

1 **CO<sub>2</sub> flux over young and snow-covered Arctic pack ice in**  
2 **winter and spring**

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37

38 **Abstract**

39

40 Rare CO<sub>2</sub> flux measurements from Arctic pack ice show that two types of ice are  
41 significant contributors to the release of CO<sub>2</sub> from ice to the atmosphere during winter  
42 and spring: young thin ice with thin layer of snow, and old (several weeks) snow  
43 covered thick ice. Young thin sea ice is characterized by high salinity and then porosity  
44 and thin layer of snow. Snow covered thick ice can remain relatively warm ( $>-7.5^{\circ}\text{C}$ )  
45 due to a thick insulating snow cover despite air temperatures were as low as  $-40^{\circ}\text{C}$ .  
46 Brine volume fractions of these two ice type are therefore high enough to provide  
47 favorable conditions for gas exchange between sea ice and the atmosphere even in mid-  
48 winter. Although the potential CO<sub>2</sub> flux from sea ice decreased due to the presence of  
49 the snow, the snow surface is still a CO<sub>2</sub> source to the atmosphere for low snow density  
50 and thin snow conditions. We found that young sea ice formed in leads, without snow  
51 cover, is the most effective in terms of CO<sub>2</sub> flux ( $+1.0 \pm 0.6 \text{ mmol C m}^{-2} \text{ day}^{-1}$ ) since  
52 the fluxes are an order of magnitude higher than for snow-covered older ice ( $+0.2 \pm 0.2$   
53  $\text{mmol C m}^{-2} \text{ day}^{-1}$ ).

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56

57 **1 Introduction**

58

59 Arctic sea ice is changing dramatically, with rapid declines in summer sea ice extent  
60 and a shift towards younger and thinner first-year ice rather than thick multi-year ice  
61 (e.g., Stroeve et al., 2012; Meier et al., 2014; Lindsay and Schweiger, 2015). Although  
62 the effects of sea ice formation and melting on biogeochemical cycles in the ocean have  
63 previously been discussed (e.g., Vancoppenolle et al., 2013), the effects of sea ice  
64 freezing and melting on the carbon dioxide (CO<sub>2</sub>) exchange with the atmosphere are  
65 still large unknowns (Parmentier et al., 2013).

66

67 Recent CO<sub>2</sub> flux measurements on sea ice indicate that sea ice is an active component in  
68 gas exchange between ocean and atmosphere (Nomura et al., 2013; Geilfus et al., 2013;  
69 2014; Delille et al., 2014; Brown et al., 2015; Kotovitch et al., 2016). The sea-ice CO<sub>2</sub>

70 fluxes depend on (a) the difference in the partial pressure of CO<sub>2</sub> (pCO<sub>2</sub>) between the  
71 sea ice surface and air, (b) brine volume fraction at the ice-snow interface, (c) ice  
72 surface condition including the snow deposited on ice, and (d) wind-driven pressure  
73 pumping through the snow. For (a), it is known that the air–sea ice CO<sub>2</sub> flux is driven  
74 by the differences in pCO<sub>2</sub> between the sea ice surface and atmosphere (e.g. Delille et  
75 al., 2014; Geilfus et al., 2014). The brine pCO<sub>2</sub> changes due to processes within the sea  
76 ice, such as thermodynamic process (e.g., Delille et al., 2014), biological activity (e.g.,  
77 Delille et al., 2007; Fransson et al., 2013; Rysgaard et al., 2013), and calcium carbonate  
78 (CaCO<sub>3</sub>; ikaite) formation and dissolution (e.g., Papadimitriou et al., 2012). When the  
79 pCO<sub>2</sub> in the brine is higher than that of the air pCO<sub>2</sub>, brine has the potential to release  
80 CO<sub>2</sub> to the atmosphere. Brine volume fraction (b) controls permeability of sea ice  
81 (Golden et al. 1998) and then CO<sub>2</sub> fluxes (Delille et al. 2014; Geilfus et al 2014). The  
82 air–sea ice CO<sub>2</sub> flux is strongly dependent on the sea ice surface conditions (c) (Nomura  
83 et al., 2010a, 2013; Geilfus et al., 2013; 2014; Barber et al., 2014; Brown et al., 2015;  
84 Fransson et al., 2015). Nomura et al. (2013) proposed that snow conditions (e.g., water  
85 equivalent) are important factors affecting gas exchange processes on sea ice. In  
86 addition, frost flowers promote CO<sub>2</sub> flux from the ice to the atmosphere (Geilfus et al.,  
87 2013; Barber et al., 2014; Fransson et al., 2015). For (d), it is thought that for snow  
88 cover, the CO<sub>2</sub> flux is affected by wind pumping (Massman et al., 1995; Takagi et al.,  
89 2005) in which the magnitude of CO<sub>2</sub> flux through snow or overlying soil (e.g., Takagi  
90 et al., 2005) increases due to wind pumping and can increase the transport relative to  
91 molecular diffusion by up to 40% (Bowling and Massman, 2011). These results were  
92 mainly found over land-based snow (soil and forest), and thus these processes are not  
93 well understood over sea ice (Papakyriakou and Miller, 2011).

94

95 In addition to the processes described above, the CO<sub>2</sub> flux over sea ice may also be  
96 influenced by the temperature difference between the ice surface and the atmosphere.  
97 This has been shown in previous studies in dry snowpacks over land surfaces. These  
98 studies show that there is an unstable air density gradient due to heating at the bottom  
99 producing a strong temperature difference between bottom and top of snow (e.g.,  
100 Powers et al., 1985; Severinghaus et al., 2010). This produces air flow within the  
101 snowpack, which is a potentially significant contributor to mixing and transport of gas

102 and heat within the snowpack. We expect that this process would also occur in snow  
103 over sea ice, especially during the wintertime when air temperatures are coldest and the  
104 temperature difference between sea ice surface (snow bottom) and atmosphere is largest  
105 (e.g., Massom et al., 2001). Generally, the sea ice surface under thick snow cover is  
106 warm due to the heat conduction from the bottom of sea ice and the insulation effect of  
107 the snow cover, and a strong temperature difference between sea ice surface and  
108 atmosphere is observed (e.g., Massom et al., 2001). Such a temperature difference  
109 would produce an unstable air density gradient and upward transport of air containing  
110 CO<sub>2</sub> degassed at the sea-ice surface, thereby enhancing CO<sub>2</sub> exchange between sea ice  
111 and atmosphere.

112

113 In the ice covered Arctic Ocean, storm periods, with high wind speeds and open leads  
114 are important for air-to-sea CO<sub>2</sub> fluxes (Fransson et al., 2017), due to the under-  
115 saturation of the surface waters in CO<sub>2</sub> with respect to the atmosphere. On the other  
116 hand, the subsequent ice growth and frost flowers formation in these leads promote ice-  
117 to-air CO<sub>2</sub> fluxes in winter (e.g. Barber et al., 2014). Given the fact that Arctic sea ice is  
118 shrinking and shifting from multi-year ice to first-year ice, the area of open ocean and  
119 thinner seasonal ice is increasing. Therefore, the contribution of open ocean/thinner sea  
120 ice surface to the overall CO<sub>2</sub> fluxes of the Arctic Ocean is potentially increasing.  
121 However, due to the difficulty in acquiring observations over the winter pack ice, most  
122 of the winter CO<sub>2</sub> flux measurements were examined over the Arctic landfast ice.  
123 Therefore, there is a definite lack of information on conditions during wintertime,  
124 especially from Arctic pack ice.

