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Bromoform concentrations in slush-layer water in Antarctic fast ice

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Abstract:

Bromoform concentrations in water of the slush layer that developed at the interface between snow and sea ice were measured during the seasonal warming in Lützow-Holm Bay, Eastern Antarctica. Mean bromoform concentration was $5.5 \pm 2.4 \text{ pmol l}^{-1}$, which was lower than that of the under-ice water ($10.9 \pm 3.5 \text{ pmol l}^{-1}$). Temporal decrease in bromoform concentrations and salinity with increasing in temperature of the slush water suggest that the bromoform concentrations were reduced through dilution with meltwater input from the upper surface of sea ice. In contrast, bromoform concentrations in the under-ice water increased during this period while the salinity of the under-ice water decreased. It is speculated that the sea ice meltwater input contained high bromoform concentrations from the brine channels within the sea ice and from the bottom of the ice that were contributed to the increased bromoform concentrations in the under-ice water.

Key words: bromoform, sea ice, slush layer, under-ice water, Lützow-Holm Bay

Introduction

Bromoform (CHBr_3) is one of the volatile organic compounds (VOCs) originating with macroalgal and planktonic sources in the ocean, and bromoform is emitted from the ocean surface to the atmosphere (Quack & Wallace 2003). Bromoform is an important carrier of bromine to the troposphere, and its lifetime in the atmosphere is one to four weeks (Quack & Wallace 2003 and references therein). Although bromoform concentrations in the ocean have been measured widely, there is a lack of information regarding measurements of bromoform in the ice-covered seas. Sea ice may be a significant platform for the production of bromine compounds and a source for the atmosphere (Sturges *et al.* 1992, 1993, 1997, Carpenter *et al.* 2007). For example, polar ice algae is known to produce significant quantities of bromoform (Sturges *et al.* 1992).

A surface slush layer (gap layer) is found extensively in Antarctic sea ice during the ice melting season (Haas *et al.* 2001, Kattner *et al.* 2004, Ackley *et al.* 2008, Zemmelen *et al.* 2008, Papadimitriou *et al.* 2009). Snow accumulation over sea ice and the formation of superimposed ice leads to the formation of a slush layer below sea level (Haas *et al.* 2001). High biological activity has been found in this layer (Kattner *et al.* 2004), suggesting that bromoform production could also be high in this layer. In addition, bromoform produced in the slush layer would accumulate because of limited gas diffusion through the snow and superimposed ice (Delille 2006, Nomura *et al.* 2010).

The bromoform dynamics in the sea ice column has been well studied (Sturges *et al.* 1993, 1997). From these studies, it is clear that the biogenic production of bromoform by ice algae could be an important contributor to atmospheric bromoform levels in the polar regions (e.g., Sturges *et al.* 1992). However, although slush water is expected to potentially include the highest levels of bromoform in surface ice, bromoform concentrations in slush water have not yet been examined. In addition, it is not clear which processes dominate the temporal variability of bromoform concentrations in slush water.

In this study, we examined for the first time the magnitude and temporal variations of bromoform concentrations in slush water, and their links to the physical and biogeochemical parameters of slush water and sea ice. Our results should provide an important insight into the organic bromine cycle in ice-covered seas.

Materials and methods

Slush-water samples were collected in the austral spring/summer (26 December 2009 to 5 February 2010) from the multi-year land-fast ice in Lützow-Holm Bay, Eastern Antarctica (Fig. 1). During this period, several sampling stations were established within a 0.5-km² area. Under-ice water was also collected from the same area.

For slush-water sampling, snow and superimposed ice were removed with a scoop, and ice crystals in the slush layer were removed with a net. The water in the slush layer was pumped through a Teflon tube by a diaphragm pump (EWP-01, As One Corporation, Osaka, Japan) and collected in a 30-ml amber glass vial (Nichiden-Rika Glass Co. Ltd, Kobe, Japan). Mercuric chloride solution (Saturated-HgCl₂; 200 µl) was added to the samples to stop biological activity. Vials were sealed with a Teflon-lined septum (Nichiden-Rika Glass Co. Ltd) and an aluminum cap (Nichiden-Rika Glass Co. Ltd). Slush-water samples were stored in a refrigerator (+4 °C) until further analysis. To check the spatial heterogeneity of slush water, duplicate samplings were carried out for slush water within 2 m² on each sampling day.

For under-ice water sampling, a hole was made using an ice corer (Geo Tecs Co., Ltd., Chiba, Japan) with an internal diameter of 9.0 cm, and under-ice water was collected with a Teflon water sampler (GL Science Inc., Tokyo, Japan) at 1 and 7 m below the bottom of the sea ice. Under-ice water samples were collected approximately 30 minutes after drilling of the ice cores to avoid the effects of the disturbance caused by drilling. Samples were treated and stored in the same manner as for slush-water samples.

