Springtime CO₂ exchange over seasonal sea ice in the Canadian Arctic Archipelago

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ABSTRACT. Springtime measurements of CO₂ exchange over seasonal sea ice in the Canadian Arctic Archipelago using eddy covariance show that CO₂ was generally released to the atmosphere during the cold (ice surface temperatures less than about –6 °C) early part of the season, but was absorbed from the atmosphere as warming advanced. Hourly maximum efflux and uptake rates approached 1.0 and ~3.0 µmol m⁻² s⁻¹, respectively. These CO₂ flux rates are far greater than previously reported over sea ice and are comparable in magnitude to exchanges observed within other systems (terrestrial and marine). Uptake generally occurred for wind speeds in excess of 6 m s⁻¹ and corresponded to local maxima in temperature at the snow–ice interface and net radiation. Efflux, on the other hand, occurred under weaker wind speeds and periods of local minima in temperature and net radiation. The wind speeds associated with uptake are above a critical threshold for drifting and blowing snow, suggesting that ventilation of the snowpack and turbulent exchange with the brine-wetted grains are an important part of the process. Both the uptake and release fluxes may be at least partially driven by the temperature sensitivity of the carbonate system speciation in the brine-wetted snow base and upper sea ice. The period of maximum springtime CO₂ uptake occurred as the sea-ice permeability increased, passing a critical threshold allowing vertical brine movement throughout the sea-ice sheet. At this point, atmospheric CO₂ would have been available to the under-ice sea-water carbonate system, with ramifications for carbon cycling in sea-ice-dominated polar waters.

INTRODUCTION

Despite recent reductions in sea-ice concentration in the north polar seas (Maslanik and others, 2007; Comiso, 2008), ice cover remains a defining feature of the Northern Hemisphere, covering between approximately 6 × 10⁶ and 15 × 10⁶ km² (US National Snow and Ice Data Center Sea Ice Index). Nonetheless, observations indicate that the cover is getting thinner (Rothrock and others, 2008; Kwok and others, 2009) and younger (Maslanik and others, 2007; Nghiem and others, 2007). Less is known about the role of sea ice on local, regional and global biogeochemical cycles relative to our understanding of the importance of sea ice to surface and atmospheric heat budgets (Serreze and Francis, 2006; Perovich and others, 2008). Most carbon-cycle research has not considered the possibility of either direct air–sea gas exchange in the presence of sea ice or indirect air–ice–sea gas exchange, where sea ice actively participates in CO₂ transfer. Therefore, global and regional budgets of air–sea CO₂ exchange have ignored ice-covered regions (Bates and Mathis, 2009, and references therein), relying instead on the assumption that a sea-ice cover is impermeable to gases. A growing body of recent research questions this assumption, as large fluxes of CO₂ have been reported over sea ice in winter (Miller and others, in press) and spring (Semiletov and others, 2004; Delille, 2006; Zemmelink and others, 2006), while other work has highlighted a complex sea-ice carbonate system coupled to that of the upper mixed layer (Delille and others, 2007; Rysgaard and others, 2007, 2009; Miller and others, in press).

It is not unreasonable to expect sea ice to exchange CO₂ with the atmosphere given the rich biological communities in the ice (Thomas and Dieckmann, 2010) and the temperature sensitivity of carbon speciation in both sea water (Zeebe and Wolf-Gladrow, 2001) and sea-ice brine (Papadimitriou and others, 2003). Ice algae can consume large quantities of carbon (Gosselin and others, 1997; Arrigo and others, 2010), and bacteria are so ubiquitous throughout the ice column that sea ice could even be net heterotrophic, despite photosynthetic CO₂ uptake at the bottom of the ice (Deming, 2010). Chemically, as sea water freezes, the increase in salinity of the resulting brine decreases gas solubility (Kilawee and others, 1998; Mock and others, 2002; Tison and others, 2002) and this salting-out effect generally dominates over the decrease in temperature, which increases gas solubility. This net decrease in solubility as salinity increases and temperature decreases in ice brines appears to be true also of CO₂, despite complications associated with its aquatic acid-base chemistry. Calcium carbonate precipitation from the brines (Papadimitriou and others, 2003; Dieckmann and others, 2008, 2010) would also serve to increase pCO₂. Therefore, pCO₂ in sea-ice brines can be extremely high (Miller and others, in press) and, depending on the location within the sea-ice sheet, CO₂ can be exported either to the underlying sea water (Rysgaard and others, 2007; Miller and others, in press) or the overlying atmosphere (Nomura and others, 2006; Miller and others, in press).