125

126 The Norwegian young sea ICE (N-ICE2015) campaign in winter and spring 2015  
127 provided opportunities to examine CO<sub>2</sub> fluxes between sea ice and atmosphere in a  
128 variety of snow and ice conditions in pack ice north of Svalbard. Formation of leads and  
129 their rapid refreezing allowed us to examine air–sea ice CO<sub>2</sub> fluxes over thin young sea  
130 ice, occasionally covered with frost flowers in addition to the snow-covered older ice  
131 that covers most of the pack ice area. The objectives of this study were to understand  
132 the effects of i) thin sea ice and frost flowers formations on the air–sea ice CO<sub>2</sub> flux in  
133 leads, ii) effect of snow-cover on the air–sea ice CO<sub>2</sub> flux over thin, young ice in the

134 Arctic Ocean during winter and spring seasons, and iii) of the effect of the temperature  
135 difference between sea ice and atmosphere (including snow cover) on the air–sea ice  
136 CO<sub>2</sub> flux.

137

138

## 139 **2 Materials and Methods**

140

### 141 **2.1 Study area**

142

143 This study was performed during N-ICE2015 campaign with R/V Lance in the pack ice  
144 north of Svalbard from January to June 2015 (Granskog et al., 2016). Air–sea ice CO<sub>2</sub>  
145 flux measurements were carried out from January to May 2015 during the drift of floes  
146 1, 2, and 3 of the N-ICE2015 campaign (Figures 1 and 2, Table 1). The ice pack was a  
147 mixture of young ice, first-year ice and second-year ice (Granskog et al., 2017), the two  
148 latter with a thick snow cover (Merkouriadi et al., 2017; Rösel et al., 2018). Air–sea ice  
149 CO<sub>2</sub> flux measurements were done over young ice (YI stations), first-year ice (FI  
150 stations), and old ice (multi-year ice) (OI station). In the N-ICE2015 study region modal  
151 ice thickness was about 1.3–1.5 m and modal snow thickness was about 0.5 m (Rösel et  
152 al., 2018). Formation of leads and their rapid refreezing provided us the opportunity to  
153 examine air–sea ice CO<sub>2</sub> fluxes over thin sea ice, occasionally covered with frost  
154 flowers at station YI1 (Figure 2 and Table 1). Air temperature and wind speed were  
155 measured at a 10 m weather mast on the ice floe installed about 400 m away from R/V  
156 Lance (Cohen et al., 2017).

157

158

### 159 **2.2 CO<sub>2</sub> flux measurements**

160

161 The air–sea ice CO<sub>2</sub> flux was measured with LI-COR 8100-104 chambers connected to  
162 a LI-8100A soil CO<sub>2</sub> flux system (LI-COR Inc., USA) (Figure 2). This enclosed  
163 chamber method has been widely applied over snow and sea ice (e.g., Schindlbacher et  
164 al., 2007, Geilfus et al., 2015). Two chambers were connected in a closed loop to the  
165 infrared gas analyzer (LI-8100A, LI-COR Inc., USA) to measure CO<sub>2</sub> concentration

166 through the multiplexer (LI-8150, LI-COR Inc., USA) with an air pump rate at 3 L min<sup>-1</sup>.  
167 <sup>1</sup>. Power was supplied by a car battery (8012-254, Optima Batteries Inc., USA). Four  
168 CO<sub>2</sub> standards (324–406 ppmv) traceable to the WMO scale (Inoue and Ishii, 2005)  
169 were prepared to calibrate the CO<sub>2</sub> gas analyzer prior to the observations. CO<sub>2</sub> flux was  
170 measured in the morning or in the afternoon during low-wind conditions (Table 2), to  
171 minimize the effect of wind on the flux (Bain et al., 2005).

172

173 One chamber was installed over undisturbed snow or frost flowers over the ice surface.  
174 The chamber collar was inserted 5 cm into the snow and 1 cm into ice at frost flowers  
175 site to avoid air leaks between inside and outside of chamber. The second chamber was  
176 installed on bulk sea ice after removing the snow or frost flowers. Flux measurements  
177 was begun immediately in order to minimize the changes of the ice surface condition. In  
178 order to evaluate the effect of removing snow on sea ice surface temperature, ice surface  
179 temperature was monitored during CO<sub>2</sub> flux measurements at station FI6. To measure  
180 the sea ice surface temperature, temperature sensor (RTR 52, T & D Corp., Japan) was  
181 installed in the top of the ice (1 cm) surface after snow removal. During first CO<sub>2</sub> flux  
182 measurements (about 30 minutes), ice surface temperature was stable at  $-5.8^{\circ}\text{C}$ ,  
183 suggesting that the effect of removing snow on the variation of sea ice surface  
184 temperature was negligible within 30 minutes. The ice surface temperature decreased  
185 from  $-5.8^{\circ}\text{C}$  to  $-8.0^{\circ}\text{C}$  at 200 minutes after removal of snow. Therefore, in this paper,  
186 the data of the initial 30 minutes of CO<sub>2</sub> flux measurement after removal of snow or  
187 frost flowers was used. The chamber was closed for 20 minutes in a sequence. The 20-  
188 minute time period was used because CO<sub>2</sub> fluxes over sea ice are much smaller than  
189 over land. The CO<sub>2</sub> concentrations within the chamber were monitored to ensure that  
190 they changed linearly throughout the measurement period (example given in Figure 3).  
191 The CO<sub>2</sub> flux ( $\text{mmol C m}^{-2} \text{ day}^{-1}$ ) (positive value indicates CO<sub>2</sub> being released from ice  
192 surface to air) was calculated based on the changes of the CO<sub>2</sub> concentration within the  
193 headspace of the chamber with LI-COR software (Model: LI8100PC Client v.3.0.1.).  
194 The mean coefficient of variation for CO<sub>2</sub> flux measurements was less than 3.0% for  
195 CO<sub>2</sub> flux values larger than  $\pm 0.1 \text{ mmol C m}^{-2} \text{ day}^{-1}$ . For CO<sub>2</sub> flux values smaller than  
196  $\pm 0.1 \text{ mmol C m}^{-2} \text{ day}^{-1}$ , the mean coefficient of variation for CO<sub>2</sub> flux measurements

197 was higher than 3.0%, suggesting that the detection limit of this system is about 0.1  
198 mmol C m<sup>-2</sup> day<sup>-1</sup>.

199

200 In this paper, we express the CO<sub>2</sub> flux measured over the snow and frost flowers as  
201 F<sub>snow</sub> and F<sub>ff</sub>, respectively, and the flux measured directly over the sea ice surface either  
202 on snow-free ice or after removal of snow and frost flowers as F<sub>ice</sub>. F<sub>snow</sub> and F<sub>ff</sub> are the  
203 natural flux (snow and frost flowers are part of the natural system), and F<sub>ice</sub> is the  
204 potential flux in cases when snow or frost flowers are removed. While removal of snow  
205 and frost flowers is an artificial situation, comparisons between F<sub>ice</sub> and F<sub>snow</sub> or F<sub>ff</sub>  
206 provide information about the effect of snow on the CO<sub>2</sub> flux. Therefore, in this study,  
207 we examine both situations for CO<sub>2</sub> flux.

