in Ba depletion; those authors hint at intensified biological activity as the main cause for this depletion. We emphasize that inorganic barite precipitation is only speculated here. However, whatever the exact mechanism, be it of biological or inorganic nature, Ba<sub>d</sub> depletion seems to be associated with sea-ice formation.

It is worth noting that barite precipitation associated with sea-ice formation occurs in autumn and winter, hence the process as such cannot explain the observed Bad depletion of the seasonal summer mixed layer. However, barite or particulate Ba formed within the seaice will eventually be released in the surface waters, either during draining and flushing of brine waters or ultimately when the ice is melted at the end of the winter. The released barite cannot dissolve in the supersaturated surface waters but, when associated with aggregates of ice algae, it may settle to the undersaturated deep ocean where it will eventually dissolve. Overall, such a process would subtract Ba<sub>d</sub> from surface waters and redistribute it in the deep ocean. As to the seasonal depletion, the melting of sea-ice at the end of winter will dilute the upper part of the winter mixed layer more than the lower part, which also has an impact on the Ba<sub>d</sub> concentrations. Clearly, the mechanism described above would lead to variability of the Bad depletion. Due to ice motion (which is different from the movement of the underlying water) and spatially variable rates of ice formation, the variation of Bad depletion is further enhanced and even enrichment (i.e., negative depletion; see Fig. 8B) of Bad is possible. We speculate that barite dissolution may even occur within the melting ice matrix.

It is interesting to note that the larger surface Ba depletion around 62°S on the Prime Meridian (Fig. 8A) appears to coincide with elevated  $Ba_d$  concentrations in the deep basin between 2000 m and the bottom (Fig. 3A). This may well hint at a mechanism whereby barite is being formed in the surface layer and is subsequently transferred to the deep ocean. At greater depths where undersaturation prevails (Fig. 6), dissolution of barite causes elevated concentrations of  $Ba_d$ . It should be added that this mechanism is not only active in the central Weddell Gyre; it is merely most prominent there. In other parts of the basin, the  $Ba_d$  concentration is to some degree determined by dissolution of barite — if the rate of surface depletion of  $Ba_d$  is smaller, also less barite will be dissolved at depth. The vertical correspondence between Ba depletion in the surface and Ba enrichment at depth is not perfect because water currents

(strength and direction) are depth-dependent. It makes sense that the correspondence is best in the central gyre, because the currents are weakest there (Fahrbach et al., 1994). It should be realized that the residence time of the deep water where  $Ba_d$  enrichment occurs is many years, and thus the enrichment process occurs on an accordingly long time scale.

#### 5.3. Bottom layer enrichment

Both silicate and Bad are enriched in the bottom layer of the Weddell Sea; actually, the highest absolute concentrations of both species were observed there (Figs. 3B and 4B). For silicate, this has been previously reported by Edmond et al. (1979). Note, in contrast, that newly formed WSBW (which is the source of all bottom water in the Weddell Gyre) along the slope of the Antarctic Peninsula (Figs. 3B and 4B) is characterized by minima of silicate and Ba<sub>d</sub>. This provides evidence that enrichment of bottom water occurs locally in the Weddell-Enderby basin. Even the spatial distributions of bottom layer enrichment of silicate and dissolved Ba largely conform: Strong enrichments were found along most of the Weddell section (Figs. 3B and 4B), and at the Prime Meridian especially in the southern part (66.5-68.5°S; Fig. 3A and 4A). In the northern part of the Prime Meridian section at 55.5-56.5°S, a silicate maximum was observed at the bottom, but this appears to be a local deep water phenomenon, as no Ba<sub>d</sub> analogue was found at that location. Hoppema et al. (1998) have demonstrated that strong enrichment of the bottom water with silicate occurs especially in the region off Kapp Norvegia, supported by a local bottom water recirculation cell. The same process could also explain the increased bottom water Bad concentrations in that region.

Enrichment of  $Ba_d$  in the bottom layer may have two causes. First, particulate barite could rapidly dissolve at these depths because of the pressure dependence of the solubility product of barite (Monnin et al., 1999; Monnin and Cividini, 2006). The saturation index SI for  $BaSO_4$  close to the seafloor at depths >4000 m drops to values <0.6 (Fig. 6) and enhanced dissolution of barite may only occur when  $SI \ll 1$  due to kinetic effects. However, this explanation is unlikely since we found evidence for  $Ba_d$  enrichment in the deep water column (see above), which hints at dissolution at depths well above the bottom layer. Second, there may be input of  $Ba_d$  from the sediments to the overlying water. Biogenic barium is abundantly present in sediment cores from the Weddell Sea and environs (e.g., Ó Cofaigh and Dowdeswell, 2001).