Observations show that sea ice is depleted in both brine and CO₂ in late spring (Delille and others, 2007; Rysgaard and others, 2009), thereby encouraging CO₂ uptake. On warming, Tison and others (2008) reported evidence of convective overturning within Antarctic sea ice, whereby dense high-salinity brine was replaced by sea water. This process requires that sea ice be permeable to liquid.
transport, which typically occurs at sea-ice brine volumes exceeding 5% (Golden and others, 1998) and provides a mechanism for replenishing sea-ice nutrients, including carbon. In addition, dissolution of any trapped CaCO3 in the sea ice on warming would consume CO2, decreasing pCO2 and thereby encouraging an air-to-sea-ice CO2 flux (Delille and others, 2007; Rysgaard and others, 2007).

The above-cited flux studies document exchanges that are episodic and potentially quite large and which seem to occur widely over sea ice in both hemispheres. Hence, it is important for CO2 fluxes associated with sea ice to be considered in polar marine carbon-cycle studies (observations and modeling). Parameterizations are unlikely until the community is able to pinpoint the (seasonally varying) processes that control the exchanges. As yet, few measurements of CO2 fluxes over sea ice exist, no doubt because the world’s sea-ice cover constitutes a truly extreme environment, making field studies both expensive and logistically challenging. Further, equipment and sensors often are required to operate at the edge of their tolerances, making accurate measurements difficult. More case studies are required to consolidate our understanding of the relationships that exist between sea-ice CO2 fluxes and the physical, chemical and biological components of the system.

In this paper we report the results of a study over seasonal sea ice during the spring of 2002 in the Canadian Arctic Archipelago (Fig. 1), where we measured heat, CO2 and momentum fluxes in conjunction with measurements of the site’s microclimate, including radiation balance and temperature structure. We have been able to identify relationships between the near-surface CO2 flux and site energetics, but we are unable to relate the observed fluxes to changes in the ice cover’s organic and inorganic carbon stores.

METHODS

The research site was situated over first-year sea ice in McDougall Sound approximately 3 km from the southwestern coast of Truro Island (75.246°N, 97.283°W) near the center of the Arctic Archipelago (Fig. 1). The flux station was situated on a pan of smooth landfast sea ice of uniform consolidation. Snow coverage on the pan varied from 5 to 50 cm, depending on the location of drifts. The ice thickness around the site ranged from 1.45 to 1.50 m at the start of the experiment and increased to 1.57–1.60 m over the course of the experiment.

The air–surface CO2 flux was measured between 9 May and 24 June 2002 using the eddy covariance technique. The flux system, a CSAT3 three-dimensional (3-D) ultrasonic anemometer (Campbell Scientific Inc., Logan, UT, USA) and LI-7500 open-path gas analyzer (LI-COR Inc., Lincoln, NE, USA), was mounted facing north at a height of 2.45 m above the snow surface. The gas analyzer was inclined at an angle of approximately 30°. Wind fetch (i.e., upstream distance of relatively uniform surface) from the measurement site exceeded 8 km across the angle arc between 300° and 60° from the flux system, as determined by a snowmobile survey and by positioning the site relative to land and adjacent flocs on a synthetic aperture radar (SAR) image. Information on bulk meteorology, radiation fluxes, and snow and ice temperature was available from sensors mounted on and around this installation. Sensors were powered by a solar panel/battery array and periodically by a gas generator situated 250 m directly to the south of the instrument tower.

Sensors were inspected daily to ensure that they remained in proper alignment and free of accumulated frost, snow or water. Turbulent fluctuations were scanned at 10 Hz using a data logger (model 21X, Campbell Scientific Inc., Logan, UT), while meteorological sensors, radiometers and thermocouple-based thermometers were scanned at 5 s intervals.