The temperature of the slush and under-ice water was measured using a needle-type temperature sensor (Testo 110 NTC, Brandt Instruments, Inc., Prairieville, LA, USA). Slush and under-ice water samples were also collected and placed into 12-ml glass screw-cap vials (Nichiden-Rika Glass Co. Ltd) for salinity analysis, and into 500-ml Nalgene polycarbonate bottles (Thermo Fisher Scientific Inc., Waltham, MA, USA) for chlorophyll *a* (chl-*a*) measurement. Salinity samples were kept in a refrigerator (+4 °C). The seawater samples for chl-*a* analysis were filtered through 25-mm Whatman GF/F filters immediately after returning to the laboratory near the sampling station. Chlorophyll pigments on the filters were extracted in dimethylformamide (Suzuki & Ishimaru 1990) for 24 h at approximately –80 °C.

The bromoform concentrations were determined by a purge-and-trap (P&T) and gas-chromatograph–mass spectrometry (GC-MS) method (Yokouchi *et al.* 2006, Ooki & Yokouchi 2011a, Ooki & Yokouchi 2011b). The total volume of seawater in the sample bottle was transferred to a custom-made bubbling vessel by helium carrier at 40 ml min^{–1}. The dissolved bromoform in the water was purged with the helium carrier at 80 ml min^{–1} and 45 °C for 30 min, and simultaneously transferred to a pre-concentration-GC-MS system (Agilent 5973, 6390; Agilent Technologies Inc., Santa Clara, CA, USA) (Yokouchi *et al.* 2006, Ooki & Yokouchi 2011a, Ooki & Yokouchi 2011b). The purge efficiency for bromoform was 84%. A diluted standard solution containing bromoform at 1.2 pmol l^{–1} was introduced to the P&T-GC-MS system every 24 h to calibrate for the bromoform concentrations in seawater samples. The precision of standard solution measurements was ±2% (*n* = 4), and the detection limit (S/N = 10) was 0.4 pmol l^{–1}.

Salinity was measured with a salinity analyzer (SAT-210, Toa Electronics Ltd., Tokyo, Japan). The salinity analyzer was calibrated with International Association for the Physical Science of the Ocean standard seawater (P series; Ocean Scientific International Ltd., Hampshire, UK). Chl-*a* concentrations were determined using a fluorometer (Model 10AU, Turner Designs, Inc., Sunnyvale, CA, USA), following methods described by Parsons *et al.* (1984).

Results

Bromoform and chl-a concentrations, salinity and temperature

Bromoform and chl-*a* concentrations, salinity and temperature in slush and under-ice water obtained during the study period are summarized in Table 1. The mean bromoform concentration in slush water was 5.5 pmol l⁻¹, which was lower than that of under-ice water (10.9 pmol l⁻¹). In contrast, the mean chl-*a* concentration in slush water (0.4 µg l⁻¹) was approximately double that of under-ice water (0.2 µg l⁻¹). The mean salinity of the slush water (9.7) was approximately one-third that of the under-ice water (32.5). The mean temperature of the slush water (+0.5 °C) was higher than that of the under-ice water (-0.4 °C) due to the downward flushing of meltwater from the top, which was generated by the melting of the snow on sea ice at higher air temperature (Fig. 2). Although the range of values for some of these parameters includes temporal variations during the study period, the slush water was consistently characterized by lower bromoform concentrations and lower salinity compared to the under-ice water (Table 1).

Temporal variations in bromoform and chl-a concentrations, salinity and temperature

Sampling occurred from late spring (26 December) to mid summer (5 February), providing an opportunity to observe temporal variations of physico-chemical properties during warming and ice-melting conditions (Fig. 2). The snow depth over the sea ice decreased from 100 cm at the beginning of this period to less than 10 cm at the end, whereas ice thickness was almost constant during the period (176 ± 25 cm). There was a layer of superimposed ice (about 10 cm thick) between the snow and slush layers during the sampling period.

Temporal variations in bromoform and chl-*a* concentration, salinity and temperature in slush and under-ice water were shown in Figure 3. Bromoform concentrations in the slush water were relatively constant (8–9 pmol l⁻¹) until 5 January, and then they decreased to below 5 pmol l⁻¹ and remained constant for the rest of the sampling period (Fig. 3a). The salinity of the slush water decreased dramatically from 28.6 on 26 December to 0.8 on 2 February (Fig. 3e). On the other hand, the slush-water temperature increased from -1.9 °C to +2.6 °C during the study period (Fig. 3g). The trends of decrease in salinity and increase in temperature of slush water were similar to that of bromoform concentrations during this period (Figs. 3a, e & g), with the bromoform concentrations being correlated with the salinity (shown in Figure 4), and with the temperature of slush water ($r = 0.78$, $P < 0.0001$, $n = 22$). For the relationship between salinity and bromoform concentrations in slush water (Fig. 4), there were two different regimes: 1) the decrease in bromoform concentrations in the slush water only holds until a decrease in salinity to around 5, 2) at lower salinity (salinity < 5), the bromoform concentration in slush water increased to 5.9 pmol l⁻¹ with decreasing salinity. Chl-*a* concentrations in slush water varied between 0.0 and 1.1 µg l⁻¹ (Fig. 3c), and were not correlated with the changes in bromoform concentrations ($r = 0.09$, $P = 0.68$, $n = 22$).

