



Estimates of ikaite export from sea ice to the underlying seawater in a sea ice–seawater mesocosm

Nicolas-Xavier Geilfus^{1,2}, Ryan J. Galley¹, Brent G. T. Else³, Karley Campbell¹, Tim Papakyriakou¹, Odile Crabeck¹, Marcos Lemes¹, Bruno Delille⁴, and Søren Rysgaard^{1,2,5}

¹Centre for Earth Observation Science, University of Manitoba, Winnipeg, Canada

²Arctic Research Centre, Aarhus University, Aarhus, Denmark

³Department of Geography, University of Calgary, Calgary, Canada

⁴Unité d'Océanographie Chimique, Université de Liège, Liège, Belgium

⁵Greenland Climate Research Centre, Greenland Institute of Natural Resources, Nuuk, Greenland

Correspondence to: Nicolas-Xavier Geilfus (geilfusn@umanitoba.ca)

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Abstract. The precipitation of ikaite and its fate within sea ice is still poorly understood. We quantify temporal inorganic carbon dynamics in sea ice from initial formation to its melt in a sea ice–seawater mesocosm pool from 11 to 29 January 2013. Based on measurements of total alkalinity (TA) and total dissolved inorganic carbon (TCO₂), the main processes affecting inorganic carbon dynamics within sea ice were ikaite precipitation and CO₂ exchange with the atmosphere. In the underlying seawater, the dissolution of ikaite was the main process affecting inorganic carbon dynamics. Sea ice acted as an active layer, releasing CO₂ to the atmosphere during the growth phase, taking up CO₂ as it melted and exporting both ikaite and TCO₂ into the underlying seawater during the whole experiment. Ikaite precipitation of up to 167 μmol kg^{−1} within sea ice was estimated, while its export and dissolution into the underlying seawater was responsible for a TA increase of 64–66 μmol kg^{−1} in the water column. The export of TCO₂ from sea ice to the water column increased the underlying seawater TCO₂ by 43.5 μmol kg^{−1}, suggesting that almost all of the TCO₂ that left the sea ice was exported to the underlying seawater. The export of ikaite from the ice to the underlying seawater was associated with brine rejection during sea ice growth, increased vertical connectivity in sea ice due to the upward percolation of seawater and meltwater flushing during sea ice melt. Based on the change in TA in the water column around the onset of sea ice melt, more than half of the total ikaite precipitated in the ice during sea ice growth was still contained in the ice when the

sea ice began to melt. Ikaite crystal dissolution in the water column kept the seawater *p*CO₂ undersaturated with respect to the atmosphere in spite of increased salinity, TA and TCO₂ associated with sea ice growth. Results indicate that ikaite export from sea ice and its dissolution in the underlying seawater can potentially hamper the effect of oceanic acidification on the aragonite saturation state ($\Omega_{\text{aragonite}}$) in fall and in winter in ice-covered areas, at the time when $\Omega_{\text{aragonite}}$ is smallest.

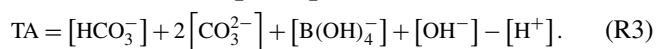
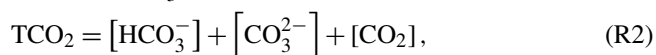
1 Introduction

Currently, each year, 7 Pg anthropogenic carbon is released to the atmosphere, 29 % of which is estimated to be taken up by the oceans through physical, chemical and biological processes (Sabine et al., 2004). The Arctic Ocean takes up −66 to −199 Tg C year^{−1}, (where a negative value indicates an uptake of atmospheric CO₂) contributing 5–14 % to the global ocean CO₂ uptake (Bates and Mathis, 2009), primarily through primary production and surface cooling (MacGilchrist et al., 2014). However, polar ocean CO₂ uptake estimates consider sea ice as an impermeable barrier, ignoring the potential role of ice-covered areas on gas exchange between the ocean and atmosphere. Recent studies have shown that sea-ice-covered areas participate in the variable sequestration of atmospheric CO₂ into the mixed layer below the ice (e.g., Papakyriakou and Miller, 2011; Geil-

fus et al., 2012, 2014, 2015; Nomura et al., 2013; Delille et al., 2014). Studies are required to elucidate the processes responsible for this as well as their magnitudes, both temporally and spatially.

The carbonate chemistry in sea ice and brine is spatially and temporally variable, which leads to complex CO₂ dynamics with the potential to affect the air–sea CO₂ flux (Parmentier et al., 2013). Release of CO₂ from sea ice to the atmosphere has been reported during sea ice formation from open water (Geilfus et al., 2013a) and in winter (Miller et al., 2011; Fransson et al., 2013), while uptake of CO₂ by sea ice from the atmosphere has been reported after sea ice melt onset (e.g., Semiletov et al., 2004; Nomura et al., 2010, 2013; Geilfus et al., 2012, 2014, 2015; Fransson et al., 2013). In combination, these works suggest that the temporal cycle of sea ice formation and melt affects atmospheric CO₂ uptake by the ocean in variable ways. Sea ice may also act as an important control on the partial pressure of CO₂ (*p*CO₂) in the sea surface through a sea ice pump (Rysgaard et al., 2007). During the earliest stages of sea ice formation, a small fraction of CO₂-supersaturated brine is expelled upward onto the ice surface, promoting a release of CO₂ to the atmosphere (Geilfus et al., 2013a). As sea ice forms and grows thicker, salts are partly rejected from the sea ice to the underlying seawater and partly trapped within the sea ice structure, concentrated in brine pockets, tubes and channels. As a result, the concentration of dissolved salts, including inorganic carbon, increases within the brine and promotes the precipitation of calcium carbonate crystals such as ikaite (CaCO₃ · 6H₂O) (Marion, 2001). These crystals have been reported in both natural (Dieckmann et al., 2008; Nomura et al., 2013; Sogaard et al., 2013) and experimental sea ice (Geilfus et al., 2013b; Rysgaard et al., 2014) and have been suggested to be a key component of the carbonate system (Rysgaard et al., 2007; Fransson et al., 2013; Delille et al., 2014).

During ikaite precipitation within sea ice, total alkalinity (TA) in brine is reduced by 2 moles due to the reduction of bicarbonate (HCO₃[−]) while total dissolved inorganic carbon (TCO₂) in brine is only reduced by 1 mol (Reactions R1–R3):



The specific conditions leading to ikaite precipitation, as well as the fate of these precipitates in sea ice, are still not fully understood. Ikaite crystals may remain within the ice structure, while the CO₂ formed during their precipitation is likely rejected with dense brine to the underlying seawater and sequestered below the mixed layer. During sea ice melt, the dissolution of these crystals triggered by increased ice temperatures and decreased bulk ice salinity will consume CO₂

and drive a CO₂ uptake from the atmosphere to the ice. Such a mechanism could be an effective sea ice pump of atmospheric CO₂ (Delille et al., 2014). In addition, ikaite stored in the ice matrix could become a source of TA to the near-surface ocean upon its subsequent dissolution during sea ice melt (Rysgaard et al., 2007, 2009).

The main air–sea fluxes of CO₂ and TCO₂ are driven by brine rejection to the underlying seawater and its contribution to intermediate and deep-water formation (Semiletov et al., 2004; Rysgaard et al., 2007, 2009; Fransson et al., 2013) or below sea ice in ice tank studies (e.g., Killawee et al., 1998; Papadimitriou et al., 2004). As sea ice thickens, reduced near-surface ice temperatures result in reduced brine volume content, increased brine salinity and increased solute concentration in the brine. In the spring–summer, as the ice temperature increases, sea ice brine volume increases and sea ice becomes vertically permeable to liquid (Golden et al., 2007), enhancing the potential CO₂ exchange between the atmosphere, sea ice and ocean. Eventually internal ice melt promotes brine dilution, which decreases brine salinity, TA and TCO₂, and leads to lower *p*CO₂ in the brine. In addition, the dissolution of ikaite decreases brine *p*CO₂ (Reaction R1) (Geilfus et al., 2012, 2015). These conditions all favor sea ice as a sink for atmospheric CO₂ (Nomura et al., 2010, 2013; Geilfus et al., 2012, 2015). Melting sea ice stratifies surface seawater, leading to decreased TA, TCO₂ and *p*CO₂ in the sea surface, enhancing air–sea CO₂ fluxes (Rysgaard et al., 2007, 2009).

Although we now have a basic understanding of the key mechanisms of carbon cycling in sea ice, significant unknowns remain. One of the major unknowns is the fate of ikaite, TCO₂ and CO₂ released from sea ice during winter. It is unclear what proportion of precipitated ikaite crystals in sea ice remains in the matrix to be released upon melt or what proportion is expelled with brine drainage during ice formation and growth. Examining the chemical signatures of the water column beneath sea ice may provide an indication of the importance of the different processes. However, the signal of carbon components released from 1 to 2 m of sea ice growth is difficult to detect in a water column several hundred meters deep.

In this study, we followed the evolution of the inorganic carbon dynamics within experimental sea ice from sea ice formation to melt in a sea ice–seawater mesocosm pool (~435 m³) at the University of Manitoba, Winnipeg, Canada. The benefits of this type of environment are multiple. An artificial pool equipped with a movable bridge makes it possible to collect undisturbed samples from thin growing sea ice. We gain the ability to carefully track carbonate parameters in the ice, in the atmosphere and in the underlying seawater, while growing sea ice in a large volume of seawater, so that conditions closely mimic the natural system. During this experiment, we examined physical and chemical processes, in the absence of biology, responsible for changes in the inorganic carbon system of sea ice and the underlying

seawater, and quantified fluxes of inorganic carbon between the atmosphere, sea ice and the water column. We also discuss that dissolution of ikaite crystals exported from sea ice in the underlying seawater can potentially hamper the effect of oceanic acidification on $\Omega_{\text{aragonite}}$.