208

209

### 210 **2.3 Sampling of snow, frost flowers, brine, and sea ice**

211

212 For salinity measurements, separate samples were taken for snow only, snow and frost  
213 flowers, and sea ice surface scrapes. The samples were taken using a plastic shovel,  
214 placed into plastic bags and stored in an insulated box for transport to the ship-lab for  
215 further processing. Samples were melted slowly (2–3 days) in the dark at +4°C. The  
216 temperature of the snow and frost flowers samples were measured during CO<sub>2</sub> flux  
217 measurements (approximately 60 minutes after the onset of the CO<sub>2</sub> flux measurement)  
218 using a needle-type temperature sensor (Testo 110 NTC, Brandt Instruments, Inc.,  
219 USA). The accuracy of this sensor is ±0.2°C. Snow density was obtained using a fixed  
220 volume sampler (Climate Engineering, Japan) and weight measurement. The depth of  
221 the snow pack and frost flowers was also recorded using a ruler.

222

223 Brine was also collected at stations FI3–6 for salinity, dissolved inorganic carbon (DIC)  
224 and total alkalinity (TA) measurements. Brine was collected from sackholes as  
225 described in Gleitz et al. (1995). The sackholes were drilled using a 9 cm diameter ice  
226 corer (Mark II coring system, Kovacs Enterprises, Inc., USA) to a depth of 30 cm. The  
227 sackholes were then covered with a lid of 5 cm-thick urethane to reduce heat and gas  
228 transfer between brine and atmosphere. When brine accumulated at the bottom of the



229 sackholes (approximately 15 minutes), it was collected with a plastic syringe (AS ONE  
230 Corporation, Japan) and kept in 500 mL unbreakable plastic bottles (I-Boy, AS ONE  
231 Corporation, Japan) in order to facilitate safe transport to the sampling sites in cold and  
232 harsh conditions. The brine bottles were filled without head-space and immediately  
233 stored in an insulated box to prevent freezing. Immediately after return to the ship, the  
234 brine samples were transferred to 250 mL borosilicate bottles (DURAN Group GmbH,  
235 Germany) for DIC and TA measurements using tubing to prevent contact with air. The  
236 samples were preserved with saturated mercuric chloride ( $\text{HgCl}_2$ , 60  $\mu\text{L}$  for a 250 mL  
237 sample) and stored in the dark at  $+10^\circ\text{C}$  until analyses was performed at the Institute of  
238 Marine Research, Norway.

239

240 Sea ice was collected by same ice corer as described for brine collection and at the same  
241 location as snow and frost flowers were collected. Sea ice temperature was measured by  
242 same sensor as described for snow. For the ice cores, the temperature sensor was  
243 inserted in small holes drilled into the core. The core was then cut with a stainless steel  
244 saw into 10 cm sections and stored in plastic bags for subsequent salinity measurements.  
245 The ice core sections were kept at  $+4^\circ\text{C}$  and melted in the dark prior to measurement.

246

247

## 248 **2.4 Sample analysis**

249

250 Salinities for melted snow, frost flowers, sea ice, and brine were measured with a  
251 conductivity sensor (Cond 315i, WTW GmbH, Germany). For calibration of salinity  
252 measurement, a Guildline PORTASAL salinometer Model 8410A, standardized by  
253 International Association for the Physical Sciences of the Oceans (IAPSO) standard  
254 seawater (Ocean Scientific International Ltd, UK) was used. Accuracy of this sensor  
255 was  $\pm 0.003$ .

256

257 Analytical methods for DIC and TA determination are fully described in Dickson et al.  
258 (2007). DIC in brine was determined using gas extraction of acidified sample followed  
259 by coulometric titration and photometric detection using a Versatile Instrument for the  
260 Determination of Titration carbonate (VINDTA 3C, Germany). TA of brine was

261 determined by potentiometric titration of 40 mL sample in open cell with 0.05 N  
262 hydrochloric acid using a Titrino system (Metrohm, Switzerland). The average standard  
263 deviation for DIC and TA, determined from replicate sample analyses from one sample,  
264 was within  $\pm 2 \mu\text{mol kg}^{-1}$  for both DIC and TA. Accuracy of the DIC and TA  
265 measurements were  $\pm 2 \mu\text{mol kg}^{-1}$  for both DIC and TA estimated using Certified  
266 Reference Materials (CRM, provided by A. G. Dickson, Scripps Institution of  
267 Oceanography, USA). The  $\text{pCO}_2$  of brine ( $\text{pCO}_{2\text{b}}$ ) was derived from in situ temperature,  
268 salinity, DIC and TA of brine using the carbonate speciation program CO2SYS (Pierrot  
269 et al., 2006). We used the carbonate dissociation constants ( $K_1$  and  $K_2$ ) of Mehrbach et  
270 al. (1973) as refit by Dickson and Millero (1987), and the  $\text{KSO}_4$  determined by Dickson  
271 (1990). The conditional stability constants used to derived  $\text{pCO}_2$  are strictly only valid  
272 for temperatures above  $0^\circ\text{C}$  and salinities between 5 and 50. Studies in spring ice  
273 indicated that seawater thermodynamic relationships may be acceptable in warm and  
274 low-salinity sea ice (Delille et al., 2007). In sea ice brines at even moderate brine  
275 salinities of 80, Brown et al. (2014) found that measured and calculated values of the  
276  $\text{CO}_2$  system parameters can differ by as much as 40%. On the other hand, because the  
277  $\text{CO}_2$  system parameters are much more variable in sea ice than in seawater, sea ice  
278 measurements demand less precision than those in seawater. Fransson et al. (2015)  
279 performed one of few detailed analyses of the internal consistency using four sets of  
280 dissociation constants and found that the deviation between measured and calculated  
281 DIC varied between  $\pm 6$  and  $\pm 11 \mu\text{mol kg}^{-1}$ , respectively. This error in calculated DIC  
282 was considered insignificant in relation to the natural variability in sea ice.

283

284 The  $\text{pCO}_2$  of atmosphere was calculated from  $\text{CO}_2$  concentration (ppmv) at Ny-Ålesund,  
285 Svalbard (<http://www.esrl.noaa.gov/gmd/dv/iadv/>) taking into account saturated water  
286 vapor and atmospheric pressure during sampling day.

287

288 The water equivalent was computed for snow by multiplying snow thickness by snow  
289 density (Jonas et al., 2009). Brine volume of sea ice was calculated from the  
290 temperature and salinity of sea ice according to Cox and Weeks (1983) and Petrich and  
291 Eicken (2010).

292

293

## 294 **3 Results**

295

### 296 **3.1 Air temperature**

297

298 Air temperature is shown in Figure 4. During the study period, air temperature varied  
299 significantly from a low of  $-41.3^{\circ}\text{C}$  (30 January) to a high of  $+1.7^{\circ}\text{C}$  (15 June) (Hudson  
300 et al., 2015). Even in wintertime (from January to March), rapid increases of air  
301 temperature from below  $-30^{\circ}\text{C}$  up to  $-0.2^{\circ}\text{C}$  (e.g., 18 February), were observed. In  
302 springtime (from April to June), the air temperature increased continuously, and from 1  
303 June, air temperatures were near  $0^{\circ}\text{C}$ , although rapid increases (and subsequent  
304 decreases) of air temperature to near  $0^{\circ}\text{C}$  were observed on two occasions in mid-May  
305 (Cohen et al., 2017).