Concentrations of bromoform in under-ice water (5–18 pmol l⁻¹) were higher compared with the slush-water concentrations (Figs. 3a & b). Chl-*a* concentrations in under-ice water remained low (<0.3 pmol l⁻¹) until 14 January, and then they gradually increased (Fig. 3d). Salinity of under-ice water decreased from about 34 early in the study period (26 December–5 January) to below 30 at the end (29 January–5 February) (Fig. 3f). The temperature of under-ice water increased slightly during the study period (Fig. 3h). There were no notable differences between the under-ice water at 1 m and 7 m for any of the parameters during the study period (Figs. 3b, d, f & h).

Discussion

The bromoform concentrations in under-ice water measured in this study (5.9–18.3 pmol l⁻¹) were lower than those of Arctic under-ice water in spring (about 80 pmol l⁻¹; Sturges *et al.* 1997) and coastal Antarctic surface water in spring/summer (about 57 pmol l⁻¹; Carpenter *et al.* 2007). The higher bromoform concentrations previously measured in polar seawater were caused by inputs of ice meltwater containing high levels of bromoform produced by ice algae, or by *in situ* production of bromoform in the seawater during algae blooms (e.g., Sturges *et al.* 1992, Carpenter *et al.* 2007). The increasing levels of bromoform in the ocean have generally been associated with the increasing abundance of diatoms (Klick & Abrahamsson 1992, Baker *et al.* 1999). An *in situ* culture experiment has shown that Arctic ice algae have the potential to produce substantial quantities of bromoform at high chl-*a* concentrations (>700 µg l⁻¹) within the bottom layer of sea ice (Sturges *et al.* 1992). However, because chl-*a* concentrations were generally low during our study (mean, 0.2 µg l⁻¹), the contribution of ice algae to bromoform production would have been minor. This is likely to be one of reasons why bromoform concentration in under-ice water in this study were so much lower than those of previous studies (Sturges *et al.* 1992, Carpenter *et al.* 2007).

A surface slush layer is found widely distributed in Antarctic sea ice during the ice melting season (Haas *et al.* 2001, Kattner *et al.* 2004, Ackley *et al.* 2008, Zemmeling *et al.* 2008, Papadimitriou *et al.* 2009). Snow accumulation over sea ice and the formation of superimposed ice leads to the formation of a slush layer below sea level (Haas *et al.* 2001). In this layer, high biological activity has been found (Kattner *et al.* 2004), suggesting that the biogenic production of bromoform should also be high in this layer. However, in this study we found chl-*a* concentrations in the slush layer (mean 0.4 µg l⁻¹; Table 1) to be much lower than those in the productive slush layer in the Weddell Sea, Antarctica (3.1–16.5 µg l⁻¹; Kattner *et al.* 2004). Snow depth was basically high (mean 24 cm) compared to those in the Weddell Sea, Antarctica (mean 16 cm; Kattner *et al.* 2004). These results suggest the reduction of the light intensity in the slush layer, thereby reducing the light available for growth of ice algae living in slush water. In addition, the salinity of slush water was lower than under-ice water, reflecting the dilution of all slush-water components including nutrients. Therefore, it was considered

that available nutrient concentration in the slush water was also low and depleted for growth of ice algae. Although measures of the light intensity and nutrient concentrations in slush layer were not examined in this study, these may be one of the factors controlling biological productivity in slush water.

The temporal decrease of bromoform concentrations in slush water was correlated with that of salinity when the salinity was higher than 5 (Fig. 4), suggesting that bromoform concentrations in the slush water decreased because of dilution by the meltwater input from the upper surface of sea ice in accordance with the increase of temperature (Figs. 2 & 3g). Therefore, the decrease in bromoform concentrations in slush water should closely reflect the effects of dilution. Same dilution process was observed for hexachlorocyclohexane in sea ice brine in the Canadian western Arctic in spring due to the melting of the ice crystal matrix and replenishment of brine with seawater (Pucko *et al.* 2010). On the other hand, at lower salinity (salinity < 5), the bromoform concentrations in the slush water increased with decreasing salinity (Fig. 4). It is speculated that the bromoform in the slush layer tended to remain even if the large volume of meltwater was added to the slush layer and diluted the bromoform because of the limited gas diffusion to the atmosphere through the snow and superimposed ice (Delille 2006, Nomura *et al.* 2010).