Trends in the turbulent fluctuations were removed through differencing, and the means, variances and covariances of the 3-D wind velocity, sonic temperature and scalar concentrations of CO2 and H2O were calculated through data-logger processing over 15 min periods. Post-processing of the data included a coordinate rotation of the covariances (McMillen, 1988) and density corrections for the CO2 flux, including the Webb–Pearman–Leuning (WPL) correction (Webb and others, 1980) following Massman and Lee (2002), with an additional correction based on Burba and others (2008) to compensate for air density fluctuations in the gas analyzer’s optical path associated with heating of the sensor. Sensor skin temperature required for the correction was estimated from air temperature using the multiple regression expression of Burba and others (2008) to compensate for air density fluctuations in the gas analyzer’s optical path associated with heating of the sensor. Sensor skin temperature required for the correction was estimated from air temperature using the multiple regression expression of Burba and others (2008) to compensate for air density fluctuations in the gas analyzer’s optical path associated with heating of the sensor. Sensor skin temperature required for the correction was estimated from air temperature using the multiple regression expression of Burba and others (2008) to compensate for air density fluctuations in the gas analyzer’s optical path associated with heating of the sensor.

Data were screened on the basis of wind direction and periods unfavorable for flux measurements, including power interruptions, frost and rime occurrence, and excessively high and low winds. Favorable wind directions were northward of the angular arc between 300° and 60° to minimize possible impacts of both the tower and generator on measured wind and CO2 concentration fluctuations. In addition, we noted that excessive tower vibration and blowing snow at wind speeds greater than approximately 12 m s−1 resulted in excessively noisy flux measurements. Diagnostic information from the ultrasonic anemometer and gas analyzer provided an objective means of screening the data for periods of frost. Data were rejected if the diagnostic parameter for window purity (AGC) reported by the LI-7500 deviated from its clean window value. At low wind speed, turbulent mixing may be insufficient to extend the surface mixed layer to the height of the flux sensors. These data were removed using a cut-off based on a wind velocity of 2 m s−1, which was determined graphically. Small gaps in the dataset (<3 h) were filled by linear interpolation. After screening, our
35 day flux record was distilled down to 18 days, leaving only those diurnal periods missing <3 hours of data.

Uncertainties in the eddy covariance estimates of the CO₂ flux have been discussed by Moncrieff and others (1996) and several others (including Goulden and others, 1996; Anthoni and others, 1999), and accuracy estimates range from ±12% to 25% of daytime values. The open-path eddy covariance system is well suited for remote deployment and has the advantage that the gas analyzer’s optical path can be situated very close to the sonic anemometer. Although many studies have shown excellent agreement between open-path-derived CO₂ fluxes and those derived from other eddy covariance systems (Leuning and King, 1992; Hirata and others, 2007; Ibrom and others, 2007; Burba and others, 2008; Haslwanter and others, 2009; Järvi and others, 2009), other studies have shown that open-path systems can underestimate the CO₂ flux (i.e. underestimate efflux and overestimate uptake), thought to be associated with: (1) sensor heating during winter season applications (Burba and others, 2008; Lafleur and Humphries, 2008; Ono and others, 2008; Järvi and others, 2009); (2) lens contamination from dust and water during unattended deployments (Serrano-Orti and others, 2008); or (3) a cross-sensitivity between the open-path’s concentration measurements of CO₂ and H₂O in marine applications (Pytherch and others, 2010). Burba and others (2008) found that wintertime open-path CO₂ flux measurements were within ±10% of closed-path measurements over several ecosystems when the effect of instrument heating on the ambient flux is corrected using their empirical relationship, which has the temperature range over which the correction of Burba and others (2008) was developed.

Sea-ice salinity and sea-water chemistry were available from water and ice samples extracted from a site approximately 4 km southwest of the flux tower. Ice cores for full salinity profiles were collected between 23 May (day of year (DOY) 143) and 3 June (DOY 156) (Johnston, 2006). The ice brine volume and salinity were estimated using the expressions of Cox and Weeks (1983) and measured sea-ice bulk density, salinity and temperature profiles (details appear in Mundy, 2003). Chlorophyll a (Chl a) accumulation within the bottom 5 cm of the sea ice was also measured three to four times a week (Mundy and others, 2005).