**Fig. 8.** Dissolved barium concentration difference between the temperature minimum layer and the near-surface in the surface layer vs. latitude at the Prime Meridian (A), and vs. longitude at the Weddell section (B). The same Ba difference plotted against the potential temperature of the temperature minimum layer at the Prime Meridian (C) and at the Weddell section (D).

The occurrence of enhanced vertical gradients of Ba<sub>d</sub> in the deepest samples combined with the fact that a significant epibenthic flux of dissolved Ba to the water column has been reported for other oceanic areas as well (e.g., McManus et al., 1994, 1998), suggests that efflux of Ba from the sediments contributes to elevated bottom water Ba concentrations. It is worthwhile adding that the spatial coincidence of Ba<sub>d</sub> and silicate enrichment lends high credibility to the case for Ba<sub>d</sub> input from the sediments, because opal dissolution occurs throughout the water column, and silicate efflux from the sediments has been reported in the Weddell basin (Edmond et al., 1979; Holby and Anderson, 1996; Rutgers van der Loeff and Van Bennekom, 1989).

A rough estimation of the Ba flux from the sediments is presented. We take the  $Ba_d$  enrichment in the bottom layer to be 5 nmol kg<sup>-1</sup>, where the vertical extent of that layer is about 500 m (Fig. 3B). The enrichment then amounts to 2500 µmol m<sup>-2</sup>. It has been generated at the time scale of the residence time of the WSBW in the bottom layer. The residence time is tentatively calculated as 6 years, using the surface area of the Weddell Gyre where WSBW occurs (assumed to be about  $1 \cdot 10^{12}$  m<sup>2</sup>), combined with the vertical extent (500 m) and the ventilation rate of WSBW (2.6 Sv =  $2.6 \cdot 10^6$  m<sup>3</sup> s<sup>-1</sup>; Fahrbach et al., 1994). This yields an efflux of  $Ba_d$  of

about 400  $\mu$ mol m<sup>-2</sup> year<sup>-1</sup>. This value falls well within the range reported for other open ocean environments, e.g., in the Equatorial Pacific: 160–1060  $\mu$ mol m<sup>-2</sup> year<sup>-1</sup> (McManus et al., 1999; Paytan and Kastner, 1996). Integrated over the Weddell Basin surface area of  $1 \cdot 10^{12}$  m<sup>2</sup>, the Ba<sub>d</sub> efflux is  $0.4 \cdot 10^9$  mol year<sup>-1</sup>.

#### 5.4. Enrichment of dissolved barium in deep water

Since water mass transformation and mixing take place in the Weddell–Enderby basin, the bottom water  $Ba_d$  enrichment is also transferred to other water masses, in particular the voluminous deep water. We can make a rough estimation of the enrichment of  $Ba_d$  in the deep Weddell basin using characteristics of the hydrography of the region. The main source of deep water to the Weddell Gyre is the inflowing CDW of the ACC near the eastern boundary. Other water masses are eventually derived from that source. At the Prime Meridian, the source water mass can be distinguished at depths of about 200–1500 m at 64–69°S (Fig. 2). Its upper boundary is the  $\theta$  maximum, while the lower boundary is distinguishable as a weak maximum of both TCO<sub>2</sub> and nutrients, occurring at  $\theta$  of about 0–0.2 °C (Hoppema et al., 1997). We compare the  $Ba_d$  concentration of the

CDW source water (marked as "inflow") with that of the remaining deep water masses at the Prime Meridian in a diagram of  $Ba_d$  against  $\theta$  (Fig. 9). The  $Ba_d$  concentration in the inflow CDW tends to be lower than that in the rest of the basin – recall that lower  $Ba_d$  concentrations were observed in the CDW of the Antarctic Circumpolar Current (see Results section and Fig. 3A). However, there is a large overlap of  $Ba_d$  values for the two sub-regions. At the highest  $\theta$  values, representing the core of the  $\theta$  maximum, which is associated with inflowing CDW,  $Ba_d$  is lowest (85–94 nmol kg<sup>-1</sup>). Beneath the  $\theta$  maximum of the inflow water,  $Ba_d$  is higher and similar to  $Ba_d$  in the low  $\theta$  maximum of the rest of the basin. In the deep Weddell basin with a  $\theta$  range of roughly 0.2 to -0.7 °C, the  $Ba_d$  concentration is constant within about 93–101 nmol kg<sup>-1</sup>. Some of the highest values of  $Ba_d$  are due to enrichment in the bottom layer.