In contrast to the changes in slush water, bromoform concentrations in under-ice water increased during the study period, tracking the increase in chl-*a* concentrations, whereas the salinity decreased (Figs. 3b, d & f). These results for salinity suggest that the under-ice water included a proportion of meltwater from the sea ice. This phenomenon is similar to the process occurring in the slush water, but there are some differences in the effects on bromoform and chl-*a* concentrations. Previously, bromoform and chl-*a* concentration at the bottom of the sea ice have been found to be higher than in the other parts of sea ice (Sturges *et al.* 1997). During ice melting, the impurities in sea-ice brine channels are flushed out before the melting of the ice itself, therefore these components were added to the under-ice water with meltwater. The temporal changes of each component in under-ice water reflect the input of the high-bromoform and chl-*a*, low-

salinity meltwater during the study period (Figs. 3b, d & f). These changes would be also enhanced by the horizontal advection of under-ice water from offshore that accumulates in Lützow-Holm Bay (Ohshima et al., 1996).

Conclusions

Bromoform concentrations in slush-layer water in Antarctic fast ice were measured during the seasonal warming in Lützow-Holm Bay. Mean bromoform concentration was $5.5 \pm 2.4 \text{ pmol l}^{-1}$, which was lower than that of the under-ice water ($10.9 \pm 3.5 \text{ pmol l}^{-1}$). Temporal decrease in bromoform concentrations and salinity with increasing in temperature of the slush water, suggesting that bromoform concentrations in the slush water decreased because of dilution by the meltwater input from the upper surface of sea ice in accordance with the increase of temperature. However, at lower salinity of slush water (salinity < 5), the bromoform concentrations in the slush water increased with decreasing salinity. It is speculated that the bromoform in the slush layer tended to remain even if the large volume of meltwater were added to the slush layer and diluted bromoform in slush water because of the limited gas diffusion to the atmosphere through the snow and superimposed ice (Delille 2006, Nomura *et al.* 2010).

In contrast to the changes in slush water, bromoform concentrations in under-ice water increased during the study period, tracking the increase in chl-*a* concentrations, whereas the salinity decreased. These results suggest that the temporal changes of each component in under-ice water reflect the input of the high-bromoform and chl-*a*, low-salinity meltwater during the study period.

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296 MS).
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References

- ACKLEY, S.F., LEWIS, M.J., FRITSEN, C.H. & XIE, H. 2008. Internal melting in Antarctic sea ice: Development of “gap layers”. *Geophysical Research Letters*, 35, L11503, doi:10.1029/2008GL033644.
- BAKER, J.M., REEVES, C.E., NIGHTINGALE, P.D., PENKETT, S.A., GIBB, S.W. & HATTON, A.D. 1999. Biological production of methyl bromide in the coastal waters of the North Sea and open ocean of the northeast Atlantic. *Marine Chemistry*, 64, 267–285.
- CARPENTER, L.J., WEVILL, D.J., PALMER, C.J. & MICHELS, J. 2007. Depth profiles of volatile iodine and bromine-containing halocarbons in coastal Antarctic waters, *Marine Chemistry*, 103, 227–236.
- DELILLE, B. 2006. Inorganic carbon dynamics and air-ice-sea CO₂ fluxes in the open and coastal waters of the Southern Ocean. *Ph. D. thesis, Univ. of Liège, Liège, Belgium*, 1–297.
- HAAS, C., THOMAS, D.N. & BAREISS, J. 2001. Surface properties and processes of perennial Antarctic sea ice in summer. *Journal of Glaciology*, 47, 613–625.
- KATTNER, G., THOMAS, D.N., HAAS, C., KENNEDY, H. & DIECKMANN, G.S. 2004. Surface ice and gap layers in Antarctic sea ice: highly productive habitats. *Marine Ecology Progress Series*, 277, 1–12.
- KLICK, S. & ABRAHAMSSON, K. 1992. Biogenic volatile iodated hydrocarbons in the ocean. *Journal of Geophysical Research*, 97(C8), 12683–12687.
- NOMURA, D., INOUE, Y.H., TOYOTA, T. & SHIRASAWA, K. 2010. Effects of snow, snow-melting and re-freezing on air-sea ice CO₂ flux. *Journal of Glaciology*, 56, 262–270.