2 Site description, sampling and analysis

The Sea-ice Environmental Research Facility (SERF) is an inground outdoor concrete pool, $18.3\text{ m} \times 9.1\text{ m}$ in surface area and 2.6 m deep, exposed to ambient temperatures, winds and solar radiation (by retracting its roof, Fig. 1). The weather conditions in the region are conducive to sea ice growth for several months every winter. Prior to the experiment, the pool is filled with artificial seawater (ASW) made by dissolving large quantities of various rock salts into local groundwater to mimic the major composition of natural seawater (see Rysgaard et al., 2014 for exact composition of the ASW). Sea ice is melted in the pool by circulating heated ethylene glycol through a closed-loop hose located at the bottom of the pool, allowing successive ice growth/melt experiments to be carried out during one winter. The experimental sea ice and brine exhibit similar physical and chemical properties to those observed in natural Arctic sea ice (Geilfus et al., 2013b; Hare et al., 2013). The experiment described herein was initiated from open water conditions on 11 January 2013 when the heater was turned off. Sea ice grew until 26 January when the heat was turned back on. The experiment ended on 30 January when the pool was 20 % ice-free.

Four 375 W pumps were installed on the bottom of the pool near each of the corners to induce a consistent current. The pumps were configured to draw water from their base and then propel it outward parallel to the bottom of the pool. The pumps were oriented successively at right angles to one another, which created a counterclockwise circulation of $2\text{--}3\text{ cm s}^{-1}$ (Else et al., 2015).

Bulk ice and seawater temperatures were recorded by an automated type-T thermocouple array fixed vertically in the pool. Seawater salinity was measured continuously using Aanderaa CT sensors (model 4319) located at 30, 100, 175 and 245 cm depth. The in situ seawater $p\text{CO}_2$ was measured every 5 s using a Contros HydroC (resolution $< 1\text{ }\mu\text{atm}$, accuracy $\pm 1\%$ of the upper range value) located at 1.3 m depth.

Air temperature and relative humidity were measured using a Vaisala HMP45C probe at a meteorological station located 2 m above the sea ice surface. Solar irradiance was continuously recorded by an Eppley Precision Spectral Pyranometer (range of $0.285\text{--}2.8\text{ }\mu\text{m}$) mounted 10 m above the sea ice surface. In addition, estimated photosynthetically active radiation (PAR) values at the ice bottom were recorded with Alec mkV/L PAR sensors throughout the study and ranged from 0 to $892\text{ }\mu\text{mol photons m}^{-2}\text{ s}^{-1}$.

Sea ice and seawater samples were obtained from a confined area located on the north side of the pool to minimize

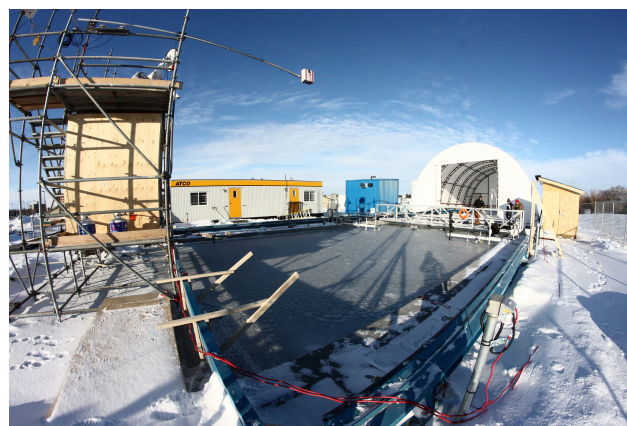


Figure 1. The Sea-ice Environmental Research Facility (University of Manitoba, Winnipeg, Canada) with thin sea ice covering the pond during the 2013 experiment. Photo: J. Sievers.

effects on other experiments (e.g., Else et al., 2015). Ice samples were collected using ceramic knives or a Kovacs Mark II coring system depending on the ice thickness. Sampling was performed from a movable bridge to avoid walking on the ice surface and to ensure only undisturbed sites were sampled. Ice cores were collected from one end of the pool (half meter away from the edge of the pool) and at least 20 cm away from previous cored sites. Ice cores were packed in clean plastic bags and kept frozen during the 20 min transport to a cold laboratory and processed within a few hours. Seawater was sampled for total alkalinity (TA) and total dissolved inorganic carbon (TCO_2) with a peristaltic pump (Cole Parmer Masterflex® Environment Sampler, equipped with PTFE tubing) through an ice core hole of the ice–water interface, at 1.25 and 2.5 m depth. Samples were stored in 12 mL gas-tight vials (Exetainer, Labco High Wycombe, United Kingdom) and poisoned with $12\text{ }\mu\text{L}$ of saturated HgCl_2 solution and stored in the dark at 4°C until analyzed.

Air–ice CO_2 fluxes were measured using a Li-Cor 8100-103 chamber associated with a LI-8100A soil CO_2 flux systems. The chamber was connected in a closed loop to the IRGA with an air pump rate of 3 L min^{-1} . The measurement of $p\text{CO}_2$ in the chamber was recorded every second over a 15 min period. The flux was computed from the slope of the linear regression of $p\text{CO}_2$ against time ($r^2 > 0.99$) according to Frankignoulle (1988), taking into account the volume of ice or snow enclosed within the chamber. The uncertainty of the flux computation due to the standard error on the regression slope was on average $\pm 3\%$.

In the cold laboratory, sea ice cores were cut into 2 cm sections using a pre-cleaned stainless steel band saw. Each section was placed in a gas-tight laminated (Nylon, ethylene vinyl alcohol and polyethylene) plastic bag (Hansen et al., 2000) fitted with a gas-tight Tygon tube and a valve for sampling. The plastic bag was sealed immediately, and excess air was gently removed through the valve using a vac-

uum pump. The bagged sea ice samples were then melted in the dark at 4 °C to minimize the dissolution of calcium carbonate precipitates (meltwater temperature never rose significantly above 0 °C). Once melted, the meltwater mixture and bubbles were transferred to gas-tight vials (12 mL Exetainer, Labco High Wycombe, United Kingdom), poisoned with 12 µL solution of saturated HgCl₂ and stored in the dark at 4 °C until analyzed.

Bulk ice and seawater salinities were measured using a Thermo Orion 3-star with an Orion 013610MD conductivity cell, and values were converted to bulk salinity (Grasshoff et al., 1983). TA was determined by potentiometric titration (Haraldsson et al., 1997), while TCO₂ was measured on a coulometer (Johnson et al., 1987). Routine analysis of Certified Reference Materials provided by A. G. Dickson, Scripps Institution of Oceanography, verified that TA and TCO₂ were analyzed within ± 3 and $\pm 2 \mu\text{mol kg}^{-1}$, respectively. Brine volume was estimated from measurements of bulk salinity, temperature and density according to Cox and Weeks (1983) for temperatures below −2 °C and according to Leppäranta and Manninen (1988) for ice temperatures within the range −2 to 0 °C.

Bulk ice samples for biological measurements were collected between 14 and 21 January. Filtered (0.2 µm) SERF seawater (FSW) was added at a ratio of three parts FSW to one part ice and the samples were left to melt in the dark. Chl *a* was determined on three occasions by filtering two aliquots of the melted ice sample onto GF/F filters (Whatman® brand) and extracting pigments in 10 mL of 90 % acetone for 24 h. Fluorescence was measured before and after the addition of 5 % HCl (with a Turner Designs fluorometer), and chl *a* concentration was calculated following Parsons et al. (1984). Measurements of bacterial production were done four times during the biological sampling period by incubating 6–10 mL subsamples of the ice–FSW solution with ³H-leucine (final concentration of 10 nM) for 3 h at 0 °C in darkness (Kirchmann, 2001). Half of the samples were spiked with trichloroacetic acid (TCA, final concentration 5 %) as controls prior to the incubation, while the remaining active subsamples were fixed with TCA (final concentration 5 %) after incubation. Following the incubation, vials were placed in 80 °C water for 15 min (Garneau et al., 2006) before filtration through 0.2 µm cellulose acetate membranes (Whatman® brand) and rinsing with 5 % TCA and 95 % ethanol. Filters were dried and dissolved in scintillation vials by adding 1 mL ethyl acetate, and radioactivity was measured on a liquid scintillation counter after an extraction period of 24 h. Bacterial production was calculated using the equations of Kirchman (1993) and a conversion factor of 1.5 kg C mol^{−1} (Ducklow et al., 2003).

3 Results

3.1 Sea ice and seawater physical conditions

Sea ice was grown in the pool from open water on 13 January 2013 and reached a maximum thickness of 24 cm on 26 January at which point the heat at the base of the pool was turned on. On 30 January the experiment ended with the pool 20 % ice-free. Three main snowfall events occurred during the experiment. The first, from 14 to 15 January, covered the sea ice surface with 1 cm of snow. The second, from 18 to 23 January, deposited 6–9 cm of snow over the entire pool. On the morning of 23 January, the snow was manually cleared off the ice surface to investigate the insulating effect of the snow on the ice temperature and ikaite precipitation (Rysgaard et al., 2014). Finally, from noon on 24 to 27 January, 8 cm of snow covered the entire pool until the end of the experiment on 30 January.

The air temperature at the beginning of the experiment ranged from −2 to −26 °C, which initiated rapid sea ice growth to 15 cm until 18 January (Fig. 2). During this initial sea ice growth, the sea ice was attached to the side of the pool, resulting in the development of a hydrostatic pressure head that caused percolation of seawater at the freezing point upwards through the sea ice volume as the sea ice grew downwards. This resulted in repeated events of increased sea ice temperature from the bottom to the surface observed between 15 and 18 January (Fig. 2). Subsequently, the ice was cut using an ice saw around the perimeter, allowing the ice to float, and a pressure release valve was installed to prevent such events (Rysgaard et al., 2014). During this period, the ice temperature oscillated between relatively warm (~ -3 °C) and cold (~ -7 °C) phases. Brine volume content (0.047) was low in the middle part of the ice cover, close to the permeability threshold of 0.05 as suggested by Golden et al. (2007). The bulk ice salinity profiles were typically C-shaped, with values ranging from 6 to 23 (Fig. 2). The underlying seawater salinity increased rapidly due to sea ice growth. From 18 to 23 January, the 9 cm snow cover insulated the ice cover from the cold atmosphere (Rysgaard et al., 2014), resulting in a fairly constant ice thickness, nearly no change in ice temperature and salinity, a brine volume content above the permeability threshold and a small increase in the underlying seawater salinity. Once the ice surface was cleared of snow on the morning of 23 January, the ice temperature decreased throughout the entire ice thickness and the ice surface salinity increased. The sea ice volume cooled from the top downwards, and the brine volume content decreased below the permeability threshold on 23 January and rapid sea ice growth rapidly increased the seawater salinity. Shortly after the snow clearing, the last snowfall event covered the ice surface with 8 cm of snow, reducing the effect of the cold atmosphere on the ice cover. On 26 January, the heater was activated to initiate sea ice melt. Sea ice temperatures increased and became isothermal around −2 °C, while