306

307

### 308 **3.2 Characteristics of snow, sea ice, and frost flowers**

309

310 The snow and ice thickness at the observation sites ranged between 0.0 and 60.0 cm and  
311 between 15.0 and  $>200$  cm, respectively (Table 1). The thin snow and ice represent  
312 newly formed ice in leads at station YI1. The thickness of the frost flowers ranged from  
313 1.0 to 2.5 cm.

314

315 Figure 5 shows vertical profiles of snow and ice temperature and salinity in the top 20  
316 cm of ice. Temperatures within the snowpack depended on the air temperature at the  
317 time of observation. However, the bottom of the snow and the surface of the sea ice  
318 were relatively warm ( $T > -7.5^{\circ}\text{C}$ ), except for the frost flowers station YI1 and the multi-  
319 year ice station OI1 (Figure 5a and Table 2). High salinities ( $S > 18.6$ ) characterized the  
320 bottom of the snow and the surface of the sea ice, except for the multi-year ice station  
321 OI1 (Figure 5b). At the multi-year ice station OI1, salinity was zero through the snow  
322 and top of sea ice. Salinity of frost flowers was up to 92.8 for the thin ice station YI1  
323 (Figure 5b). Snow density and water equivalent ranged from 268 to  $400\text{ kg m}^{-3}$  and 11  
324 to  $180\text{ kg m}^{-2}$ , respectively.

325

326

### 327 **3.3 Physical and chemical properties of brine**

328

329 The brine volume fraction, temperature, salinity, DIC, TA, and calculated pCO<sub>2</sub> are  
330 summarized in Table 2. Brine volume fraction in top 20 cm of ice was from 9 to 17%,  
331 except for the value of 0% at the multi-year ice station OI1 (Table 2). Brine  
332 temperatures and salinity ranged from -5.3 to -3.3°C and 51.8 to 86.6, respectively.  
333 DIC and TA of brine ranged from 3261 to 4841 μmol kg<sup>-1</sup> and 3518 to 5539 μmol kg<sup>-1</sup>,  
334 respectively. The pCO<sub>2</sub> of brine (pCO<sub>2b</sub>) (334–693 μatm) was generally higher than  
335 that of atmosphere (pCO<sub>2a</sub>) (401 ± 7 μatm), except for station FI4.

336

337

### 338 **3.4 CO<sub>2</sub> flux**

339

340 Table 3 summarizes the CO<sub>2</sub> flux measurements for each surface condition. For  
341 undisturbed natural surface conditions, i.e. measurements directly on the snow surface  
342 (F<sub>snow</sub>) or the frost flowers (F<sub>ff</sub>) on young ice, the mean CO<sub>2</sub> flux was +0.2 ± 0.2 mmol  
343 C m<sup>-2</sup> day<sup>-1</sup> for F<sub>snow</sub> and +1.0 ± 0.6 mmol C m<sup>-2</sup> day<sup>-1</sup> for F<sub>ff</sub>. The potential flux in  
344 cases when snow or frost flowers had been removed (F<sub>ice</sub>) was +2.5 ± 4.3 mmol C m<sup>-2</sup>  
345 day<sup>-1</sup>. The air–sea ice CO<sub>2</sub> fluxes measured over the ice surface (F<sub>ice</sub>) increased with  
346 increasing difference in pCO<sub>2</sub> between brine and atmosphere (ΔpCO<sub>2b-a</sub>) with  
347 significant correlation (R<sup>2</sup> = 0.9, p < 0.02), but this was not the case for F<sub>snow</sub> (R<sup>2</sup> = 0.0,  
348 p < 0.96) (Figure 6).

349

350

351

## 352 **4 Discussion**

353

### 354 **4.1 Effect of snow cover on the physical properties of sea ice surface**

355

356 In this study, we examined CO<sub>2</sub> fluxes between sea ice and atmosphere in a variety of  
357 air temperature conditions from  $-32$  to  $-3^{\circ}\text{C}$  and diverse snow and ice conditions (Table  
358 2). The bottom of the snow pack and the surface of the sea ice remained relatively warm  
359 ( $>-7.5^{\circ}\text{C}$ ) (Figure 5a, Table 2), except for stations OI1 and YI1, even though air  
360 temperature was sometimes below  $-40^{\circ}\text{C}$  (Figure 4). Relatively warm ice temperatures  
361 were likely due to the upward heat transport from the bottom of the ice and in some  
362 cases the thick insulating snow cover, except for stations OI1 and YI1 (Table 2).  
363 Therefore, snow acted as thermal insulator over sea ice, and in general the snow depths  
364 observed during N-ICE2015 point towards this being representative for first-year and  
365 second-year or older ice in the study region in winter 2015 (Rösel et al., 2018). The  
366 young and first-year ice surfaces were characterized by high salinities (Figure 5b).  
367 During sea ice formation, upward brine transport to the snow pack occurs (e.g., Toyota  
368 et al., 2011). In addition, brine within the sea ice was not completely drained as  
369 compared to that of multi-year ice. Furthermore, formation of frost flowers and  
370 subsequent wicking up of surface brine into the frost flowers also provides high salinity  
371 at the surface of sea ice (Kaleschke et al., 2004; Geilfus et al., 2013; Barber et al., 2014;  
372 Fransson et al., 2015) as observed in this study ( $S>92$ ) (Figure 5b). Snowfall over the  
373 frost flowers would have preserved the high salinity at the bottom of snow pack and top  
374 of sea ice for young and first-year ice.

375

376 As a result of the combination of the relatively high temperature and high salinity at the  
377 top of sea ice, brine volume fractions in the upper parts of the sea ice were high, up to  
378 17% (Table 2). It has been shown that ice permeability increases by an order of  
379 magnitude when brine volume fraction  $> 5\%$ , which would correspond to a temperature  
380 of  $-5^{\circ}\text{C}$  for a bulk ice salinity of 5 – the so called “law of fives” (Golden et al., 1998;  
381 Pringle et al., 2009; Zhou et al., 2013). Because sea ice temperatures were low and  
382 thereby reduced permeability in winter season, generally, air–sea ice CO<sub>2</sub> flux is at its  
383 minimum in the winter (e.g., Delille et al., 2014). However, in our study, the brine  
384 volume fractions were generally  $>9\%$ , except for station OI1 with fresh ice at the  
385 surface, providing conditions for active gas exchange within sea ice and between sea ice  
386 and atmosphere. This situation was likely made possible due to the thick snow cover  
387 and relatively thin and young sea ice.

388

389

## 390 **4.2 CO<sub>2</sub> fluxes over different sea-ice surface types**

391

392 The CO<sub>2</sub> flux measurements over different surface conditions indicate that the snow  
393 cover over sea ice affects the magnitude of air–sea ice CO<sub>2</sub> flux, especially for stations  
394 FI5 and FI6 (Table 3). For undisturbed natural surface conditions, the CO<sub>2</sub> flux  
395 measured directly over snow-covered first-year ice and young ice with frost flowers  
396 ( $F_{\text{snow}}$  and  $F_{\text{ff}}$ ) was lower in magnitude than that for potential flux obtained directly over  
397 the ice surface after removing snow ( $F_{\text{ice}}$ ), especially for stations FI5, FI6, and YI1.