- OHSHIMA, K.I., TAKIZAWA, T., USHIO, S. & KAWAMURA, T. 1996. Seasonal variations of the Antarctic coastal ocean in the vicinity of Lützow-Holm Bay. *Journal of Geophysical Research*, 101, 20617–20628.
- OOKI, A. & YOKOUCHI, Y. 2011a. Determination of Henry's law constant of halocarbons in seawater and analysis of sea-to-air flux of iodoethane (C₂H₅I) in the Indian and Southern oceans based on partial pressure measurements. *Geochemical Journal*, 45, e1–e7.
- OOKI, A. & YOKOUCHI, Y. 2011b. Dichloromethane in the Indian Ocean: Evidence for in-situ production in seawater. *Marine Chemistry*, doi:10.1016/j.marchem.2011.01.001, In press.
- PAPADIMITRIOU, S., THOMAS, D.N., KENNEDY, H., KUOSA, H. & DIECKMANN, G.S. 2009. Inorganic carbon removal and isotopic enrichment in Antarctic sea ice gap layers during early austral summer. *Marine Ecology Progress Series*, 386, 15–27.
- PARSONS, T.R., TAKAHASHI, M. & HARGRAVE, B. 1984. *Biological oceanographic processes*, 3rd ed. 330 pp. Oxford, Pergamon Press.
- PUCKO, M., STERN, G.A., MACDONALD, R.W. & BARBER, D.G. 2010. α - and γ -Hexachlorocyclohexane measurements in the brine fraction of sea ice in the Canadian High Arctic using a sump-hole technique. *Environmental Science & Technology*, 44, 9258–9264.
- QUACK, B. & WALLACE, D.W.R. 2003. Air-sea flux of bromoform: Controls, rates, and implications. *Global Biogeochemical Cycles*, 17, 1023, doi:10.1029/2002GB001890.
- STURGES, W.T., COTA, G.F. & BUCKLEY, P.T. 1992. Bromoform emission from Arctic ice algae. *Nature*, 358, 660–662.

- STURGES, W.T., SULLIVAN, C.V., SCHNELL, R.C., HEIDT, L.E. & POLLACK, W.H. 1993. Bromoalkane production by Antarctic ice algae. *Tellus*, 45B, 120–126.
- STURGES, W.T., COTA, G.F. & BUCKLEY, P.T. 1997. Vertical profiles of bromoform in snow, sea ice, and seawater in the Canadian Arctic. *Journal of Geophysical Research*, 102(C11), 25073–25083.
- SUZUKI, R., & ISHIMARU, T. 1990. An improved method for the determination of phytoplankton chlorophyll using N,N-dimethylformamide. *Journal of the Oceanographical Society of Japan*, 46, 190–194.
- YOKOUCHI, Y., TAGUCHI, S., SAITO, T., TOHJIMA, Y., TANIMOTO, H. & MUKAI, H. 2006. High frequency measurements of HFCs at a remote site in east Asia and their implications for Chinese emissions. *Geophysical Research Letters*, 33, L21814, doi:10.1029/2006GL026403.
- ZEMMELINK, H.J., HINTSA, E.J., HOUGHTON, L., DACEY, J.W.H. & LISS, P.S. 2008. DMS fluxes over the multi year ice of the western Weddell Sea. *Geophysical Research Letters*, 35, L06603, doi:10.1029/2007GL031847.

Figure captions

Fig 1 Map showing the location of sampling area in Lützow-Holm Bay, Eastern Antarctica.

Fig 2 Time series of daily mean air temperatures at the sampling station. Gray band indicates the sampling period.

Fig 3 Time series values for (a, b) bromoform concentration, (c, d) chl-*a* concentration, (e, f) salinity and (g, h) temperature, for slush water (a, c, e & g) and under-ice water (b, d, f & h). Error bars for slush water data (a, c, e & g) indicate the range of duplicate measurements. Error bars are only shown if they extend beyond the symbols.

Fig 4 Relationships between salinity and bromoform concentrations in slush water. Solid and dashed line represents the linear regression line. Bromoform concentrations in slush water were correlated with the salinity in slush water for salinity > 5 ($r = 0.93$, $P < 0.0001$, $n = 12$) and for salinity < 5 ($r = 0.64$, $P < 0.046$, $n = 10$).

Table captions

Table 1. Mean, minimum, Q1 (25 percentile), median, Q3 (75 percentile) and maximum of bromoform and chl-*a* concentrations, and salinity and temperature for slush water (n=22) and under-ice water (n=30 to 32) as measured during the study period.