Surface sea water was sampled through a hole in the sea ice from 1 May to 9 June and water samples were analyzed for total inorganic carbon (TIC), total alkalinity (A_T), salinity and temperature. Sea water was extracted at 0.5 m depth below the ice base using GoFlo bottles prior to 7 May and using a subpump (e.g. Pondmaster Supreme Mag Drive) thereafter (Burba and others, in press). The samples were collected into triplicate 250 mL glass bottles and poisoned with 100 μL of saturated HgCl₂ solution. The ground glass stoppers were sealed with gasket and elastic closures and the samples were stored cold, but not frozen, until analysis at the Institute of Ocean Sciences (Sidney, British Columbia, Canada) within 4 months (Dickson and Goyet, 1994). Temperature was determined using a hand-held temperature probe placed in the outflow from the pump.

The TIC analyses were conducted by colometric titration with a SOMMA extraction system (Johnson and others, 1993), while alkalinity was determined by potentiometric titration using an open cell with end-point determination using the modified Gran plot (Hansson and Jagnér, 1973; Haraldsson and others, 1997). Both sets of analyses were calibrated using certified reference materials provided by A. Dickson of Scripps Institution of Oceanography, USA, and precision, based on the standard deviations between triplicate samples, was 2 μmol kg⁻¹ for both TIC and A_T. Salinity was measured on the remains of the TIC/A_T samples at the end of November 2002 using a Guildline Portasal model 8410 salinometer and calibrated with IAPSO (International Association of Meteorology and Atmospheric Sciences) standard sea water. We estimate that the error in salinity associated with the added mercuric chloride is <0.03 (psu).

RESULTS AND DISCUSSION

Environmental physical and chemical context

The 26 day experiment encompassed the period of transition from an early spring cold snowpack through to a period of snowmelt and melt-pond formation. Air temperature was highly variable throughout the study (Fig. 2), and the first positive air temperature was recorded on 2 June (DOY 153). The temperature at the ice–snow interface ranged from −9.8 °C at the onset of the experiment to a maximum value of −3.2 °C near the end of the experiment. The ice surface warmed rapidly between 10 and 19 May (DOY 130–139) and after 26 May (DOY 148). Storms centered on 11 May (DOY 131), 21 May (DOY 141) and 25–30 May (DOY 145–150) were associated with high surface winds. Gaps in the time series correspond to periods of particularly harsh weather, including blizzard and/or ice-riming conditions.

The ice surface and snow-base environments were both saline (Fig. 3a). Upwardly decreasing salinity gradients in snow over seasonal sea ice are common (e.g. Barber and others, 1995; Langlois and others, 2007). Salinity at the ice surface and 2 cm into the snowpack varied between 5.9 and...
15.3 over our experiment and decreased sharply with distance away from the snow base. At 4 cm from the snow base, the salinity was on average 3.5 less than at the snow–ice interface, while at 6 cm from the snow base the salinity ranged from 0 to 3 (not shown). Snow salinity decreased significantly after 25 May (DOY 145) at 4 cm up from the snow base and 30 May (DOY 150) at the ice surface, presumably the result of snowmelt. Ice salinity profiles showed a weak 'C-shape' distribution (not shown), averaging 7.5 ± 0.8 in the upper 15 cm of the cores and 5.1 ± 0.6 between 30 and 120 cm. The brine salinity at the ice surface (calculated from measured temperatures) decreased over the experiment from 140 to 70 (Fig. 3b) as a result of dilution by melting. Surface sea-water salinity below the ice fluctuated between 32.65 and 32.74, with no obvious trend (not shown). Sea-water pCO₂ (as calculated from TIC and Aₜ) declined over the period 18 May–4 June (DOY 138–155), yet remained supersaturated relative to atmospheric levels until late in the experiment (Fig. 3c).

Springtime air–surface CO₂ exchange

Figure 4 shows the hourly and daily average air–surface CO₂ exchange. The corrected flux tended to show a low-level CO₂ efflux (averaging 0.36 μmol m⁻² s⁻¹) prior to 20 May (DOY 140), followed by periods of moderate to strong CO₂ uptake. Maximum hourly efflux exceeded 1.0 μmol m⁻² s⁻¹, while hourly uptake approached –3.0 μmol m⁻² s⁻¹ on 30 May (DOY 150).