398

399  $F_{\text{ff}}$  indicates that the frost flowers surface on young thin ice is a CO<sub>2</sub> source to the  
400 atmosphere and  $F_{\text{ff}}$  was higher than  $F_{\text{snow}}$ , except for station FI1. Frost flowers are  
401 known to promote gas flux, such as CO<sub>2</sub>, from the sea ice to the atmosphere (Geilfus et  
402 al., 2013; Barber et al., 2014; Fransson et al., 2015). At multi-year ice station OI1,  
403 neither snow or ice surface acted as a CO<sub>2</sub> source/sink. The surface of multi-year ice did  
404 not contain any brine (Figure 5b and Table 2), and the top of the ice was clear, colorless  
405 and very hard, suggesting superimposed formation at the top of sea ice. This situation  
406 would be similar as for freshwater-ice and superimposed-ice as these non-porous media  
407 block gas exchange effectively at the sea ice surface (Delille et al., 2014). Snow-ice and  
408 superimposed-ice were frequently found in second-year ice cores during N-ICE2015  
409 (Granskog et al., 2017), so the ‘blocking’ of gas exchange in second-year and multi-  
410 year ice may be a widespread process in the Arctic.

411

412 The magnitude of positive  $F_{\text{snow}}$  is less than  $F_{\text{ice}}$  for stations FI5 and FI6 (Table 3)  
413 indicating that the potential CO<sub>2</sub> flux from sea ice decreased due to the presence of  
414 snow. Previous studies have shown that snow accumulation over sea ice effectively  
415 impede CO<sub>2</sub> exchange (Nomura et al., 2013; Brown et al., 2015). Nomura et al. (2013)  
416 reported that 50–90% of the potential CO<sub>2</sub> flux was reduced due to the presence of  
417 snow/superimposed-ice at the water equivalent of 57–400 kg m<sup>-2</sup>, indicating that the  
418 snow properties are an important factor that controls the CO<sub>2</sub> exchange through a  
419 snowpack. Comparisons between stations FI5 and FI6 for  $F_{\text{snow}}/F_{\text{ice}}$  ratio (0.2 for FI5

420 and 0.0 for FI6) and water equivalent (11 kg m<sup>-2</sup> for FI5 and 127 kg m<sup>-2</sup> for FI6)  
421 indicate that the potential CO<sub>2</sub> flux is reduced (80% for FI5 and 98% for FI6 of the  
422 potential CO<sub>2</sub> flux) with increasing water equivalent. Although the magnitude of the  
423 potential CO<sub>2</sub> flux through the sea ice surface decreased by the presence of snow for  
424 stations FI5 and FI6 (Table 3), the snow surface still presents a CO<sub>2</sub> source to the  
425 atmosphere for low snow density and shallow depth conditions (e.g., +0.6 mmol C m<sup>-2</sup>  
426 day<sup>-1</sup> for FI5).

427

428 For F<sub>ice</sub>, there were negative CO<sub>2</sub> fluxes at stations FI3 and FI4 (-0.6 mmol C m<sup>-2</sup> day<sup>-1</sup>  
429 for FI3 and -0.8 mmol C m<sup>-2</sup> day<sup>-1</sup> for FI4) (Table 3). These fluxes corresponded to low  
430 or negative  $\Delta p\text{CO}_2_{b-a}$  as compared to that in atmosphere (Table 2 and Figure 6).  
431 Negative CO<sub>2</sub> fluxes should correspond to negative  $\Delta p\text{CO}_2_{b-a}$ . Therefore, the  
432 uncertainty for the calculation of carbonate chemistry may be one reason for the  
433 discrepancy in pCO<sub>2</sub> calculation in these conditions (Brown et al., 2014).

434

435

### 436 **4.3 Comparison to earlier studies on sea-ice to air CO<sub>2</sub> flux**

437

438 The CO<sub>2</sub> fluxes measured over the undisturbed natural surface conditions (F<sub>snow</sub> and F<sub>ff</sub>)  
439 in this study ranged from +0.1 to +1.6 mmol C m<sup>-2</sup> day<sup>-1</sup> (Table 3), which are at the  
440 lower end of the reported range based on the chamber method and eddy covariance  
441 method for natural and artificial sea ice (-259.2 to +74.3 mmol C m<sup>-2</sup> day<sup>-1</sup>)  
442 (Zemmelink et al., 2006; Nomura et al., 2006, 2010a, 2010b, 2013; Miller et al., 2011;  
443 Papakyriakou and Miller, 2011; Geilfus et al., 2012, 2013, 2014; Barber et al., 2014;  
444 Delille et al., 2014; Sørensen et al., 2014; Brown et al., 2015; Kotovitch et al., 2016).  
445 Direct comparison to previous studies is complicated because CO<sub>2</sub> flux measurements  
446 with both chamber and eddy covariance techniques were used during different condition  
447 for season and ice surface characteristics. In addition, discrepancies between chamber  
448 and eddy covariance measurements of air-ice CO<sub>2</sub> fluxes have been repeatedly observed.  
449 The footprint size of CO<sub>2</sub> exchange measured with the two approaches (Zemmelink et  
450 al., 2006, 2008; Burba et al., 2008; Amiro, 2010; Miller et al., 2011; Papakyriakou and  
451 Miller, 2011; Sørensen et al., 2014; Miller et al., 2015) may be one reason for the large

452 difference. The eddy covariance method reflects a flux integrated over a large area, that  
453 can contain several different surface types. Therefore, eddy-covariance appears to be  
454 more useful for understanding fluxes at large spatial and temporal scales. On the other  
455 hand, the chamber method reflects the area where chamber was covered, and it is useful  
456 for understanding the relationship between fluxes and ice surface conditions on smaller  
457 scales. The different spatial scales of the two methods may be therefore one reason for  
458 the discrepancy in CO<sub>2</sub> flux measurements.

459

460 Comparison of the natural CO<sub>2</sub> flux range (+0.1 to +1.6 mmol C m<sup>-2</sup> day<sup>-1</sup> for F<sub>snow</sub> and  
461 F<sub>fr</sub>) (Table 3) with previous estimates derived from the chamber method (-5.2 to +6.7  
462 mmol C m<sup>-2</sup> day<sup>-1</sup>) (Nomura et al., 2006, 2010a, 2010b, 2013; Geilfus et al., 2012,  
463 2013; 2014; Barber et al., 2014; Delille et al., 2014; Brown et al., 2015; Kotovitch et al.,  
464 2016) (these studies include both natural and potential fluxes) shows that CO<sub>2</sub> fluxes  
465 during NICE2015 experiment are at the lower end of positive values. However, our  
466 potential CO<sub>2</sub> flux (F<sub>ice</sub>) was a larger CO<sub>2</sub> source (up to +11.8 mmol C m<sup>-2</sup> day<sup>-1</sup>) than  
467 reported in previous studies (+6.7 mmol C m<sup>-2</sup> day<sup>-1</sup>). In our study, the maximum  
468 potential flux (e.g., +11.8 mmol C m<sup>-2</sup> day<sup>-1</sup>) was obtained for F<sub>ice</sub> at station FI6 (Table  
469 3). In this situation, ΔpCO<sub>2 b-a</sub> (293 μatm) was the highest (Table 2 and Figure 6), and it  
470 is reasonable to consider this as the highest magnitude of positive CO<sub>2</sub> flux within our  
471 study. However, a previous study by closed chamber method showed that even for a  
472 similar ΔpCO<sub>2 b-a</sub> (297 μatm) and magnitude for the brine volume fraction (10–15%),  
473 the CO<sub>2</sub> flux was +0.7 mmol C m<sup>-2</sup> day<sup>-1</sup> for artificial sea ice with no snow in the tank  
474 experiment (Nomura et al., 2006).