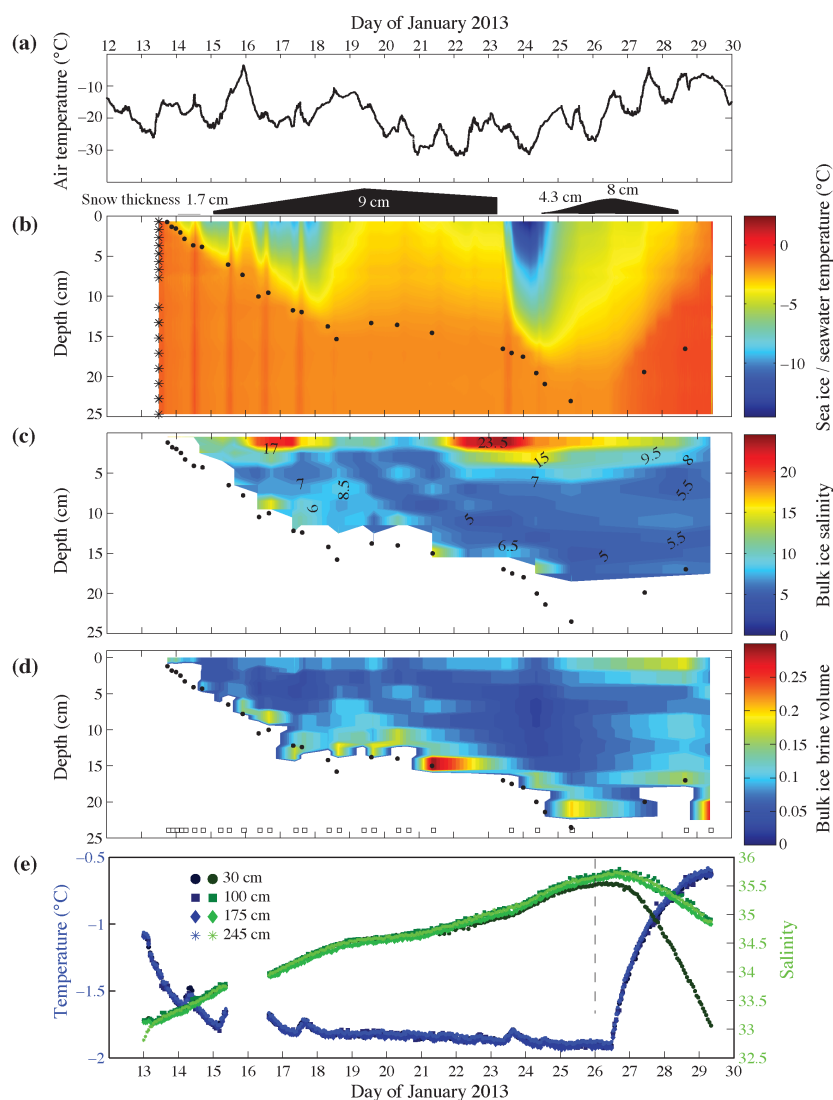


Figure 2. Evolution of (a) air temperature (°C) at 2 m height, (b) snow thickness (black shaded areas) and sea ice/seawater temperature (°C), (c) bulk ice salinity, (d) brine volume content within sea ice and (e) seawater temperature (blue) and salinity (green). Measurements were performed at 30, 100, 175 and 245 cm water depths. The darker the color is, the closer to the surface. In panels (b–d) sea ice thickness is illustrated by black dots. Stars on panel (b) represent the depth at which the temperature profiles are derived from. Open squares in the lower part of (d) mark the sampling times. The dashed line on panel (e) indicates when the heat at the bottom of the pool was turned back on.

the bulk ice salinity decreased and the brine volume content increased up to 0.13. The sea ice melt decreased the seawater salinity. The pool was well mixed during the whole growth phase with similar salinity and temperature observed at the four depths. However, once the heat was turned on, the pool became stratified with respect to salinity changes, as the salinity at 30 cm depth started to diverge from the deeper depths (Fig. 2).

3.2 Carbonate system

TA and TCO₂ in seawater, noted as TA_(sw) and TCO_{2(sw)}, were sampled at the sea ice–seawater interface, at 1.25 and

2.5 m depth. An analysis of variance (ANOVA) test over the three depths revealed that the means are not statistically different ($p < 0.01$), so we consider the average concentration of the three depths in the following analysis. During sea ice growth, TA_(sw) increased from 2449 to 2644 $\mu\text{mol kg}^{-1}$ (black line, Fig. 3a), while TCO_{2(sw)} increased from 2347 to 2516 $\mu\text{mol kg}^{-1}$ (black line, Fig. 3b). Once the ice started to melt, TA_(sw) decreased to 2607 $\mu\text{mol kg}^{-1}$, and TCO_{2(sw)} decreased to 2461 $\mu\text{mol kg}^{-1}$. As the experiment stopped before the ice was completely melted in the tank, both the seawater salinity and TA_(sw) do not reach their initial values by the end of the experiment (Table 1, Figs. 2, 3). To discard the effect of salinity changes, we normalized TA_(sw)

Table 1. Seawater conditions on 11 January, before any sea ice formation ($t = 0$), on 25 January, just before the heat was turned back on and on 29 January, at the end of the experiment. Note that seawater salinity and $\text{TA}_{(\text{sw})}$ do not reach the initial seawater values as sea ice was still present at the end of the experiment.

Date	Temperature (°C)	Salinity	$\text{TA}_{(\text{sw})}$	$\text{nTA}_{(\text{sw})}$	$\text{TCO}_{2(\text{sw})}$	$\text{nTCO}_{2(\text{sw})}$
			(μmol kg ⁻¹)			
11 January	−1.4	33.5	2453	2416	2341	2306
25 January	−1.9	35.5	2659	2471	2524	2346
29 January	−0.6	34.4	2607	2500	2461	2361

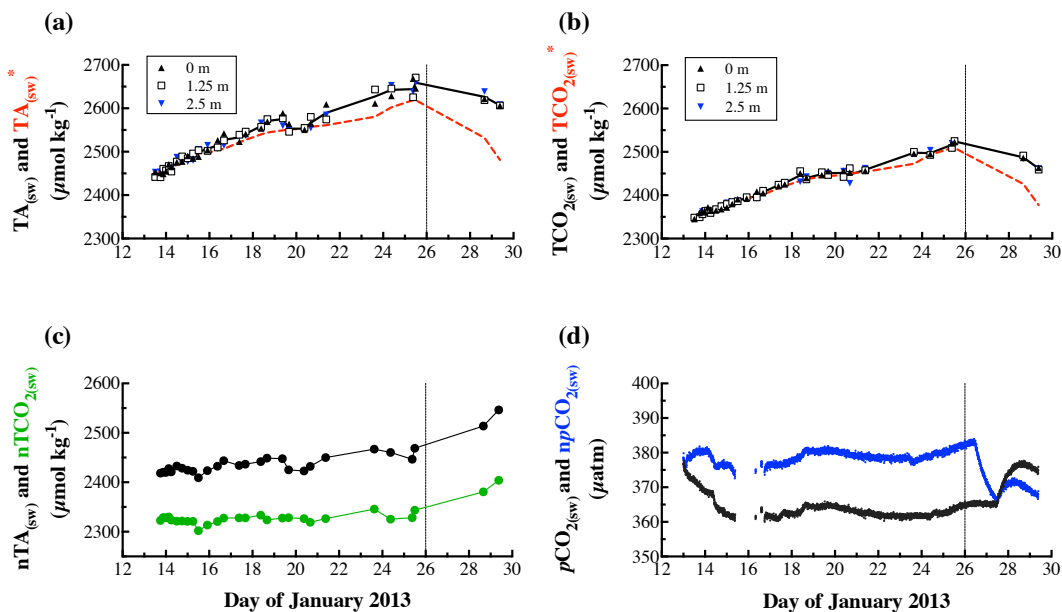


Figure 3. Evolution of (a) $\text{TA}_{(\text{sw})}$ and $\text{TA}_{(\text{sw})}^*$ (μmol kg⁻¹), (b) $\text{TCO}_{2(\text{sw})}$ and $\text{TCO}_{2(\text{sw})}^*$ (μmol kg⁻¹), (c) $\text{nTA}_{(\text{sw})}$ (black) and $\text{nTCO}_{2(\text{sw})}$ (green) (μmol kg⁻¹) and (d) the seawater $p\text{CO}_2$ (μatm) measured in situ (black) and corrected to a constant temperature of -1°C (blue). In panels (a) and (b) the black line is the average over the three depths, while the dotted red line is the expected concentrations according to the variation of salinity observed and calculated from the mean values of the three depths ($\text{TA}_{(\text{sw})}^*$ and $\text{TCO}_{2(\text{sw})}^*$, respectively). The vertical black dotted line on 26 January marks when the heat was turned back on.

and $\text{TCO}_{2(\text{sw})}$ to a salinity of 33 (noted as $\text{nTA}_{(\text{sw})}$ and $\text{nTCO}_{2(\text{sw})}$) according to Reactions (R4) and (R5):

$$\text{nTA}_{(\text{sample})t} = \frac{\text{TA}_{(\text{sample})t}}{S_{(\text{sample})t}} \times 33, \quad (\text{R4})$$

$$\text{nTCO}_{2(\text{sample})t} = \frac{\text{TCO}_{2(\text{sample})t}}{S_{(\text{sample})t}} \times 33, \quad (\text{R5})$$

where t is the time of the sampling and S the salinity of the sample (seawater or sea ice). During ice growth, $\text{nTA}_{(\text{sw})}$ and $\text{nTCO}_{2(\text{sw})}$ increased slightly to 2446 and 2328 μmol kg⁻¹, respectively (Fig. 3c). However, once the ice started to melt, $\text{nTA}_{(\text{sw})}$ increased to 2546 μmol kg⁻¹, and $\text{nTCO}_{2(\text{sw})}$ increased to 2404 μmol kg⁻¹.