The maximum uptake rates in Figure 4 are in excess of any other values reported over sea ice to date (Semiletov and others, 2004; Zemmelink and others, 2006; Miller and others, in press) and are two to four times larger than hourly fluxes observed over the North Atlantic (McGillis and others, 2001). The fluxes observed by Semiletov and others (2004) approached –0.8 μmol m⁻² s⁻¹ over first-year sea ice near Barrow, Alaska, while Zemmelink and others (2006) reported fluxes up to –0.3 μmol m⁻² s⁻¹ over ablating Antarctic multi-year ice. In both of these studies, the CO₂ fluxes were measured using eddy covariance systems with open-path CO₂ sensors, consistent with this study, but without the correction of Burba and others (2008). Wintertime CO₂ fluxes reported by Miller and others (in press) over fast ice in the western Canadian Arctic showed considerable variability and were also during isolated periods quite large. The fluxes were generally directed upward (median flux of 0.14 μmol m⁻² s⁻¹). In that work, we speculated that the exchange was driven by some combination of sea-ice brine carbonate chemistry and biological activity. By comparison, wintertime hourly efflux in cold terrestrial systems can range anywhere between 0.25 μmol m⁻² s⁻¹ for wetlands (e.g. Lafleur and others, 2003) and approximately 0.45 μmol m⁻² s⁻¹ for upland boreal forest soils in Alaska (Sullivan and others, 2008).

Fick’s law for gas diffusion dictates that we would require a CO₂ concentration at the snow base of approximately 38 mmol m⁻³, roughly twice atmospheric levels, to sustain our average early experiment diffusive CO₂ efflux (0.36 μmol m⁻² s⁻¹) through the snow, assuming a snow depth of 0.3 m and diffusivity of 0.05 cm² s⁻¹. Diffusivity for gas transport in wind-packed snow can range between 0.02 and 0.08 cm² s⁻¹ (Albert and Shultze, 2002). An upward flux of 0.36 μmol m⁻² s⁻¹ is not inconsistent with...
field observations elsewhere, which indicate that sea-ice pCO$_2$ in the spring can exceed atmospheric values (Delille and others, 2007) and perhaps significantly so (Miller and others, in press). In laboratory experiments, Nomura and others, (2006) observed low-level CO$_2$ efflux from growing sea ice at a rate (0.004–0.012 μmol CO$_2$ m$^{-2}$ s$^{-1}$) that increased with increasing brine salinity (decreasing brine temperature). Ice temperature in their experiments ranged between −2.7°C and −4.1°C, which is warmer than our near-surface sea ice for all but the final days of our experiment (Fig. 2). Consequently, brine salinity in their study was lower by almost a factor of 2, which could partly explain why their observed average efflux was higher.

As the under-ice sea water was supersaturated in pCO$_2$ relative to atmospheric values for most of our study (Fig. 3c), the prevailing ocean–air pCO$_2$ gradient could also support an upward flux of CO$_2$. Through much of the study, the ice in McDougall Sound was likely impermeable to brine because the brine volume constituted <5% (the cut-off for brine permeability in columnar sea ice; Golden and others, 1998) of the sea-ice interior prior to 26 May (DOY 146) (Fig. 5) and hence CO$_2$ probably could not have been transported by the brine. However, little is known about the comparable temperature cut-off for gas diffusion in sea ice, aside from the laboratory study of Gosink and others (1976) who observed that atmospheric CO$_2$ could have been moving through the ice throughout our study, something that we cannot confirm.

Spring heating raised the ice temperature above −5°C by 28 May (DOY 148), with a corresponding increase in sea-ice brine volume above 5% (Fig. 5). Beyond this date, the brine should have been draining freely and mixing with the underlying sea water (Golden and others, 1998; Tison and others, 2008). The timing of this transition closely corresponded to the period of maximum CO$_2$ uptake observed between 28 and 30 May (DOY 148–150; Fig. 4), meaning that atmospheric CO$_2$ could have offset partial of any pCO$_2$ deficit developing at the ice–water interface as a result of algal production or melt dilution. In regard to the former, sea ice in the Canadian Arctic Archipelago ranks amongst the most productive in the Arctic, with reported primary production rates between 20 and 463 mg C m$^{-2}$ d$^{-1}$ (Arrigo and others, 2010). Chl a concentrations within the bottom 5 cm of the ice cover close to our research site (Mundy and others, 2005) were between 20 and 45 mg m$^{-2}$ by 29 May (DOY 149). Data shown by Lavoie and others (2005) indicate that pronounced ice algal accumulation in 2002 (commensurate with our study) had started at about 10 May (DOY 130) and persisted into the first week of June (approximately DOY 160) in sea ice approximately 70 km southeast of our study site and of comparable thickness and snow depth. Maximum Chl a concentration ranged between 55 and 80 mg m$^{-2}$ at their site, with net particulate organic carbon (POC) accumulation rates ranging between 28 and 55 mg C m$^{-2}$ d$^{-1}$, depending on snow thickness. Algae are the main contributors of POC at the base of Arctic first-year sea ice (Riedel and others, 2008). Accumulation rates were obtained by applying a POC:Chl a ratio of 20 (Riedel and others, 2008) to Chl a time-rate-of-change estimates obtained graphically from figures provided by Lavoie and others (2005), over the period between 10 May (DOY 130) and 25 May or 10 June (DOY 145 or 161) for thick and thin snow, respectively. Uptake, according to our tower measurements, averaged approximately 23 mg C m$^{-2}$ d$^{-1}$ between 10 and 30 May (DOY 130–150). This value is close in magnitude to the POC accumulation rates that we derived from Lavoie and others (2005) and on the low end of primary production rates reported for sea ice in the Canadian Arctic Archipelago.