475

476 The CO<sub>2</sub> flux between the sea ice and overlying air can be expressed by the following  
477 equation,

478

$$479 F_{\text{CO}_2} = r_b k \alpha \Delta p\text{CO}_{2 \text{ b-a}},$$

480

481 where  $r_b$  is the ratio of surface of the brine channel to sea ice surface, and we assume  
482 that the value of  $r_b$  is equal to brine volume fraction,  $k$  is the gas transfer velocity,  $\alpha$   
483 is the solubility of CO<sub>2</sub> (Weiss, 1974), and  $\Delta p\text{CO}_{2 \text{ b-a}}$  is the difference in pCO<sub>2</sub> between



484 brine and atmosphere. The equation is based on the fact that CO<sub>2</sub> transfer between  
485 seawater and air is controlled by processes in the near-surface water (Liss, 1973). The  
486 gas transfer velocity ( $k$ ) calculated from  $F$ ,  $r_b$ ,  $\alpha$  and  $\Delta p\text{CO}_2_{b-a}$  was  $5.12 \text{ m day}^{-1}$  for  $F_{ice}$   
487 at station FI6 and  $0.29 \text{ m day}^{-1}$  for the tank experiment examined in Nomura et al.  
488 (2006). This result clearly indicates that the gas transfer velocity for  $F_{ice}$  at station FI6 is  
489 higher than that of tank experiment examined in Nomura et al. (2006) even with very  
490 similar  $\Delta p\text{CO}_2_{b-a}$  and brine volume fraction.

491

492 Here, we surmise that the gas transfer velocity and thereby CO<sub>2</sub> flux is greatly enhanced  
493 by the temperature difference between sea ice surface and atmosphere. Previous studies  
494 indicate that there is an unstable air density gradient in a dry snowpack due to basal  
495 heating and the strong temperature difference develops between bottom and top of snow  
496 (e.g., Powers et al., 1985; Severinghaus et al., 2010), which enhances the flow of air  
497 through the snowpack. We propose that the mixing and transport of gas within the  
498 snowpack could also occur over sea ice. Because temperatures at the bottom of snow  
499 and the top of sea ice were relatively warm due to a thick insulating snow over sea ice,  
500 there was a strong temperature difference between sea ice surface and atmosphere when  
501 air temperature was low (Figure 5a and Table 2). For station FI6, temperature difference  
502 between sea ice surface and atmosphere was  $20.2^\circ\text{C}$  after snow removal. On the other  
503 hand, in the tank experiment by Nomura et al. (2006), the temperature difference  
504 between sea ice surface (top 1.5 cm) and air in the headspace was only  $4.5^\circ\text{C}$ .

505

506 Figure 6 shows the relationship between mean air–sea ice CO<sub>2</sub> fluxes and temperature  
507 difference between ice and atmosphere. The strong dependence of CO<sub>2</sub> flux with  
508 temperature difference ( $T_{ice}-T_a$ ) was observed, especially for  $F_{ff}$  and  $F_{ice}$  ( $R^2 > 0.7$ ,  $p <$   
509  $0.01$ ) (Figure 6). Due to the high brine volume fractions (Table 2), sea ice surface had  
510 enough permeability for gas exchange. In addition, ice temperatures were similar for  
511 young and first-year ice (Figure 6, Table 2), indicating that  $p\text{CO}_2$  at the top of sea ice  
512 and CO<sub>2</sub> flux would be of similar order of magnitude if thermodynamic processes  
513 dominated. Therefore, our results suggest that the CO<sub>2</sub> fluxes even over the frost  
514 flowers as a natural condition, would be enhanced by the upward transport of air  
515 containing high CO<sub>2</sub> from the surface of sea ice to the atmosphere due to the strong

516 temperature difference between sea ice surface and atmosphere. Although the presence  
517 of snow on sea ice has potential to produce a larger temperature difference between sea  
518 ice surface and atmosphere and promote the upward transport, the magnitude of the CO<sub>2</sub>  
519 flux decreased due to the presence of snow. However, for young sea ice with frost  
520 flowers (e.g., station Y11), ice surface temperature was warm (Table 2), suggesting that  
521 CO<sub>2</sub> flux would be enhanced by the large temperature difference between sea ice  
522 surface and atmosphere.

523

524

525

## 526 **5 Conclusions**

527

528 We measured CO<sub>2</sub> fluxes along with sea ice and snow physical and chemical properties  
529 over first-year and young sea ice north of Svalbard in the Arctic pack ice. Our results  
530 suggest that young thin snow-free ice, with or without frost flowers, is a source of  
531 atmospheric CO<sub>2</sub> due to the high pCO<sub>2</sub> and salinity and relatively high sea ice  
532 temperature. Although the potential CO<sub>2</sub> flux from sea-ice surface decreased due to the  
533 presence of snow, snow surface still presents a modest CO<sub>2</sub> source to the atmosphere  
534 for low snow density and shallow depth situations. The highest ice to air fluxes were  
535 observed over thin young sea ice formed in leads. During N-ICE2015 the ice pack was  
536 dynamic, and formation of open water was associated with storms, where new ice was  
537 formed. The subsequent ice growth in these leads becomes important for the ice-to-air  
538 CO<sub>2</sub> fluxes in winter due to the fact that the flux from young ice is an order of  
539 magnitude larger than from snow-covered first-year and older ice.

540

541

542

## 543 **6 Data availability**

544

545 Data used in this paper will be available at Norwegian Polar Data Centre

546 (data.npolar.no).

547

548

549

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551

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818

819 **Figure captions**

820

821 Figure 1. Location map of the sampling area north of Svalbard during N-ICE2015.  
822 Image of the sea ice concentrations (a) and station map (b) were derived from Special  
823 Sensor Microwave Imager (SSM/I) satellite data for mean of February 2015 and from  
824 Sentinel-1 (Synthetic Aperture Radar Sensor) satellite data, respectively.

825

826 Figure 2. Photographs of the CO<sub>2</sub> flux chamber system at station YI1 north of Svalbard  
827 on Friday 13 March 2015. CO<sub>2</sub> flux chamber was installed over the frost flowers on the  
828 new thin ice in the refreezing lead.

829

830 Figure 3. Example of the temporal variation in CO<sub>2</sub> concentration ( $\Delta$ CO<sub>2</sub>) in the  
831 chambers installed at station YI1 that is use to calculate the CO<sub>2</sub> flux.  $\Delta$ CO<sub>2</sub> indicates  
832 the change in CO<sub>2</sub> concentration inside the chamber since the chamber was closed.

833

834 Figure 4. Time series of air temperature measured at the weather mast over the ice floe  
835 (10 m height) (Hudson et al., 2015). Blank period indicates no data. Colored symbols  
836 indicate the date for the chamber flux measurements. The horizontal dashed line  
837 indicates air temperature = 0°C.