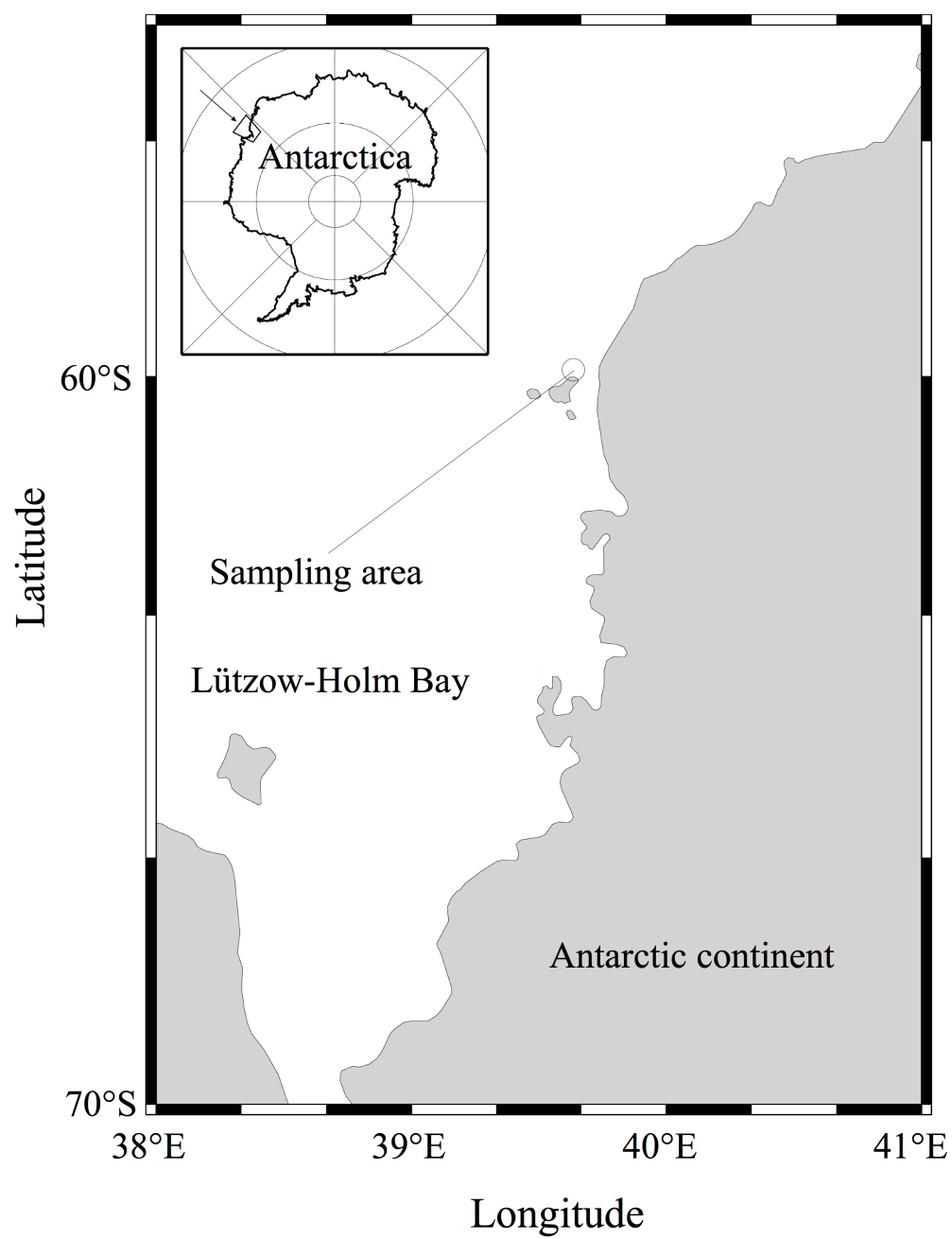


Figure 1. Nomura et al.