Although we expect the ocean mixed layer to be the main supplier of nutrients to the skeletal layer at the ice base (Thomas and others, 2010, and references therein), it is not inconceivable for atmospheric CO$_2$, via brine drainage, to offset short-term CO$_2$ deficits at the ice base given that: (1) both TIC and CO$_2$(aq) in sea-ice brine can be depleted under conditions of high primary production and large accumulation of algal biomass (Thomas and others, 2010, and references therein); (2) the nutrient supply from the mixed layer to biomass at the ice base can be restricted by the thickness of the molecular sublayer (Lavoie and others, 2005); and (3) downward flushing of nutrients into sea ice associated with snowmelt has been documented and thought able to augment sea-ice biomass accumulation (e.g. Granskog and others, 2003).

Finally, if CaCO$_3$ had precipitated in the ice when it formed, the late-spring melting process would have dissolved the salts, lowering pCO$_2$ within the ice and further encouraging CO$_2$ drawdown from the atmosphere. Collectively, these observations support the argument that the observed late-season CO$_2$ uptake was a result of a permeable ice cover and a lowering of pCO$_2$ in the ice and sea water through photosynthesis, melt and possibly calcium carbonate dissolution.

The magnitude of fluxes reported here raises concerns about possible measurement biases associated with the open-path system. It is evident (Fig. 4) that prior to 20 May (DOY 140) the heating correction constituted a large component of the corrected flux, often changing the flux direction from predominately a regime of low-level uptake to low-level efflux. Accordingly, we are less confident of measurements during this period given the possibility that

Fig. 5. The evolution of the 1200 h (local time: North American Central Standard Time) temperature and brine volumetric fraction in the upper meter of sea ice.

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the heating correction to the CO2 flux represents only a small component of the strong midday uptake that occurred on 21, 28 and 30 May and 3 June (DOY 141, 148, 150 and 154). We are also reasonably confident that the measured uptake at these times is not the result of an obstruction of the open-path lens (Serrano-Orti and others, 2008) or cross-sensitivity to the water vapour flux (Pyrtherch and others, 2010). In regard to the former, we do observe a decrease in atmospheric CO2 concentration of 4–5% corresponding to CO2 uptake on 28 and 30 May (DOY 148 and 150) (Fig. 6a), which if related to lens obstruction would artificially lower our flux measurement by as much as 30% through a reduction in the WPL correction (Serrano-Orti and others 2008). The sensor lens was checked for cleanliness daily, and diurnal fluctuations in ambient CO2 of >10% are not uncommon over ecosystems with strong 24 hour cycles of net photosynthesis (Anthoni and others, 1999; Yang and others, 1999), hence we expect our measurements over these periods to be within the sensor’s specifications. High-frequency measurements of CO2 and H2O concentration are not available from this study to explore possible sensor cross-sensitivity, but data from subsequent experiments over winter- or springtime sea ice show no coherent relationship between high-frequency measurements of CO2 mixing ratio and relative humidity. Further, our measured latent heat flux (±7 W m–2) between 26 and 31 May (DOY 146–151; Fig. 6b) is close to a factor of 10 smaller than reported in Pyrtherch and others (2010) over open water, indicating that if there was cross-sensitivity in the gas concentration measurements, it was not strongly correlated to vertical velocity or, hence, the CO2 flux.