The in situ $p\text{CO}_2$ of the underlying seawater ($p\text{CO}_{2(\text{sw})}$) decreased from 377 to 360 μatm as the seawater temperature in the pool decreased to the freezing point. The $p\text{CO}_{2(\text{sw})}$

then oscillated from 360 to 365 μatm during sea ice growth. One day after the heater was turned on, the $p\text{CO}_{2(\text{sw})}$ increased to a similar concentration as at the beginning of the experiment, before decreasing to 373 μatm by the end of the experiment (Fig. 3d).

Within bulk sea ice, $\text{TA}_{(\text{ice})}$ ranged from 300 to 1907 μmol kg⁻¹, while $\text{TCO}_{2(\text{ice})}$ ranged from 237 to 1685 μmol kg⁻¹. Both $\text{TA}_{(\text{ice})}$ and $\text{TCO}_{2(\text{ice})}$ exhibited C-shaped profiles with higher concentrations at the surface and bottom layers of the ice cover (Fig. 4). The concentration of $\text{TA}_{(\text{ice})}$ (average of 476 μmol kg⁻¹) and $\text{TCO}_{2(\text{ice})}$ (average of 408 μmol kg⁻¹) did not show significant variability during our survey, except at the surface of the ice. A first maximum was observed on 17 January with a concentration of 1907 μmol kg⁻¹ for $\text{TA}_{(\text{ice})}$ and 1685 μmol kg⁻¹ for $\text{TCO}_{2(\text{ice})}$. A second maximum was observed on 23 January with a concentration of 1433 μmol kg⁻¹ for $\text{TA}_{(\text{ice})}$ and

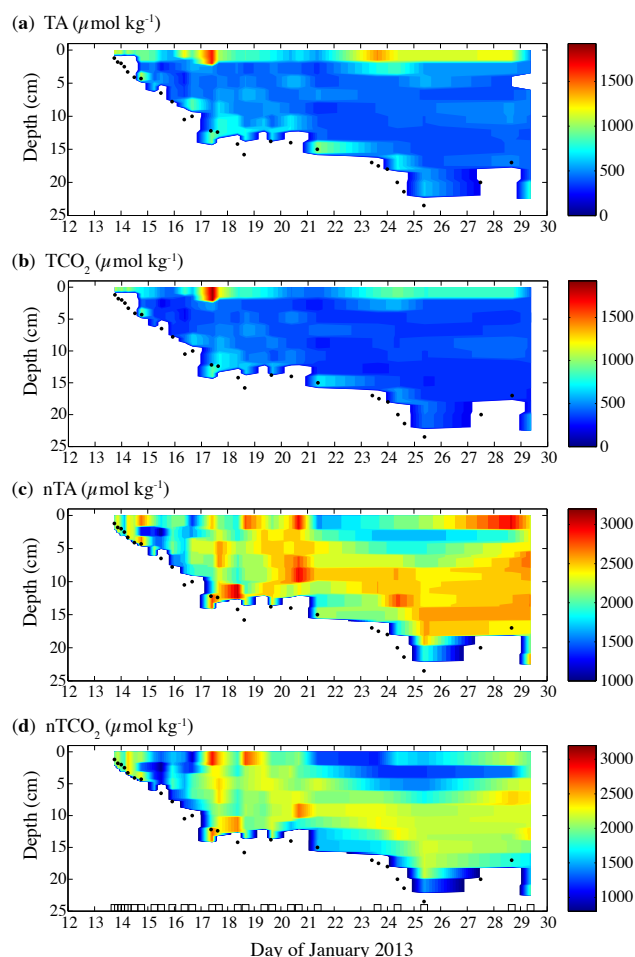


Figure 4. Evolution of (a) $TA_{(ice)}$ ($\mu\text{mol kg}^{-1}$), (b) $TCO_{2(ice)}$ ($\mu\text{mol kg}^{-1}$), (c) $nTA_{(ice)}$ ($\mu\text{mol kg}^{-1}$) and (d) $nTCO_{2(ice)}$ ($\mu\text{mol kg}^{-1}$). Sea ice thickness is illustrated by black dots. Open squares in the lower part of (d) mark the sampling times.

$861 \mu\text{mol kg}^{-1}$ for $TCO_{2(ice)}$. These maxima matched the high bulk ice salinity (Fig. 2), so we also normalized $TA_{(ice)}$ and $TCO_{2(ice)}$ (noted as $nTA_{(ice)}$ and $nTCO_{2(ice)}$, Fig. 4) to a salinity of 33 (according to the Reactions R4 and R5) to discard the effect of salinity changes and facilitate comparison with the underlying seawater. During initial sea ice formation (up to 17 January), the concentration of both $nTA_{(ice)}$ (from 1083 to 2741, average $1939 \mu\text{mol kg}^{-1}$) and $nTCO_{2(ice)}$ (from 853 to 2440, average $1596 \mu\text{mol kg}^{-1}$) were at their minima in the experimental time series. From 17 to 21 January, both $nTA_{(ice)}$ and $nTCO_{2(ice)}$ increased throughout the ice column (average $nTA_{(ice)}$ $2375 \mu\text{mol kg}^{-1}$ and $nTCO_{2(ice)}$ $2117 \mu\text{mol kg}^{-1}$). However, from 21 January until the initial sea ice melt, $nTA_{(ice)}$ and $nTCO_{2(ice)}$ decreased in the top 5 cm of the ice cover (average $nTA_{(ice)}$ $2125 \mu\text{mol kg}^{-1}$ and $nTCO_{2(ice)}$ $1635 \mu\text{mol kg}^{-1}$).

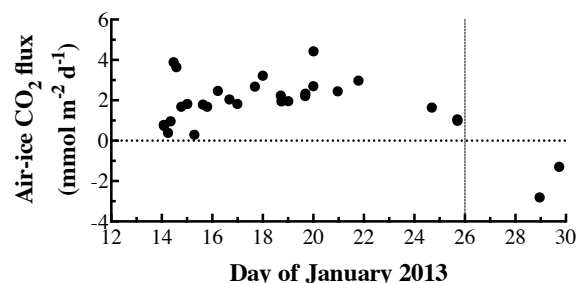


Figure 5. Air–ice CO_2 fluxes ($\text{mmol m}^{-2} \text{d}^{-1}$). Positive air–ice CO_2 flux means outgassing from the ice, and negative CO_2 flux means uptake of atmospheric CO_2 . The vertical black dotted line on 26 January marks when the heat was turned back on.

3.3 Air–ice CO_2 fluxes

The CO_2 fluxes measured at the variably snow-covered sea ice surface (Fig. 2b), which ranged from 0.29 to $4.43 \text{ mmol m}^{-2} \text{d}^{-1}$, show that growing sea ice released CO_2 to the atmosphere (Fig. 5). However, as soon as the ice started to warm up and then melt, the sea ice switched from source to sink for atmospheric CO_2 , with downward fluxes from -1.3 to $-2.8 \text{ mmol m}^{-2} \text{d}^{-1}$. These ranges of air–ice CO_2 exchanges are of the same order of magnitude as fluxes reported on natural sea ice using the same chamber technique in the Arctic during the initial sea ice growth (from 4.2 to $9.9 \text{ mmol m}^{-2} \text{d}^{-1}$ in Geilfus et al., 2013a) and during the spring–summer transition (from -1.4 to $-5.4 \text{ mmol m}^{-2} \text{d}^{-1}$ in Geilfus et al., 2015). In Antarctica, air–ice CO_2 fluxes were reported during the spring–summer transition from 1.9 to $-5.2 \text{ mmol m}^{-2} \text{d}^{-1}$ by Delille et al. (2014), from 0.3 to $-2.9 \text{ mmol m}^{-2} \text{d}^{-1}$ (Geilfus et al., 2014) and from 0.5 to $-4 \text{ mmol m}^{-2} \text{d}^{-1}$ (Nomura et al., 2013).

4 Discussion

4.1 Key processes affecting the carbonate system

The dynamics of inorganic carbon in the ocean and sea ice are mainly affected by temperature and salinity changes, precipitation and dissolution of calcium carbonate and biological activities (Zeebe and Wolf-Gladrow, 2001). During this experiment, neither organic matter nor biota were purposely introduced into the pool; the observed range of bulk ice microbial activity (5.7×10^{-9} on 14 January to $7.5 \times 10^{-7} \text{ g CL}^{-1} \text{h}^{-1}$ on 21 January) and algal chl *a* (0.008 on 14 January to $0.002 \mu\text{g L}^{-1}$ on 21 January) were too low to support any biological activity (Rysgaard et al., 2014). Therefore biological activity is unlikely to have played a role. During the same 2013 time series at SERF, Rysgaard et al. (2014) discussed the precipitation of ikaite within the ice cover in detail, reporting high concentrations of ikaite ($> 2000 \mu\text{mol kg}^{-1}$) at the surface of the ice in brine skim and

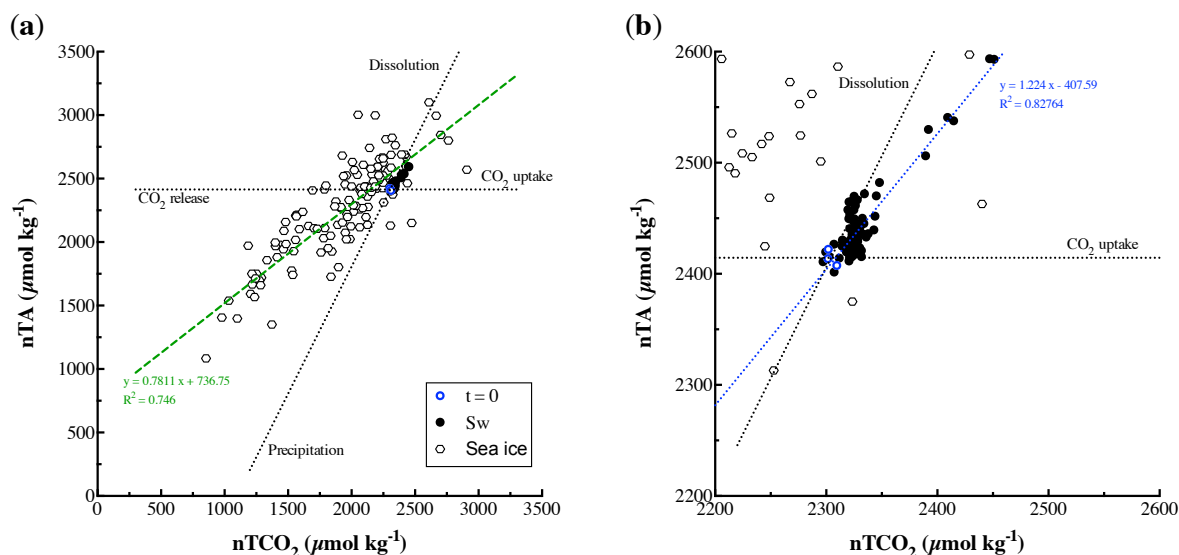


Figure 6. (a) Relationship between nTCO₂ and nTA (μmol kg⁻¹) in bulk sea ice (white hexagons) and seawater (black dots). (b) Zoomed-in image of seawater data. The different dotted lines represent the theoretical evolution of nTA and nTCO₂ ratio following the precipitation/dissolution of calcium carbonate and release/uptake of CO_{2(g)}. A linear regression is shown in green for the ice samples (a) and blue for the seawater samples (b).

frost flowers, and ikaite precipitation up to 350 μmol kg⁻¹ within bulk sea ice. Within sea ice, ikaite precipitation is associated with low ice temperatures, high bulk salinity and high TA_(ice) and TCO_{2(ice)} concentrations (Figs. 2, 3).