838

839 Figure 5. Vertical profiles of temperature (a) and salinity (b) in snow and sea ice (top 20  
840 cm). The horizontal line indicates snow–ice interface. Shaded area indicates sea ice. The  
841 triangle in (a) indicates the air temperature for each station. For stations FI7 and YI2  
842 and 3, we have no salinity data.

843

844 Figure 6. Relationships between mean air–sea ice CO<sub>2</sub> fluxes and temperature  
845 difference between ice (T<sub>ice</sub>) and atmosphere (T<sub>a</sub>) (circle) and ice temperature (T<sub>ice</sub>)  
846 (top 20 cm) (cross) for F<sub>snow</sub> (blue), F<sub>ff</sub> (black) and F<sub>ice</sub> (red) for young and first-year sea  
847 ice. Relationships between mean air–sea ice CO<sub>2</sub> fluxes and the difference of pCO<sub>2</sub>  
848 ( $\Delta p\text{CO}_2_{b-a}$ ) between brine (pCO<sub>2 b</sub>) and atmosphere (pCO<sub>2 a</sub>) (triangle) for F<sub>snow</sub> (solid  
849 gray) and F<sub>ice</sub> (open gray).

850

851

## 852 **Table captions**

853

854 Table 1. Station, date for CO<sub>2</sub> flux measurement, position, floe number, surface  
855 condition, ice type and thickness of snow, frost flowers, and sea ice.

856

857 a. Sea ice coring and snow sampling was conducted on 5 March 2015.

858

859 b. Sea ice coring and snow sampling was conducted on 10 March 2015.

860

861

862 Table 2. Station, snow density and water equivalent, brine volume fraction, and  
863 temperature for sea ice (top 20 cm), brine temperature, salinity, DIC, TA, pCO<sub>2</sub> (pCO<sub>2</sub>  
864 b), and atmospheric temperature, wind speed, pCO<sub>2</sub> (pCO<sub>2 a</sub>)<sup>a</sup> and  $\Delta p\text{CO}_2_{b-a}$ .

865

866 a.  $p\text{CO}_2\text{ a}$  ( $\mu\text{atm}$ ) was calculated from  $\text{CO}_2$  concentration (ppmv) at Ny-Ålesund,  
867 Svalbard (<http://www.esrl.noaa.gov/gmd/dv/iadv/>) taking into account saturated water  
868 vapor and atmospheric pressure during sampling day.

869

870 b. Mean values for snow column.

871

872 c. "-" indicates no data. Due to technical reasons, data of snow, sea ice, and brine were  
873 not obtained.

874

875

876 Table 3.  $\text{CO}_2$  flux measured over the snow ( $F_{\text{snow}}$ ), frost flowers ( $F_{\text{ff}}$ ), and ice surface  
877 ( $F_{\text{ice}}$ ). Values measured directly over undisturbed surfaces (either with frost flowers or  
878 on snow surface) at a given station are indicated in bold.

879

880 a. Data of first  $\text{CO}_2$  flux measurement after removal of snow or frost flowers.

881

882 b. "-" indicates no data.

883

884 c. Number of measurements in bracket.

885

886 d. Data from station OI1 was not included.

Table 1. Station, position, date for CO<sub>2</sub> flux measurement, floe number, surface condition, ice type and thickness of snow, frost flower, and sea ice.

Station	Position	Date of 2015	Floe number	Surface condition	Ice type <sup>c</sup>	Thickness (cm)		
						Snow	Frost flower	Sea ice
FI1	83°03.77N, 17°34.94E	28 January	1	Frost flower	First-year ice	0.0	1.0	37.0
FI2	83°03.77N, 17°34.94E	28 January	1	Snow	First-year ice	8.0	No	35.0
FI3	83°08.00N, 24°09.02E	5 and 8 March <sup>a</sup>	2	Snow	First-year ice	29.0	No	98.0
FI4	83°10.56N, 22°09.42E	9 March	2	Snow	First-year ice	36.0	No	92.0
FI5	83°06.02N, 21°38.29E	10 and 11 March <sup>b</sup>	2	Snow	First-year ice	3.0	No	48.0
FI6	82°55.36N, 21°25.92E	12 March	2	Snow	First-year ice	37.0	No	69.0
FI7	81°22.18N, 08°59.93E	13 May	3	Snow	First-year ice	26.5	No	127.0
YI1	82°52.52N, 21°16.54E	13 March	2	Frost flower	Young ice	0.0	1.0	15.0
YI2	81°46.53N, 13°16.00E	5 May	3	Snow and frost flower mixed	Young ice	2.5	2.5	17.5
YI3	81°32.45N, 11°17.20E	9 May	3	Snow and frost flower mixed	Young ice	2.0	2.0	22.0
OI1	83°07.18N, 24°25.59E	6 March	2	Snow	Old ice (multi-year ice)	60.0	No	>200

a. Sea ice coring, brine and snow sampling was conducted on 5 March 2015.

b. Sea ice coring, brine and snow sampling was conducted on 10 March 2015.

c. Ice type was categorized based on WMO (1970).

Table 2. Station, snow density and water equivalent, brine volume fraction and temperature for sea ice (top 20 cm), brine temperature, salinity, DIC, TA, pCO<sub>2</sub> (pCO<sub>2,b</sub>) and atmospheric temperature, wind speed, pCO<sub>2</sub> (pCO<sub>2,a</sub>)<sup>a</sup> and ΔpCO<sub>2,b-a</sub>.

Station	Snow		Sea ice (top 20 cm)		Brine					Atmosphere				
	Density <sup>b</sup> (kg m <sup>-3</sup> )	Water equivalent (kg m <sup>-2</sup> )	Brine volume fraction (%)	Temperature (°C) (range)	Temperature (°C)	Salinity	DIC (μmol kg <sup>-1</sup> )	TA (μmol kg <sup>-1</sup> )	pCO <sub>2,b</sub> (μatm)	Temperature (°C)	Wind speed (m second <sup>-1</sup> )	pCO <sub>2,a</sub> (μatm)	ΔpCO <sub>2,b-a</sub> (μatm)	
F11	— <sup>c</sup>	— <sup>c</sup>	— <sup>c</sup>	— <sup>c</sup>	— <sup>c</sup>	— <sup>c</sup>	— <sup>c</sup>	— <sup>c</sup>	— <sup>c</sup>	— <sup>c</sup>	-31.6	4.0	405	— <sup>c</sup>
F12	— <sup>c</sup>	— <sup>c</sup>	— <sup>c</sup>	— <sup>c</sup>	— <sup>c</sup>	— <sup>c</sup>	— <sup>c</sup>	— <sup>c</sup>	— <sup>c</sup>	— <sup>c</sup>	-31.6	4.0	405	— <sup>c</sup>
F13	399	104	9	-6.8 (-7.4 to -6.3)	-5.2	84.8	4628	5539	427	-3.3	9.0	400	27	
F14	400	180	9	-4.7 (-5.5 to -3.7)	-5.3	86.6	4433	5490	334	-3.5	6.2	386	-52	
F15	268	11	17	-3.5 (-3.8 to -3.1)	-3.3	51.8	3261	3518	472	-18.1	6.8	389	83	
F16	343	127	13	-4.8 (-5.7 to -3.8)	-4.8	84.0	4841	5493	693	-25.0	3.6	400	293	
F17	— <sup>c</sup>	— <sup>c</sup>	— <sup>c</sup>	-6.1 (-6.1 to -5.8)	— <sup>c</sup>	— <sup>c</sup>	— <sup>c</sup>	— <sup>c</sup>	— <sup>c</sup>	— <sup>c</sup>	-13.0	5.8	405	— <sup>c</sup>
Y11	— <sup>c</sup>	— <sup>c</sup>	17	-6.6 (-12.3 to -2.6)	— <sup>c</sup>	— <sup>c</sup>	— <sup>c</sup>	— <sup>c</sup>	— <sup>c</sup>	— <sup>c</sup>	-26.0	2.6	402	— <sup>c</sup>
Y12	— <sup>c</sup>	— <sup>c</sup>	— <sup>c</sup>	-3.6 (-5.1 to -1.8)	— <sup>c</sup>	— <sup>c</sup>	— <sup>c</sup>	— <sup>c</sup>	— <sup>c</sup>	— <sup>c</sup>	-16.2	4.5	407	— <sup>c</sup>
Y13	— <sup>c</sup>	— <sup>c</sup>	— <sup>c</sup>	-3.9 (-6.4 to -2.0)	— <sup>c</sup>	— <sup>c</sup>	— <sup>c</sup>	— <sup>c</sup>	— <sup>c</sup>	— <sup>c</sup>	-14.2	6.7	410	— <sup>c</sup>
O11	— <sup>c</sup>	— <sup>c</sup>	0	-10.8 (-11.0 to -10.9)	— <sup>c</sup>	— <sup>c</sup>	— <sup>c</sup>	— <sup>c</sup>	— <sup>c</sup>	— <sup>c</sup>	-13.5	4.7	397	— <sup>c</sup>