Diurnal variation in air–surface CO2 exchange

Prominent shifts from efflux to uptake were observed several times, including on 10 May (DOY 130), 21–22 May (DOY 141–142) and then 26, 28 and 30 May and 2 June (DOY 146, 148, 150 and 155) (Fig. 4). Two periods with large CO2 flux (FCO2) ranges are shown in Figure 7a, showing the measured flux between 20 and 24 May (DOY 140–142) and between 26 and 30 May (DOY 146–150). Also shown are the corresponding variations in surface net radiation (Rn, Fig. 7b), ice surface temperature (T, Fig. 7c), horizontal wind speed (u, Fig. 7d) and air temperature (Ta, Fig. 7e). The interface temperature reflects the thermodynamic state of the surface. Net radiation provides a measure of heating and cooling of the surface via radiation, while horizontal wind speed is related to surface shear and forced convection.

Uptake on 21 May (DOY 141) started during the night of 20 May (DOY 140) and increased over the course of the morning and afternoon of 21 May (Fig. 7a). The uptake occurred during the first high wind event following the intense heating of the snowpack on 19 May (DOY 139) (Fig. 2). The start of the CO2 uptake during the evening of 20 May (DOY 140) coincided with the rise in wind speed (Fig. 7d), and uptake stopped when the wind speed dropped again below 6 m s–1 in the early morning of 22 May (DOY 142). The Gaussian-shaped timeline of hourly Rn centered midday on 21 May (DOY 141; Fig. 7b) indicates clear-sky conditions, which allowed the surface to cool intensely over much of the night. Peak uptake on 21 May (DOY 141) also corresponded in time to the 24 hour maximum in Rn and a rise in T. The efflux observed at about midnight at the start of 22 May (DOY 142) corresponded to the 24 hour minimum in Rn.

Strong CO2 uptake was also observed on 26, 28 and 30 May (DOY 146, 148 and 150), with efflux occurring just about midnight on 27 and 28 May (DOY 147 and 148). The periods of efflux corresponded to night-time minima in Rn, air temperature and cooling at the snow base, again associated with clear-sky conditions. Over this period, the hourly pattern in FCO2 mirrored wind speed. High winds (generally >6 m s–1) gave rise to large CO2 uptake rates, and low winds corresponded to periods of little exchange or efflux. The largest uptake rate observed over the entire experiment occurred on 30 May (DOY 150) under conditions of extremely high wind speed (>9 m s–1) and high interface (greater than −5.5 °C) and air (approximately −5 °C) temperatures.

We conclude that, in general, CO2 uptake and efflux mirrored wind speed, with winds in excess of approximately 5–6 m s–1 supporting uptake, while lighter winds were associated with efflux. The relationship between wind speed and FCO2 is nonlinear in nature, as evident in Figure 8 for the period 26–30 May (DOY 146–150). High winds ventilate the snow (Colbeck, 1989; Albert and Shultz, 2002) and above a variable threshold (4.4–11.0 m s–1) give rise to blowing snow (Li and Pomeroy, 1997). Snow ventilation allows exchange between the atmosphere and saline snow

Fig. 6. The time series of average (a) atmospheric pCO2 and (b) latent heat flux (Rε) between 26 and 31 May (DOY 146–151). The flux of CO2 (FCO2) is provided in both plots for reference.
base/ice surface. In addition, blowing snow has a high surface area:mass ratio and is subject to intense atmospheric turbulence, hence the potential for turbulent transfer of trace gases (in addition to heat) to and from the snow grains (Pomeroy and others, 1999) or, in our case, brine-wetted snow grains. The efficiency of gas adsorption/desorption on ice grains will be proportional to the changing snow surface area (Legagneux and others, 2002). Hence, blowing snow during conditions conducive to down-directed sensible heating (i.e. a warming atmosphere) allows for rapid heating of the brine-wetted snow base and ice surface by convection in the snow. On the other hand, in the absence of high winds, CO₂ exchange with the snow base and ice surface is controlled by diffusion, and the upper snow may moderate the exchange by storing air (and CO₂).

Uptake also corresponded to local peaks in both net radiation and temperature, while efflux generally occurred in association with local minima in net radiation and temperature.