The main processes affecting the carbonate system can be described by changes in TA and TCO₂ (Zeebe and Wolf-Gladrow, 2001). An exchange of CO_{2(gas)} affects TCO₂, while TA remains constant and the precipitation–dissolution of calcium carbonate affects both TA and TCO₂ in a ratio of 2 : 1 (see Reactions R1–R3, Fig. 6). To calculate the theoretical changes in TA and TCO₂ during the course of the experiment, we used seawater samples from 11 January prior to sea ice formation ($t = 0$, Table 1) as the origin point (blue circle on Fig. 6). Sea ice data are located between the theoretical calcium carbonate precipitation line and the CO₂ release line (Fig. 6a), while seawater data mainly fall on the calcium carbonate dissolution line (Fig. 6b), suggesting that the carbonate system within sea ice is affected by both the precipitation of ikaite and a release of CO_{2(gas)}, while the underlying seawater is mainly affected by the dissolution of calcium carbonate.

4.2 Estimation of the precipitation–dissolution of ikaite

During the experiment, Rysgaard et al. (2014) observed ikaite within sea ice using direct microscopic observations. The precipitation–dissolution of ikaite and gas exchange are the only two processes taking place during the experiment. As illustrated in Fig. 6, an exchange of CO₂ does not affect TA, while the precipitation–dissolution of ikaite affects TA and TCO₂ in a ratio 2 : 1. Therefore, we use TA to estimate

how much ikaite is precipitated or dissolved within the ice cover and the underlying seawater.

Assuming no biological effect, ikaite precipitation/dissolution and gas exchange, TA and TCO₂ are considered conservative with salinity. Therefore, we can calculate the expected TA and TCO₂ (noted as TA_(ice)^{*} and TCO_{2(ice)}^{*} in the ice cover and TA_(sw)^{*} and TCO_{2(sw)}^{*} for the water column) based on the initial seawater conditions (TA_(sw), TCO_{2(sw)} and $S_{(sw)}$ at $t = 0$, Table 1) and the sample salinity (bulk sea ice or seawater) measured during the experiment:

$$TA_{(sample) t}^* = \frac{TA_{(sw) t=0}}{S_{(sw) t=0}} \times S_{(sample) t}, \quad (R6)$$

$$TCO_{2(sample) t}^* = \frac{TCO_{2(sw) t=0}}{S_{(sw) t=0}} \times S_{(sample) t}, \quad (R7)$$

where t is the time of the sampling. Within the ice cover, TA_(ice), TCO_{2(ice)} and the bulk ice salinity are averaged throughout the ice column on each sampling day (Fig. 7a, b, black line), while for the underlying seawater, we used the averaged TA_(sw), TCO_{2(sw)} and salinity for all the measured depths (Fig. 2a, b, black line). The difference between TA_(sample)^{*} and the observed TA is only due to the precipitation or dissolution of ikaite crystals. In case of ikaite precipitation (i.e., TA_(sample)^{*} > TA_(sample)), half of this positive difference corresponds to the amount of ikaite precipitated within the ice. This ikaite may either remain or may be exported out of the ice. A negative difference (i.e., TA_(sample)^{*} < TA_(sample)) indicates ikaite dissolution.

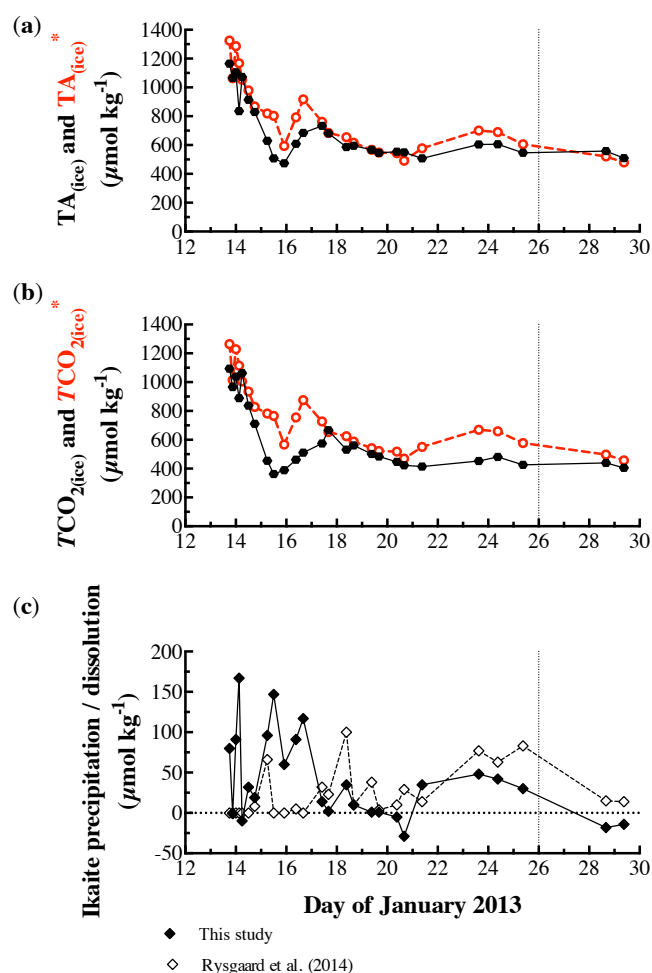


Figure 7. Evolution of (a) $TA_{(ice)}$ averaged throughout the ice thickness on each sampling day (black dots) and $TA_{(ice)}^*$ (dashed red line) ($\mu\text{mol kg}^{-1}$) and (b) $TCO_{2(ice)}$ averaged throughout the ice thickness on each sampling day (black diamonds) and $TCO_{2(ice)}^*$ (dashed red line) ($\mu\text{mol kg}^{-1}$). (c) Estimation of the ikaite precipitation/dissolution from half of the difference between $TA_{(ice)}^*$ and $TA_{(ice)}$ ($\mu\text{mol kg}^{-1}$) (black diamonds) compared to the average amount of ikaite precipitated throughout the ice thickness for each sampling day from Rysgaard et al. (2014) (white dots). The vertical black dotted line on 26 January marks when the heat was turned back on.

4.2.1 Sea ice

Greater $TA_{(ice)}^*$ and $TCO_{2(ice)}^*$ compared to the averaged observed $TA_{(ice)}$ and $TCO_{2(ice)}$ (Fig. 7a, b) are expected as ikaite is precipitated and CO_2 is released from the ice to the atmosphere (Figs. 5, 6). Half the difference between $TA_{(ice)}^*$ and $TA_{(ice)}$ is a result of ikaite precipitation (Fig. 7c, black diamonds). Highly variable ikaite precipitation was observed (Fig. 7c). Ikaite precipitation was up to $167 \mu\text{mol kg}^{-1}$ (e.g., first days of the experiment) and as low as $1 \mu\text{mol kg}^{-1}$ (e.g., 19 January). A negative difference between $TA_{(ice)}^*$ and

$TA_{(ice)}$ (i.e., ikaite dissolution) occurred on three occasions: 14, 20 and after 26 January (beginning of the sea ice melt). On these occasions, the ice cover was relatively warm due to warmer atmospheric temperatures (14 January), thicker snow cover insulating the ice cover from the cold atmosphere (20 January) or when heat was turned back on (after 26 January, Fig. 2). Relatively high sea ice temperatures likely promote ikaite dissolution in agreement with Rysgaard et al. (2014), who linked ikaite precipitation/dissolution to ice temperature. The upward percolation of seawater observed from 15 to 18 January might complicate the effect of sea ice temperature on ikaite formation because it was in part responsible for increased ice temperatures (Fig. 2b) and therefore increased the sea ice brine volumes (Fig. 2c). Increased vertical connectivity (permeability) of the network of liquid inclusions throughout the sea ice (Golden et al., 2007; Galley et al., 2015) would have allowed the export of ikaite crystals from the ice cover to the underlying seawater. However, while we calculated a negative difference between $TA_{(ice)}^*$ and $TA_{(ice)}$, ikaite crystals were observed by Rysgaard et al. (2014). We compared the direct microscopy observations by averaging the amount of ikaite precipitated throughout the ice thickness for each sampling day from Rysgaard et al. (2014) (Fig. 7c, white dots) with our estimation of the amount of ikaite based on the difference between $TA_{(ice)}^*$ and $TA_{(ice)}$ (Fig. 7c, black diamonds). Both ikaite measurements are of the same order of magnitude; however, the average ($22 \mu\text{mol kg}^{-1}$) and maximum ($100 \mu\text{mol kg}^{-1}$) of direct observations presented by Rysgaard et al. (2014) were lower than our estimated average ($40 \mu\text{mol kg}^{-1}$) and maximum of up to $167 \mu\text{mol kg}^{-1}$ over this whole experiment. Deviations are likely due to methodological differences. Here, sea ice samples were melted to subsample for TA and TCO_2 . Ikaite crystals may have dissolved during melting, leading to an underestimation of the total amount of ikaite precipitated in the ice. However, the difference between $TA_{(ice)}^*$ and $TA_{(ice)}$ provides an estimation of how much ikaite is precipitated in the ice cover, including those crystals potentially already exported to the underlying seawater. The method used by Rysgaard et al. (2014) avoids the bias of ikaite dissolution during sea ice melt with the caveat that crystals need to be large enough to be optically detected. If no crystals were observed, Rysgaard et al. (2014) assumed that no crystals were precipitated in the ice, though ikaite crystals could have been formed and then exported into the underlying seawater prior to microscopic observation of the sample, which may explain the difference observed between both methods during initial sea ice formation (15–18 January) when the ice was still very thin. In addition, the succession of upward percolation events could have facilitated the ikaite export from the ice cover to the underlying seawater. Estimations from both methods show similar concentrations when the ice (i) warmed due to snowfall (18–23 January) and (ii) cooled once the snow was removed (on 23 January). Once the ice started to melt (26 January), Rysgaard et al. (2014) reported a decrease in

the ikaite precipitation, while in this study we reported a negative difference between $TA_{(ice)}^*$ and $TA_{(ice)}$, possibly indicating that ikaite dissolved in the ice.