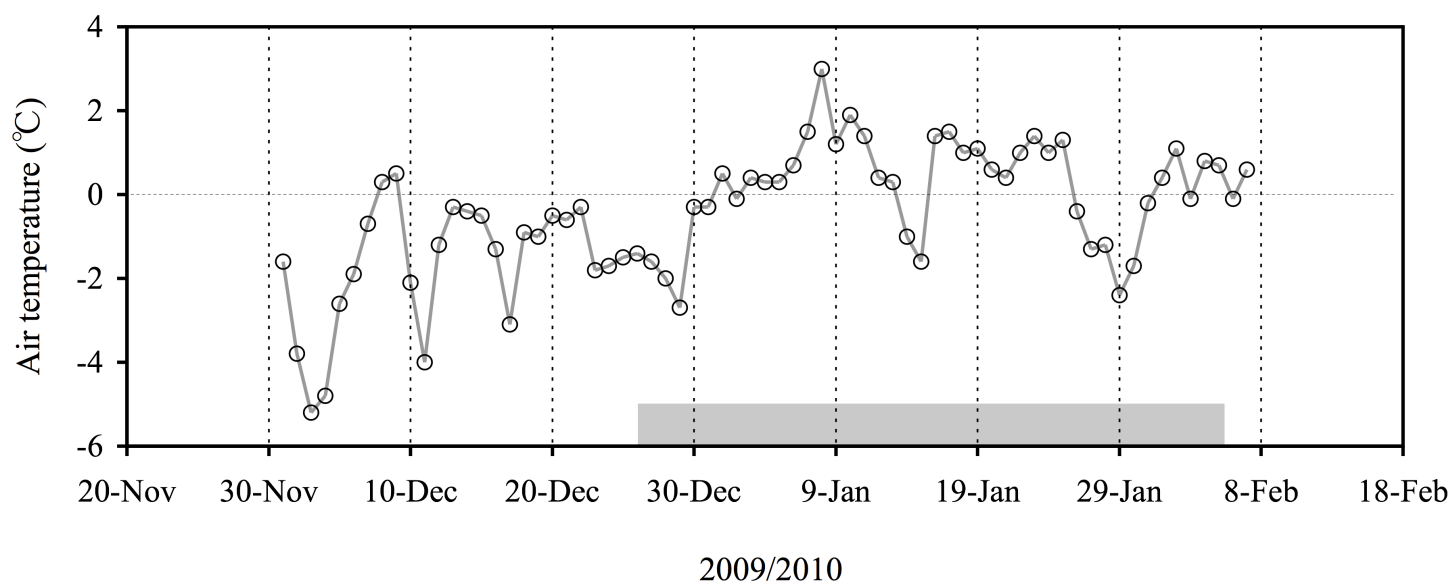


Figure 2. Nomura et al.

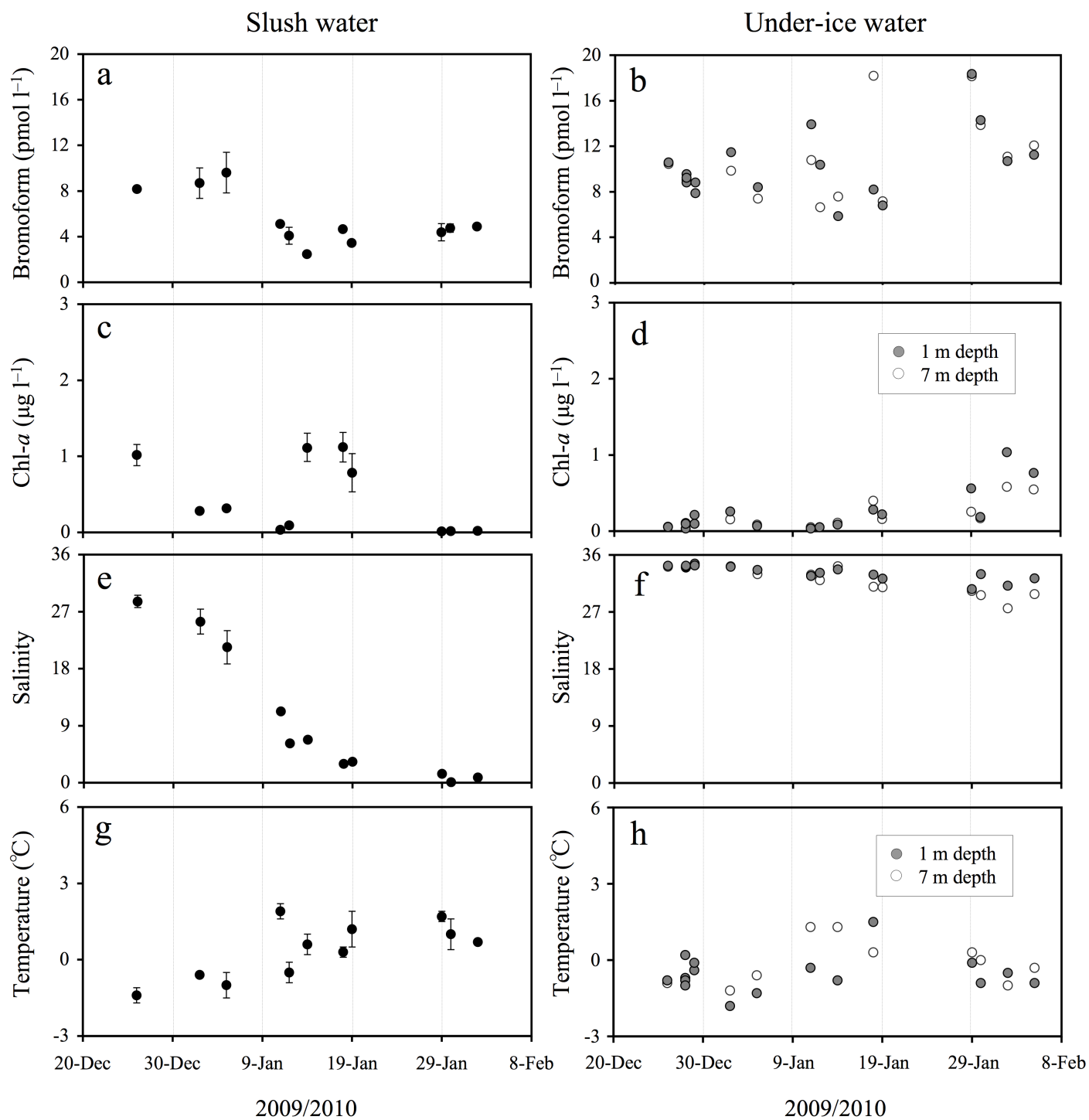


Figure 3. Nomura et al.