![Graphs showing hourly variation in CO₂ flux, surface net radiation, snow–ice interface temperature, wind speed, and air temperature for different periods.](image)

*Fig. 7. Hourly variation in (a) CO₂ flux (FCO₂), (b) surface net radiation (R₀), (c) snow–ice interface temperature (Tᵢ), (d) wind speed (u) and (e) air temperature (Tᵃ) for the periods 20–23 May (DOY 140–143) and 26–30 May (DOY 146–150). No data are available from 29 May (DOY 149).*
We measured springtime CO2 fluxes over seasonal sea ice observed daytime CO2 uptake, given the dependency of intuitive that net photosynthesis could contribute to our explore relationships to biology with this dataset, it is indicative of a biologically controlled system of diurnal daytime uptake and nocturnal efflux observed here is combination of: increasing gas solubility in the brines in the upper sea ice and snow base; strong air-to-brine pCO2 gradients associated with nocturnal degassing and CO2 depletion in the brine (Delille and others, 2007; Rysgaard and others, 2007); and possibly dissolution of calcium carbonate (Papadimitriou and others, 2003). The pattern of daytime uptake and nocturnal efflux observed here is indicative of a biologically controlled system of diurnal photosynthesis and respiration. Although we are unable to explore relationships to biology with this dataset, it is intuitive that net photosynthesis could contribute to our observed daytime CO2 uptake, given the dependency of photosynthesis on photosynthetically active radiation.

SUMMARY AND CONCLUSIONS

We measured springtime CO2 fluxes over seasonal sea ice in the Canadian Arctic Archipelago that were much greater than reported observations from other studies over sea ice (Semiletov and others, 2004; Zemmelink and others, 2006; Miller and others, in press). Efflux tended to dominate the exchange early in the experiment, when sea-ice surface temperature was less than approximately –6°C, giving rise to episodes of increasingly large rates of uptake as the sea ice warmed.

Diurnal variation of the CO2 flux (both the direction and magnitude) corresponded to fluctuations in basic micro-meteorological variables. Uptake of CO2 was generally associated with warming in the presence of high winds, while cooling and low winds gave rise to CO2 effluxes. In general, uptake rarely occurred when the wind speed was <4.5 m s\(^{-1}\). The high wind speeds associated with uptake (>6 m s\(^{-1}\)) suggest that ventilation of, and turbulent exchange with, the snow are an important part of the process. The flux may also be influenced by calcium carbonate precipitation and dissolution at the brine-wetted snow base and upper sea-ice surface.

Early in the season (ice temperature less than –6°C), the sea ice appears to be a source of atmospheric CO2, consistent with observations from a winter study (Miller and others, in press), also over seasonal sea ice. Later in the season, the magnitude of uptake increases as the ice surface temperature increases past –5°C, corresponding to an increase in brine volume and permeability, permitting vertical drainage and coupling with the underlying sea water.

These findings raise the prospect of atmospheric CO2 directly contributing to the sea-water carbon drawdown in the presence of an ice cover, with consequences for both basic carbon budgets and biophysical modeling of ice ecosystem environments. Further CO2 flux studies over sea ice, particularly those involving the simultaneous deployment of open- and closed-path eddy covariance systems in conjunction with measurements of time-dependent sea-ice carbon inventory, are required to confirm the robustness of existing corrections to eddy covariance CO2 flux measurements and the spatial relevance of the processes described here.

ACKNOWLEDGEMENTS

Sea-ice camps are inherently expensive to operate and numerous investigators and agencies participated in the C-ICE 2002 experiment. This research was largely funded by a Natural Sciences and Engineering Research Council of Canada (NSERC) grant (Discovery) to T. Papakyriakou and the Department of Fisheries and Oceans Canada (DFO) Strategic Science Fund (L. Miller). Resources for infrastructure and logistics were shared among several agencies, including Natural Resources Canada, Environment Canada (Canadian Ice Service), NSERC and the University of Manitoba (Centre for Earth Observation Science). We thank D. Arychuk, M. Davelaar, M. Fortier, K. Johnston, C.J. Mundy, K. Morris and O. Owens for field and laboratory assistance, and the reviewers and the scientific editor for their valuable suggestions. We thank the Polar Continental Shelf Project (PCSP) for logistical support.

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