4.2.2 Water column

The main process affecting the carbonate system in the underlying seawater in this study is the export of ikaite from the ice and its dissolution in the water column (Fig. 6). While a few studies of ikaite precipitation within sea ice carried out over open ocean hypothesized that ikaite remained trapped within the sea ice matrix (Rysgaard et al., 2007, 2013; Delille et al., 2014), the observed increase of $nTA_{(sw)}$ (Fig. 3) suggests that ikaite precipitated within the ice cover was exported to the underlying seawater where the crystals were dissolved, as suggested by Fransson et al. (2013). Lower $TA_{(sw)}^*$ and $TCO_{2(sw)}^*$ compared to $TA_{(sw)}$ and $TCO_{2(sw)}$ (Fig. 3) confirm the dissolution of ikaite in the underlying seawater, as the dissolution of ikaite crystals will decrease both TA and TCO_2 (Reactions R1–R3). Therefore, half the difference between $TA_{(sw)}^*$ and $TA_{(sw)}$ corresponds to the concentration of ikaite exported from the ice and dissolved in the underlying seawater (Fig. 8a). This concentration increased over time to a maximum of $66 \mu\text{mol kg}^{-1}$.

During this experiment, $nTA_{(sw)}$ increased by $128 \mu\text{mol kg}^{-1}$, while $nTCO_{2(sw)}$ increased by $82 \mu\text{mol kg}^{-1}$ (Fig. 3c). This suggests that $64 \mu\text{mol kg}^{-1}$ of ikaite is dissolved compared to the $66 \mu\text{mol kg}^{-1}$ estimated from the difference between $TA_{(sw)}^*$ and $TA_{(sw)}$. As a result of the effect of ikaite dissolution on the 2 : 1 ratio of TA : TCO_2 , the dissolution of ikaite accounts for the entire increase of $nTA_{(sw)}$ but only accounts for $64\text{--}66 \mu\text{mol kg}^{-1}$ of the $82 \mu\text{mol kg}^{-1}$ increase in $nTCO_{2(sw)}$. Therefore, $16\text{--}18 \mu\text{mol kg}^{-1}$ (about 25 %) of the increase of $nTCO_{2(sw)}$ cannot be explained by the dissolution of ikaite. The increase of both $nTA_{(sw)}$ and $nTCO_{2(sw)}$ is more significant once the ice starts to melt (26 January). During sea ice melt, increased vertical permeability resulting in increased liquid communication through the sea ice volume from below likely, in part, dissolved ikaite crystals still residing in the ice at that time, and also will have created a downward crystal export mechanism. As the ice melt advanced, patches of open water occurred at the surface of the pool. Therefore, uptake of atmospheric CO_2 by the undersaturated seawater likely occurred, increasing the $TCO_{2(sw)}$.

The dissolution of ikaite crystals could also have a strong impact on the $pCO_{2(sw)}$. The water column was undersaturated compared to the atmosphere during the whole experiment (Fig. 3d). A release of CO_2 , from the ice to the atmosphere, was measured during sea ice growth (Fig. 5) in spite of the undersaturated $pCO_{2(sw)}$. This suggests that air–ice CO_2 fluxes are only due to the concentration gradient between the ice and the atmosphere (Geilfus et al., 2012; Nomura et al., 2013) but that sea ice exchanges CO_2 with the atmosphere independently of the seawater concentration

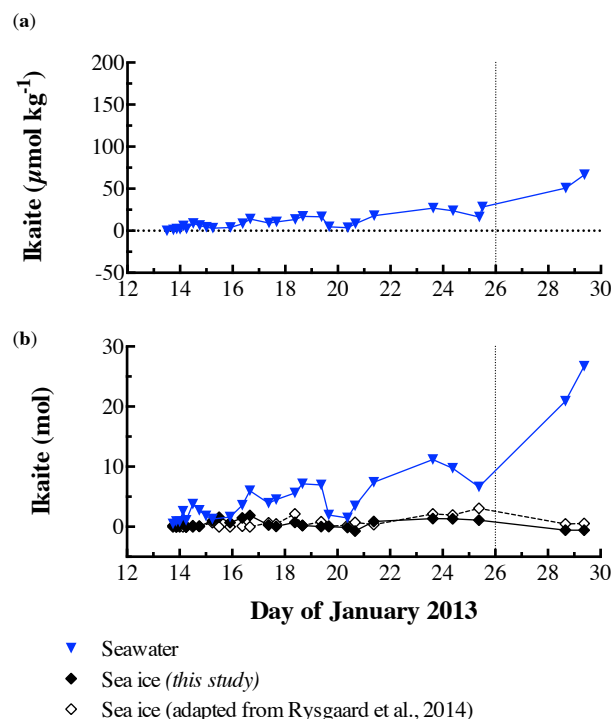


Figure 8. Evolution of (a) ikaite dissolution within the water column (in $\mu\text{mol kg}^{-1}$), (b) mass of ikaite dissolved in the underlying seawater (blue) and mass of ikaite precipitated in sea ice (black) estimated from this study and estimated from Rysgaard et al. (2014) (white). The vertical black dotted line on 26 January marks when the heat was turned back on.

(Geilfus et al., 2014). The $pCO_{2(sw)}$ is highly correlated with the seawater temperature (Fig. 2), with a rapid decrease of $pCO_{2(sw)}$ during the first days of the experiment (13–15 January) and a relative constant $pCO_{2(sw)}$ until 27 January. However, on 26 January, the heat was turned back on, affecting the seawater temperature on the same day (Fig. 2), while the impact of increasing temperature on the $pCO_{2(sw)}$ appeared 1 day later (Fig. 3d). We normalized the $pCO_{2(sw)}$ to a temperature of -1°C (after Copin-Montegut, 1988, noted as $npCO_{2(sw)}$, blue line on Fig. 3d). The $npCO_{2(sw)}$ does not show major variations during sea ice growth, with values around $380 \mu\text{atm}$. However, once the heat is turned on and the seawater temperature increased (on 26 January), $npCO_{2(sw)}$ decreased from 383 to $365 \mu\text{atm}$, while $pCO_{2(sw)}$ did not change in response to increased seawater temperatures until 27 January, suggesting that a process other than temperature change affected the $pCO_{2(sw)}$. According to Reaction (R1), the dissolution of calcium carbonate has the potential to reduce $pCO_{2(sw)}$. Therefore, during sea ice growth and the associated release of salt, TA, TCO_2 and ikaite crystals to the underlying seawater, ikaite dissolution within the seawater could be responsible for maintaining stable $pCO_{2(sw)}$ values while seawater salinity, $TA_{(sw)}$ and $TCO_{2(sw)}$ are increasing. Once the seawater temperature increased (26 January), sea

Table 2. Masses of TCO_2 in the water column ($\text{TCO}_{2(\text{sw})}$) and in the ice cover ($\text{TCO}_{2(\text{ice})}$), masses of ikaite within the ice cover estimated from this study and from Rysgaard et al. (2014), masses of ikaite dissolved in the water column ($\text{Ikaite}_{(\text{sw})}$) and masses of CO_2 exchanged between the ice and the atmosphere over the whole pool (estimation based on the air–ice CO_2 fluxes). All units are in mol.

January (DOY)	$\text{TCO}_{2(\text{sw})}$	$\text{TCO}_{2(\text{ice})}$	$\text{Ikaite}_{(\text{ice})}$ from this study	$\text{Ikaite}_{(\text{ice})}$ from Rysgaard et al. (2014)	$\text{Ikaite}_{(\text{sw})}$	$\text{CO}_{2(\text{air-ice})}$
$t = 0$	1041	–	–	–	–	–
13.75	1040	2.38	0.17	0.00	0.47	–
13.88	1044	2.09	0.00	0.00	0.87	–
14	1043	2.90	0.25	0.00	0.83	0.03
14.13	1043	3.29	0.62	0.00	2.57	0.02
14.25	1038	4.91	–0.05	0.00	1.06	0.01
14.5	1037	4.77	0.18	0.00	3.75	0.12
14.75	1039	4.36	0.12	0.05	2.73	0.07
15	1037	–	–	–	1.80	0.08
15.25	1032	4.67	0.98	0.68	1.28	0.01
15.5	1034	3.89	1.58	0.00	–1.57	0.07
15.92	1034	4.47	0.69	0.00	1.63	0.12
16.38	1024	7.36	1.45	0.08	3.60	0.19
16.67	1028	8.17	1.87	0.00	6.00	0.10
17.38	1023	15.48	0.29	0.65	3.90	0.22
17.67	1026	13.26	0.04	0.46	4.50	0.13
18.38	1030	11.39	0.74	2.14	5.61	0.38
18.67	1027	12.06	0.21	0.21	7.16	0.10
19.38	1029	11.13	0.01	0.84	6.96	0.23
19.67	1030	10.75	0.03	0.09	1.97	0.11
20.38	1028	10.25	–0.12	0.23	1.47	0.42
20.67	1022	10.36	–0.70	0.71	3.48	0.12
21.38	1025	10.50	0.88	0.35	7.42	0.35
23.63	1034	12.60	1.34	2.14	11.18	–
24.38	1026	14.84	1.30	1.94	9.75	0.21
25.38	1017	15.67	1.09	3.05	6.62	–
25.5	1029	–	–	–	11.51	0.02
28.67	1022	13.46	–0.57	0.46	20.91	–0.93
29.38	987.3	15.82	–0.56	0.55	26.72	–0.15

ice melt likely released ikaite crystals to the underlying seawater (Figs. 2, 8a) along with brine and meltwater, a process that would continuously export ikaite from the sea ice as the volume interacting with the seawater via percolation or convection increased. The dissolution of these crystals likely contributed to keeping the $p\text{CO}_{2(\text{sw})}$ low and counterbalancing the effect of increased temperature. We argued that once all the ikaite crystals are dissolved, the increased seawater temperature increased the $p\text{CO}_{2(\text{sw})}$ simultaneously with the $n\text{pCO}_{2(\text{sw})}$ (27 January, Fig. 3).