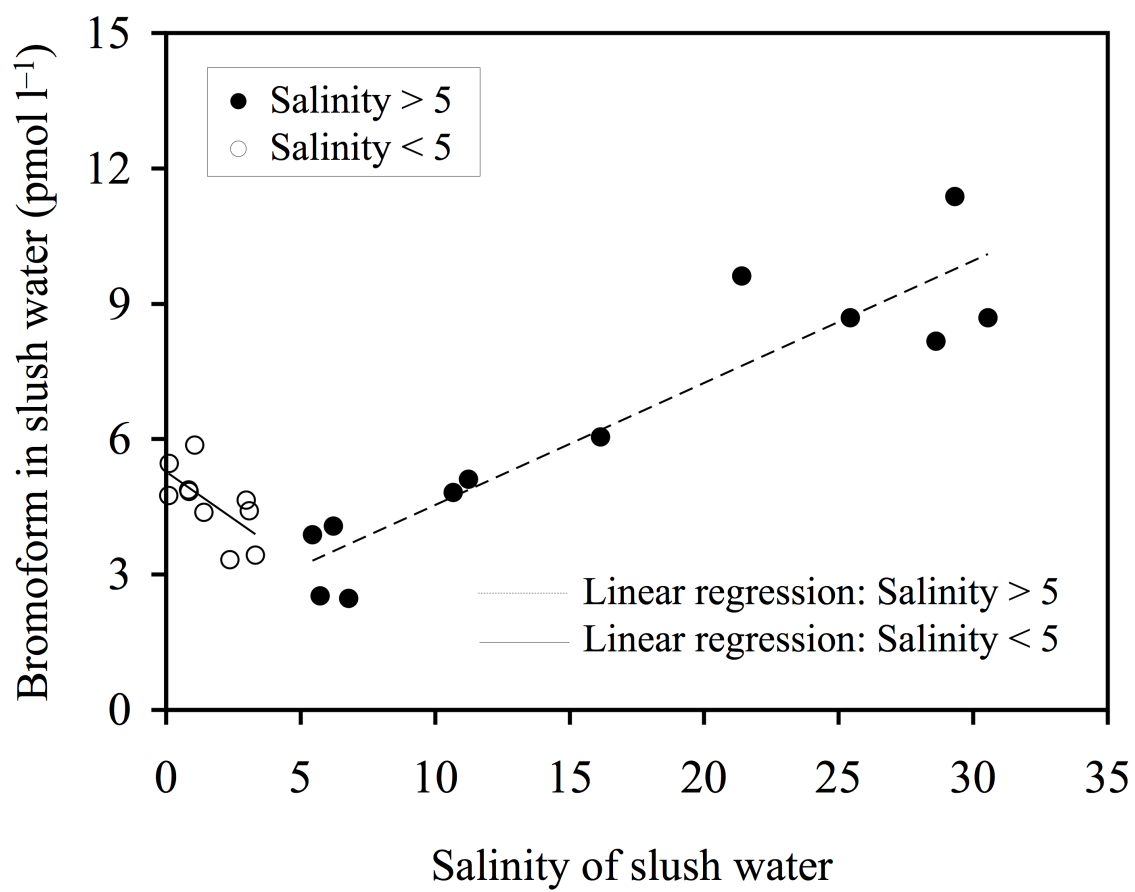


Figure 4. Nomura et al.

Table 1. Mean, minimum, Q1 (25 percentile), median, Q3 (75 percentile) and maximum of bromoform and chl-*a* concentrations, and salinity and temperature for slush water (n=22) and under-ice water (n=30 to 32) as measured during the study period.

Sample	Parameter	Mean	Minimum	Q1	Median	Q3	Maximum
Slush water	Bromoform (pmol l ⁻¹)	5.5	2.5	4.2	4.8	6.0	11.4
	Chl- <i>a</i> (µg l ⁻¹)	0.4	0.0	0.0	0.2	0.7	1.1
	Salinity	9.7	0.1	1.6	5.6	14.9	30.6
	Temperature (°C)	▼ +0.5	-1.9	-0.4	▼ +0.6	▼ +1.3	▼ +2.6
Under-ice water	Bromoform (pmol l ⁻¹)	10.9	5.9	8.4	10.4	12.5	18.3
	Chl- <i>a</i> (µg l ⁻¹)	0.2	0.0	0.1	0.1	0.3	1.0
	Salinity	32.5	27.6	31.1	32.9	34.1	34.7
	Temperature (°C)	-0.4	-1.8	-0.9	-0.7	-0.1	▼ +1.5