a. pCO<sub>2,a</sub> (μatm) was calculated from CO<sub>2</sub> concentration (ppmv) at Ny-Ålesund, Svalbard (<http://www.esrl.noaa.gov/gmd/dv/iadv/>) taking into account the saturated water vapor and atmospheric pressures at sampling day.

b. Mean values for column.

c. "—" indicates no data. Due to the technical reason, data of snow, sea ice, and brine were not obtained.

Table 3. CO<sub>2</sub> flux measured over the snow (F<sub>snow</sub>), frost flowers (F<sub>ff</sub>) and ice surface (F<sub>ice</sub>).

Station	CO <sub>2</sub> flux (mmol C m <sup>-2</sup> day <sup>-1</sup> )		
	Natural flux (mean ± 1SD)		Potential flux
	F <sub>snow</sub>	F <sub>ff</sub>	F <sub>ice</sub> <sup>a</sup>
FI1	– <sup>b</sup>	<b>+0.1 ± 0.1</b> (n=7) <sup>c</sup>	– <sup>b</sup>
FI2	<b>+0.4 ± 0.3</b> (n=13) <sup>c</sup>	– <sup>b</sup>	– <sup>b</sup>
FI3	<b>+0.1 ± 0.1</b> (n=7) <sup>c</sup>	– <sup>b</sup>	–0.6
FI4	<b>+0.1 ± 0.1</b> (n=6) <sup>c</sup>	– <sup>b</sup>	–0.8
FI5	<b>+0.6 ± 0.3</b> (n=5) <sup>c</sup>	– <sup>b</sup>	+2.6
FI6	<b>+0.2 ± 0.1</b> (n=5) <sup>c</sup>	– <sup>b</sup>	+11.8
FI7	<b>+0.1 ± 0.1</b> (n=10) <sup>c</sup>	– <sup>b</sup>	±0.0
YI1	– <sup>b</sup>	<b>+1.6 ± 0.2</b> (n=6) <sup>c</sup>	+7.3
YI2	– <sup>b</sup>	<b>+1.3 ± 0.2</b> (n=9) <sup>c</sup>	+1.0
YI3	– <sup>b</sup>	<b>+1.0 ± 0.4</b> (n=8) <sup>c</sup>	+1.1
OI1	<b>+0.1 ± 0.0</b> (n=6) <sup>c</sup>	– <sup>b</sup>	+0.2
Mean <sup>d</sup>	<b>+0.2 ± 0.2</b> (n=46) <sup>c</sup>	<b>+1.0 ± 0.6</b> (n=30) <sup>c</sup>	+2.5 ± 4.3 (n=9) <sup>c</sup>

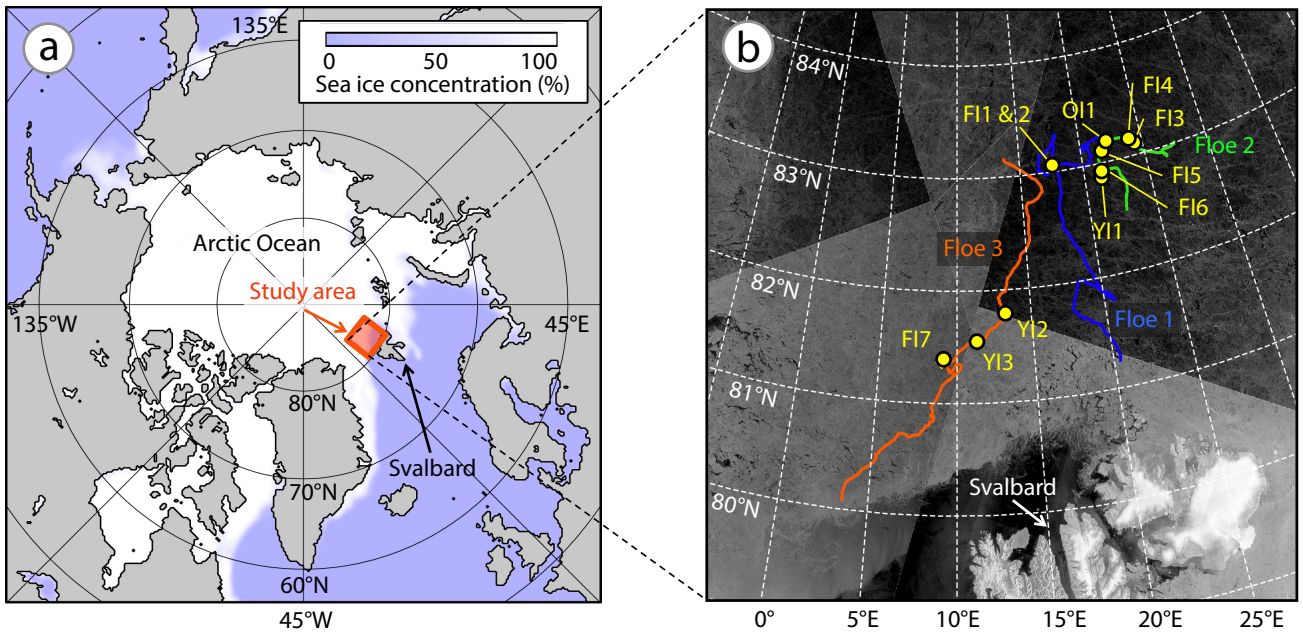
a. Data of first measurement after removal of snow or frost flower.

b. "–" indicates no data.

c. Number of measurements in bracket.

d. Data of station OI1 was not included.





Nomura et al., Figure 1 (Double column size)