Since Bad in water of about 0.5 °C in the central and northern Weddell Gyre along the Prime Meridian coincides with that of 0.5 °C in the inflow area (64–69°S) (Fig. 9), it appears that the upper portion of the inflow water with a high  $\theta$  maximum (>0.5 °C) is upwelled into the surface layer during its traverse of the gyre. Of note is that at the Prime Meridian, we do not catch the full extent of upwelling of low-Bad subsurface water in the inflow. Part of the upwelling has already taken place in the eastern Weddell Gyre between the eastern boundary and 0°E, changing water properties considerably (Bakker et al., 2008; Gouretski and Danilov, 1993). In the deep Weddell basin ( $\theta$  of 0.2 to -0.7 °C), the Ba<sub>d</sub> concentration is higher than in the  $\theta$ maximum layer, but it is largely independent of  $\theta$  (with a range of about 5 nmol kg<sup>-1</sup>; Fig. 9). Further Ba<sub>d</sub> enrichment occurs in the bottom layer (see above). This suggests that the Bad level of the deep water is fed by input from above (surface water) and below (water with  $\theta < -0.7$  °C). Within the deep basin, mixing and uplifting of deep water takes place, which relocates dissolved Ba.

#### 5.6. Mass balance of Ba in the deep Weddell Gyre

We construct a tentative mass balance of  $Ba_d$  in the deep Weddell Gyre that must be consistent with the apparent enrichment of  $Ba_d$ within the Gyre compared to the source waters from the ACC. Orsi et al. (1999) estimated the total Antarctic Bottom Water export of the Atlantic sector of the Southern Ocean to be about 11 Sv. This is too high for the Weddell Gyre, because their definition of the Atlantic sector includes large regions east of the gyre. A value of 8–9 Sv may be more representative of the subsurface Weddell Gyre, as reported by Klatt et al. (2005), although also this should be considered as an upper boundary. This outflow must be balanced by an inflow of CDW and surface water from the ACC. The  $Ba_d$  concentration difference between the inflowing CDW with low  $Ba_d$  and outflowing deep water with high  $Ba_d$  amounts to about 5 nmol kg<sup>-1</sup> (see Fig. 9). Taking the water export values of Klatt et al. (2005), we calculate the net Ba loss from the Weddell Gyre to the ACC to be about  $40-45 \text{ mol s}^{-1}$ , or 1.3–  $1.4 \cdot 10^9$  mol year<sup>-1</sup>. Additionally, the deep Weddell Gyre loses water (and Ba<sub>d</sub>) to the surface layer through upwelling and diffusion. Different estimates are being reported for the rate of vertical advection in the Weddell Sea: Hoppema et al. (2001) give a lower end range between 0.5 and 1.3 Sv, while Weppernig et al. (1996), Mensch et al. (1996) and Haine et al. (1998) estimate vertical advection of 1.5-1.7 Sv. Taking a mean deep ocean to Winter Water  $Ba_d$  difference of 15 nmol·kg<sup>-1</sup>, vertical advection would convey 0.6–  $0.8 \cdot 10^9$  mol year<sup>-1</sup> from the deep water to the surface. The total loss from the deep Weddell Gyre thus amounts to  $1.9-2.2 \cdot 10^9$  mol  $Ba_d$  year<sup>-1</sup>. Assuming steady state, this figure must be identical to the supply of Ba<sub>d</sub> to the deep water. The efflux of Ba<sub>d</sub> from the Weddell Sea sediments contributes  $0.4 \cdot 10^9$  mol year<sup>-1</sup> to the deep water (see above). We assume that the remaining  $1.5-1.8 \cdot 10^9$  mol Ba<sub>d</sub> year<sup>-1</sup> is contributed by supply from the surface layer (Fig. 10), where Bad is removed and precipitated and brought to the deep Weddell as particulate matter, where it dissolves in the undersaturated water (Fig. 6).

We computed a net loss of  $Ba_d$  of the Weddell Gyre, but it should be stressed that this is only a loss from the *deep* gyre (Fig. 10). This excess  $Ba_d$  originates predominantly from the surface layer. The surface layer of the Weddell Gyre is eventually generated through upwelling of CDW. It is apparent that the  $Ba_d$  concentration in the surface layer is much lower than that of the CDW (Fig. 3), indicating that  $Ba_d$  is lost from the surface layer (see also above). The Ba depleted surface water is transferred equatorwards from the Weddell Gyre through Ekman transport.