4.3 Ikaite export from the ice cover to the water column

We estimated the amount of ikaite precipitated and dissolved within sea ice and seawater based on the sea ice (and seawater) volume (in m^3), the sea ice and seawater density, the concentration of ikaite precipitated and dissolved within the ice cover (Fig. 7c) and the concentration of ikaite dissolved in the water column (Fig. 8a). Within the ice cover, the amount

of ikaite precipitated/dissolved ranged from -0.7 to 1.97 mol (Fig. 8b, Table 2), with a maximum just after the snow was cleared on 23 January. In the underlying seawater, the amount of ikaite dissolved in the pool increased from 0.47 mol on the first day of the experiment to 11.5 mol on 25 January when sea ice growth ceased. Once the ice started to melt, the amount of dissolved ikaite increased up to 20.9 (28 January) and 26.7 mol (29 January, Table 2). The estimation of ikaite dissolution in the pool is significantly higher than the estimated amount of ikaite precipitated (and potentially exported) within the ice cover, especially during sea ice melt. Within the ice cover, the ikaite values presented here represent a snapshot of the ikaite content in the ice at the time of sampling. In the underlying seawater, ikaite dissolution increased $\text{TA}_{(\text{sw})}$ cumulatively over time.

The difference between $\text{TA}_{(\text{ice})}^*$ and $\text{TA}_{(\text{ice})}$ provides an estimation of ikaite precipitated within the ice, including potential ikaite export to the underlying seawater, so it cannot be used to determine how much ikaite remained in the ice

vs. how much dissolved in the water column. However, Rysgaard et al. (2014) indicate that ikaite precipitated within the ice based on direct observations. Using the ikaite concentration reported in Rysgaard et al. (2014) (and shown in Fig. 7c), the sea ice volume (in m^3) and density, we calculate that 0–3.05 mol of ikaite precipitated within the ice cover during sea ice growth (Fig. 8b, Table 2). This amount decreased to 0.46 and 0.55 mol during the sea ice melt (28 and 29 January, respectively). Increased ikaite dissolution in the water column when the ice began to melt (from 11.5 to 20.9 mol) indicates that 9.4 mol of ikaite was stored in the ice and rejected upon the sea ice melt. This amount is about 3 times the amount of ikaite precipitated in the ice estimated by Rysgaard et al. (2014) at the end of the growth phase (3.05 mol, Table 2), suggesting more work is needed for the best estimate of ikaite precipitation within sea ice.

Once the ice started to melt, the increased ikaite dissolution from 11.5 to 20.9 mol (28 January) and to 26.7 mol (29 January) suggests that about the same amount of ikaite is dissolved during the sea ice growth as during the first 2 days of the sea ice melt. The amount of ikaite dissolved in the water column after melt commenced continued to increase cumulatively, suggesting that ikaite is continuously exported to the underlying seawater as increased sea ice temperatures permit more of the volume to communicate with the underlying seawater. Therefore, we can assume that more than half of the amount of ikaite precipitated within the ice remained in the ice cover before ice melt began.

4.4 Air–ice–seawater exchange of inorganic carbon

SERF is a semi-closed system where the only way for the surface (water or sea ice) to gain or lose CO_2 is through exchange with the atmosphere, making it reasonable to track the exchange of TCO_2 in the atmosphere–sea ice–seawater system. The ice cover always had lower $\text{TCO}_{2(\text{ice})}$ during the experiment ($\text{TCO}_{2(\text{ice})}^* > \text{TCO}_{2(\text{ice})}$) compared to what would be expected if the CO_2 simply followed brine rejection in a conservative process (i.e., $\text{TCO}_{2(\text{ice})}^*$) (Fig. 7b). This could be due to (i) CO_2 released to the atmosphere from sea ice, (ii) decreased $\text{TCO}_{2(\text{ice})}$ due to the precipitation of ikaite within sea ice and/or (iii) sea ice exchanging TCO_2 with the underlying seawater.

The number of moles of TCO_2 exchanged during this experiment was calculated using the sea ice (and seawater) volume (in m^3) and density (in kg m^{-3}). The total amount of $\text{TCO}_{2(\text{ice})}$ lost from the ice cover (the difference between $\text{TCO}_{2(\text{ice})}^*$ and $\text{TCO}_{2(\text{ice})}$) ranged from 0.11 to 6.02 mol (average 2.38 mol, Fig. 9, black dots). The greatest sea ice TCO_2 losses occurred on 15–16 January, during initial sea ice growth, and from 23 to 25 January, during ice cooling due to snow removal. The exchange of CO_2 between the ice and the atmosphere is known (Fig. 5). The number of moles of CO_2 exchanges between the ice and the atmosphere was calculated (noted as $\text{CO}_{2(\text{air}–\text{ice})}$ in Table 2) using the time

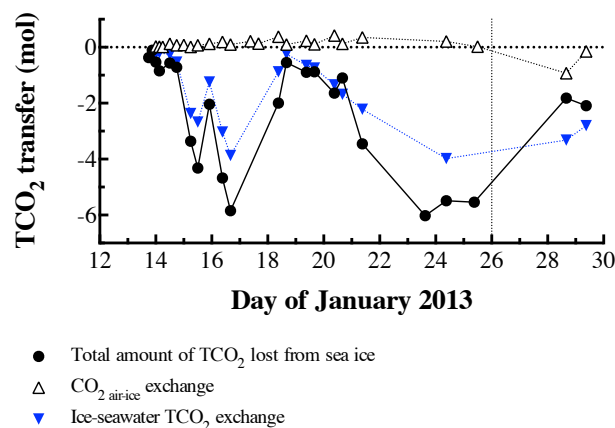


Figure 9. Total amount of TCO_2 lost from the ice cover (black dots), amount of CO_2 exchange between the atmosphere and the ice cover ($\text{CO}_{2\text{air}–\text{ice}}$, white triangle) and sea ice–seawater TCO_2 exchanges (blue triangle) measured in moles for each day, integrated over the whole tank. The dotted line on 26 January marks when the heat was turned back on.

step between each flux measurement, the ice thickness and the density. During sea ice growth, 0.01–0.42 mol of CO_2 was released from the ice-covered pool to the atmosphere. During sea ice melt, uptake of atmospheric CO_2 by the ice-covered pool ranged from -0.15 to -0.93 (Fig. 9, white triangles). On average, over the duration of the experiment, the ice cover released 0.08 mol of CO_2 to the atmosphere. Assuming we know how much ikaite is contained within the ice cover (Fig. 8b), we can estimate how much TCO_2 is exported from the ice to the underlying seawater (Fig. 9, blue triangles) by subtracting the air–ice CO_2 exchange and the ikaite precipitation from the total reduction of $\text{TCO}_{2(\text{ice})}$ observed within the ice cover (Fig. 9, black dots). The sea ice–seawater TCO_2 export ranged from 0.2 to 3.98 mol (average = 1.7 mol), confirming that sea ice primarily exports TCO_2 to the underlying seawater. TCO_2 export from the ice to the water column ranged from 23 % of the total sea ice TCO_2 early in the ice growth (14 January) to 100 % after the onset of melt. These estimations are comparable to the study of Sejr et al. (2011), who suggested that sea ice exports 99 % of its total TCO_2 to the seawater below it. On average over the whole experiment, sea ice exported 1.7 mol of TCO_2 to the underlying seawater (Fig. 9), which corresponds to a $\text{TCO}_{2(\text{sw})}$ increase of $43.5 \mu\text{mol kg}^{-1}$ considering the average sea ice thickness and density during the experiment and the volume of the pool. However, $\text{TCO}_{2(\text{sw})}$ increased by $115 \mu\text{mol kg}^{-1}$ over the whole experiment (Fig. 3b), leaving an increase of $71.5 \mu\text{mol kg}^{-1}$ in the $\text{TCO}_{2(\text{sw})}$ that cannot be explained by the sea ice–seawater exchange of TCO_2 . We postulate that as the ice melt advanced, patches of open water that opened at the surface of the pool which were undersaturated compared to the atmosphere (Fig. 3d) imported the additional TCO_2 directly from the atmosphere in the form

of $\text{CO}_2(\text{g})$. Considering the pool volume, the $71.5 \mu\text{mol kg}^{-1}$ increase of $\text{TCO}_{2(\text{sw})}$ could be explained by an air–sea water CO_2 uptake of $8.5 \text{ mmol m}^{-2} \text{ d}^{-1}$ over 3 days of sea ice melt in a 20 % ice-free pool. High air–sea gas exchange rates have been observed over partially ice-covered seas (Else et al., 2011, 2013). This mechanism is also corroborated by models that account for additional sources of turbulence generated by the presence of sea ice (Loose et al., 2014).

The design of the experiment allowed for constrained measurements of inorganic carbon fluxes between sea ice and the water column not possible in a natural environment where large-volume mixing processes alter the underlying seawater, making it more complicated to identify changes. We build a CO_2 budget based only on the sea ice growth phase because only 2 days of data for the melt phase are available, and the experiment stopped while the pool was 20 % ice-free (Rysgaard et al., 2014; Else et al., 2015). The initial seawater (origin point, $t = 0$) contained 1041 mol of $\text{TCO}_{2(\text{sw})}$ on 11 January, while on the last day of sea ice growth (25 January) the seawater contained 1017 mol of $\text{TCO}_{2(\text{sw})}$ (Table 2), with the difference (24 mol of TCO_2) in all likelihood transferred from the water column to the ice cover or the atmosphere. However, the TCO_2 content within the ice cover at the end of the growing phase was 15.6 mol and the ice cover released 3.1 mol of CO_2 to the atmosphere (Table 2). Therefore, 4.9 of the 24 mol of TCO_2 exchanged from the water column is unaccounted for, but may be explained by air–ice CO_2 fluxes. The chamber measurement technique for air–ice CO_2 flux may underestimate the exchange of CO_2 , and the air–seawater CO_2 fluxes are unknown until the ice started to grow (13 January). These missing moles of TCO_2 may also be explained by our assumption of uniform sea ice thickness in the SERF. Using the seawater conditions at the end of the experiment, 1 cm of seawater in the pool contains 4.21 mol of TCO_2 , making it difficult to close our budget.

4.5 Potential impact of sea ice growth and ikaite export on aragonite saturation state of the underlying seawater

The Arctic Ocean is a region where calcifying organisms are particularly vulnerable to ocean acidification since low temperatures and low salinity lower the carbonate saturation state. As a result, several areas of the Arctic Ocean are already undersaturated with respect to aragonite (Chierici and Fransson, 2009; Yamamoto-Kawai et al., 2009; Bates et al., 2011). This undersaturation is enhanced in winter as the temperature decreases and $p\text{CO}_2$ increases as a result of respiration. Calcifying organisms might therefore be most susceptible to the effects of acidification in the winter, corresponding to the annual minimum in aragonite saturation state ($\Omega_{\text{aragonite}}$). Sea ice retreat is thought to enhance the impact of ocean acidification by freshening and ventilating the surface water (Yamamoto-Kawai et al., 2008; Yamamoto et al., 2012; Popova et al., 2014). However, any understanding

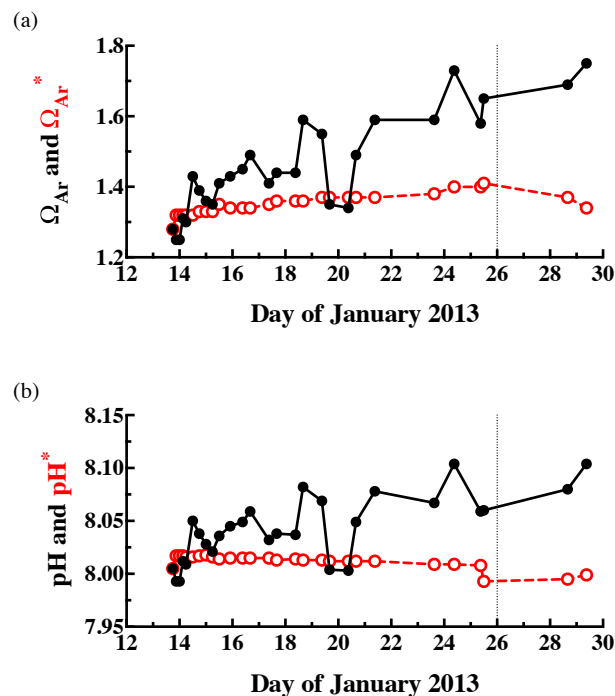


Figure 10. Evolution of (a) $\Omega_{\text{aragonite}}$ in the water column, calculated based on $\text{TA}_{(\text{sw})}$ and $\text{TCO}_{2(\text{sw})}$ (black dots) and calculated based on $\text{TA}_{(\text{sw})}^*$ and $\text{TCO}_{2(\text{sw})}^*$ (dashed red line), and (b) pH in the water column calculated based on $\text{TA}_{(\text{sw})}$ and $\text{TCO}_{2(\text{sw})}$ (black dots) and calculated based on $\text{TA}_{(\text{sw})}^*$ and $\text{TCO}_{2(\text{sw})}^*$ (dashed red line).

of the effect of ikaite precipitation in sea ice on ocean acidification is still in its infancy (e.g., Fransson et al., 2013).

Since the discovery of ikaite precipitation in sea ice (Dieckmann et al., 2008), research on its impact on the carbonate system of the underlying seawater has been ongoing. Depending on the timing and location of this precipitation within sea ice, the impact on the atmosphere and the water column in terms of CO_2 transport can be significantly different (Delille et al., 2014). Dissolution of ikaite within melting sea ice in the spring and export of this related high TA : TCO_2 ratio meltwater from the ice to the water column will decrease the $p\text{CO}_2$ and increase pH and $\Omega_{\text{aragonite}}$ of the surface layer seawater. Accordingly, during sea ice melt, an increase of $\Omega_{\text{aragonite}}$ in the surface water in the Arctic was observed (Chierici et al., 2011; Fransson et al., 2013; Bates et al., 2014). However, it was difficult to ascribe this increase to the legacy of excess TA in sea ice, ikaite dissolution or primary production.

The impact of ikaite precipitation on the surface seawater during sea ice growth is less clear. Fransson et al. (2013) suggested in winter in the Amundsen Gulf that the release of brine decreased $\Omega_{\text{aragonite}}$ by 0.8 at the sea ice–seawater interface as a result of ikaite precipitation within sea ice and the related CO_2 enrichment of brine. Conversely, dur-

ing ice melt, $\Omega_{\text{aragonite}}$ increased by 1.4 between March and May, likely due to both calcium carbonate dissolution and primary production. This contrasts with the present experiment. Figure 10 shows the evolution of $\Omega_{\text{aragonite}}$ and pH in the water column derived from $\text{TA}_{(\text{sw})}$ and $\text{TCO}_{2(\text{sw})}$ and the evolution of $\Omega_{\text{aragonite}}$ and pH predicted solely from salinity changes (i.e., using $\text{TA}_{(\text{sw})}^*$ and $\text{TCO}_{2(\text{sw})}^*$, noted as $\Omega_{\text{aragonite}}^*$ and pH^*). We used the CO2sys_v2.1.xls spreadsheet (Pierrot et al., 2006) with the dissociation constants from Goyet and Dickson (1989) and all other constants from DOE (1994). This shows the complexity of ikaite and its impact on the carbonate system and Ω in the underlying water.

During ice growth, sea ice brine rejection appears to increase both pH (from 8.00 to 8.06) and $\Omega_{\text{aragonite}}$ (from 1.28 to 1.65) of the underlying seawater, offsetting the effect of decreased temperature. A slight increase of $\Omega_{\text{aragonite}}$ was predicted due to increased salinity and a proportional increase of TA and TCO_2 as depicted in $\Omega_{\text{aragonite}}^*$. However, the effect of ikaite rejection and subsequent changes in TA strongly enhance the increase of $\Omega_{\text{aragonite}}$. Therefore, ikaite rejection from sea ice has a much stronger potential to increase $\Omega_{\text{aragonite}}$ than brine rejection during fall and winter sea ice growth, suggesting ikaite exported to seawater from sea ice may hamper the effect of oceanic acidification on $\Omega_{\text{aragonite}}$ in fall and in winter at the time when $\Omega_{\text{aragonite}}$ is at its minimum (Chierici and Fransson, 2009; Yamamoto-Kawai et al., 2009; Chierici et al., 2011). Ice formation may therefore delay harmful effects of ocean acidification on calcifying organisms by increasing $\Omega_{\text{aragonite}}$ in the critical winter period when $\Omega_{\text{aragonite}}$ reaches its minimal values. As a corollary, ice removal acts to alleviate the effect of ikaite rejection and may therefore lower $\Omega_{\text{aragonite}}$. This calls for an accounting of under-ice ikaite rejection in modeling predictions on the consequences of Arctic Ocean acidification in the context of northern hemispheric annual multiyear sea ice loss, as increased summer open water will lead to more first-year sea ice formation in fall and winter in the future.

5 Conclusion

We quantified the evolution of inorganic carbon dynamics from initial sea ice formation to its melt in a sea ice–seawater mesocosm pool from 11 to 29 January 2013. Based on our analysis of TA and TCO_2 in sea ice and seawater, the main processes affecting inorganic carbon within sea ice are ikaite precipitation and CO_2 exchange with the atmosphere, while in the underlying seawater, dissolution of ikaite was the main process affecting the inorganic carbon system.

During this experiment, sea ice exchanged inorganic carbon components (e.g., CO_2 , ikaite, TCO_2) with both the atmosphere and the underlying seawater. During sea ice growth, CO_2 was released to the atmosphere, while during ice melt, an uptake of atmospheric CO_2 was observed. We re-

port ikaite precipitation of up to $167 \mu\text{mol kg}^{-1}$ sea ice, similar to previous estimates from Rysgaard et al. (2014) based on microscopically observed values. In the underlying seawater, a net increase of $n\text{TA}_{(\text{sw})}$ over the whole experiment was observed (up to $128 \mu\text{mol kg}^{-1}$), suggesting that a portion of the ikaite crystals precipitated within sea ice was exported to the underlying seawater and then dissolved as the ice cover evolved in time. Ikaite export from ice to the underlying seawater was associated with brine rejection during sea ice growth, increased sea ice vertical connectivity due to the upward percolation of seawater and meltwater flushing during sea ice melt. Rysgaard et al. (2007) suggested that ikaite precipitation within sea ice could act as a significant sink for atmospheric CO_2 ; however to act as a sink for atmospheric CO_2 , ikaite crystals must remain in the ice structure while the CO_2 produced by their precipitation is expelled with dense brine rejection and entrained in deep seawater (Delille et al., 2014). TA changes observed in the water column once the sea ice started to melt indicate that more than half of the total amount of ikaite precipitated in the ice during the sea ice growth remained in the ice until the sea ice began to melt. Derivation of air–sea CO_2 fluxes related to the sea ice carbon pump should take into account ikaite export to the underlying ocean during sea ice growth, which might reduce the efficiency of oceanic CO_2 uptake upon sea ice melt. As sea ice melts, ikaite is flushed downward out of the ice along with the meltwater.

Ikaite export from sea ice and its dissolution had a strong impact on the underlying seawater. In this semi-closed system, sea ice growth increased the seawater salinity, $\text{TA}_{(\text{sw})}$ and $\text{TCO}_{2(\text{sw})}$. In spite of these increases, the $p\text{CO}_2$ of the underlying seawater remained undersaturated compared to the atmosphere. We conclude that ikaite dissolution within the water column is responsible for the seawater's continual $p\text{CO}_2$ undersaturation. In addition, we discuss that dissolution of ikaite crystals exported from sea ice in the underlying seawater can potentially hamper the effect of oceanic acidification on $\Omega_{\text{aragonite}}$ in fall and in winter in ice-covered areas at the time when $\Omega_{\text{aragonite}}$ is smallest.

6 Data availability

Data are available upon request from the authors.

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