Considering a Weddell surface area of  $1 \cdot 10^{12}$  m<sup>2</sup>, the supply to the deep Weddell Bad reservoir, required to make up the balance, equals 4- $5 \,\mu\text{mol}\,\text{m}^{-2}\,\text{day}^{-1}$ . This is similar, but generally somewhat larger than, the Ba fluxes recorded by deep water sediment traps elsewhere. A trap at 4023 m from site M2 (French KERFIX/ANTARES program 1994–1995) located in the ACC beyond the eastern rim of the Weddell Gyre in the Indian sector (52°S-61°32'E), recorded Ba fluxes between 0.2 and 2.8  $\mu$ mol m<sup>-2</sup> day<sup>-1</sup> (average flux = 0.94  $\mu$ mol m<sup>-2</sup> day<sup>-1</sup>; Jeandel et al., unpublished data). At the M3 site (63°S-71°E; KERFIX/ANTARES program) in the seasonal ice zone further south, Ba fluxes at 1334 m and 3444 m were smaller (range: 0.06 to 0.34  $\mu$ mol m<sup>-2</sup> day<sup>-1</sup>; average =  $0.14 \,\mu\text{mol}\,\text{m}^{-2}\,\text{day}^{-1}$  and range 0.09 to 0.38  $\mu\text{mol}\,\text{m}^{-2}\,\text{day}^{-1}$ ; average =  $0.22 \text{ }\mu\text{mol m}^{-2} \text{ day}^{-1}$ , respectively; Jeandel et al., unpublished data). Sediment traps moored in the Polar Frontal Zone along 145°E, had average Ba fluxes of 0.74 and 0.68  $\mu$ mol m<sup>-2</sup> day<sup>-1</sup> at 800 and 1580 m, respectively (Jacquet et al., 2007a). The particulate Ba removal estimated here is similar to the seasonal input of particulate Ba in the mesopelagic waters (as a result of barite precipitation in decomposing sinking aggregates) of the Polar Front along 6°W  $(1.4 \,\mu\text{mol}\,\text{m}^{-2}\,\text{day}^{-1};\text{ Dehairs et al., 1997})$ , which represents at least



Fig. 9. Dissolved barium plotted against potential temperature. Data are from the Prime Meridian section only; surface layer data are excluded.



**Fig. 10.** Schematic representation of the deep Weddell Gyre highlighting annual fluxes of dissolved Ba (in 10<sup>9</sup> mol year<sup>-1</sup>).

part of the vertical flux of Ba in the water column. Though it seems that the here derived apparent vertical fluxes are on the high side, we also need to consider that the total enrichment of  $Ba_d$  must by nature be larger than the particulate Ba fluxes captured by deep ocean traps, because the material caught in these traps is only part of the total, the rest having been dissolved on the way. Finally,  $Ba_d$  removal in the surface layer via barite precipitation associated with sea-ice formation and biological activity may be more efficient in the Weddell Gyre than in other regions because of the extensive sea-ice formation.

#### 6. Concluding remarks

Two full-depth sections have provided insight into the factors determining the Ba<sub>d</sub> distribution in the Weddell Gyre. The gyre is supplied with deep water from the ACC, which is relatively low in Ba<sub>d</sub>. The low Bad concentration could still be observed at the Prime Meridian (which is some distance away from the actual ACC source). In the gyre, this low-Bad subsurface water is upwelled into the surface layer. Further decrease of Bad in the surface layer occurs due to Ba uptake as a consequence of phytoplankton activity, possibly complemented by barite formation in sea-ice. Within the voluminous deep Weddell Gyre, Bad is enriched compared to the source water of the gyre. This is effected by barite dissolution in the undersaturated deep water (with barite originating from the depleted surface layer); and also by Bad mixing into the deep water from the strongly Bad enriched bottom layer. Bad influx from the bottom layer seems to occur predominantly in the southeastern part of the Weddell basin, in exactly the same region where bottom layer enrichment of silicate occurs. Water movement and circulation further redistribute Bad through the Weddell-Enderby basin. Tentative calculations show that enrichment of the deep water due to Bad originating from the surface layer is likely to be larger than through Ba<sub>d</sub> supply from the bottom layer.

Summarizing, a mechanism appears to exist whereby the Weddell Gyre acts as an efficient conduit for  $Ba_d$  export to the deep and abyssal world oceans. In the surface layer,  $Ba_d$  is depleted due to phytoplankton activity and, possibly, barite precipitation within sea-ice. This particulate, absorbed and adsorbed Ba is subsequently exported from the surface layer by settling and redissolves in the deep Weddell Gyre. The deep water, classified as Antarctic Bottom Water, thus enriched in  $Ba_d$ , abandons the gyre to replenish the abyssal ocean basins to the north.  $Ba_d$  depleted surface water leaves the gyre northwards through Ekman transport.

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#### Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at doi:10.1016/j.marchem.2010.07.005.

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