CO2 and O2 variability in the partially ice-covered Arctic Ocean

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CO$_2$ AND O$_2$ VARIABILITY IN THE PARTIALLY ICE COVERED ARCTIC OCEAN

By

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Limited carbon cycle research has been conducted so far in the Arctic Ocean (AO) compared to many other open-ocean and coastal environments, with relatively few studies of the inorganic carbon cycle and air-sea gas exchange. Understanding these processes in depth and understanding the physical, chemical, and biological processes that control carbon dioxide (CO\textsubscript{2}) and dissolved oxygen (DO) variability in the AO are crucial to predicting the future of the carbon cycle in the region and its impact on greenhouse gases and marine ecosystem processes, such as ocean acidification. To study the AO carbon cycle, \textit{in situ} time-series data have been collected from the Canada Basin of the AO during late summer to autumn of 2012. Partial pressure of CO\textsubscript{2} (\(p\text{CO}_2\)), DO concentration, temperature, salinity, and chlorophyll-a fluorescence (Chl-a) were measured at 6-10 m depth under little ice and multi-year ice on two drifting platforms. The \(p\text{CO}_2\) levels were always below atmospheric saturation, whereas the seawater was almost always slightly supersaturated with respect to DO. Although the two time-series data were on an average only 222 km apart they had 10 \(\pm\) 10\% and 63 \(\pm\) 16\% ice cover and differed significantly in contributions from gas exchange and net community production (NCP). Modeled variability of CO\textsubscript{2} and DO suggest that gas exchange, NCP and horizontal gradients are the main sources of the CO\textsubscript{2} and DO variability in the partially ice-covered AO. Horizontal gradients dominated the more densely ice-covered region, with no significant NCP in the surface. These results suggest that the signature imparted on CO\textsubscript{2} and DO in open water is widely disbursed under-ice and that biological production under multi-year ice is negligible due to lack of light and nutrients.
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Chapter 1

Introduction

1.1. Overview

The Arctic Ocean (AO) is changing rapidly. Decrease in sea ice thickness (Perovich et al., 2003, 2007; Maslanik et al., 2007), freshening of the sea surface (Yamamoto-Kawai et al., 2009a), changing mixed-layer dynamics (Toole et al., 2010) and increased primary production (Arrigo and van Dijken, 2012) provide evidence that steady state conditions no longer exist in the AO. Recent decreases in Arctic summer sea ice area have also increased exchange of CO$_2$ and DO between the atmosphere and the ocean. Ocean acidification might be accelerated as ice cover decreases, potentially resulting in the largest and most rapid pH decrease of all the ocean basins during this century (Steinacher et al., 2009). While most previous observations have found large $p$CO$_2$ undersaturation, there is evidence that loss of ice cover has already increased $p$CO$_2$ levels in AO surface waters (Cai et al., 2010; Else et al., 2013). It remains uncertain, however, whether the deep AO basins will uptake significant CO$_2$ under seasonal ice-free conditions. Rapid equilibration and warming of the shallow isolated surface water and weak biological CO$_2$ drawdown are hypothesized to limit CO$_2$ invasion, keeping the ice-free central AO basins from becoming large atmospheric CO$_2$ sinks (Cai et al., 2010). The net effects of all these climate change impacts are unknown, but they show clear evidence of significant biogeochemical changes in the AO.
1.2. AO and Canada Basin

The Arctic Ocean is nearly landlocked, almost completely surrounded by Eurasia and North America (Fig. 1.0). It makes up ~4.3% of the ocean area but only ~1.4% of the volume (Jakobsson, 2002). The AO consists of a deep ocean basin, the broad shelves of the Barents, Kara, Laptev, East Siberian, Chukchi, Beaufort, White and Lincoln Seas and the narrow shelf off the Canadian Arctic Archipelago and northern Greenland (Fig. 1.0). The AO is partly covered by sea ice throughout the year and almost completely covered in winter. The surface temperature and salinity of the AO vary as the ice cover melts and freezes (Aagaard and Woodgate, 2001). Its salinity is the lowest compared to other oceans, due to low evaporation, heavy fresh water inflow, ice melt, and limited connection to nearby oceans that have comparatively higher salinities.

The Canada Basin is the largest sub-basin of the Arctic Ocean, extending approximately 1100 km from the Beaufort Sea shelf to the Canadian Archipelago (Fig. 1.0), with an average depth of 3800 m. Studying the Canada Basin of the AO has become particularly important due to its uniqueness in several respects (McLaughlin et al., 2011). The Canada Basin is unique compared to other ocean basins in that it is surrounded by relatively broad and shallow (<200 m deep) continental shelves that comprise about 53% of the area of the AO (Bates and Mathis, 2009). The surface transpolar drift separates waters of the Canada Basin in the central basin of the AO from the Eurasian Basin. Its surface waters and subsurface halocline waters have distinctly different physical and chemical properties compared to the other basins. Warm and salty water from the North Atlantic enters the Canada Basin and forms the layer between about 250 and 800 m beneath the relatively cold Pacific water layer (Timmermans et al., 2008). The loss of both multiyear and first-year ice is greater in the Canada Basin com-
pared to the other three sub-basins in the AO (McLaughlin et al., 2011), making it more vulnerable to climate change.

**Figure 1.0** The International Bathymetric Chart of the Arctic Ocean (IBCAO, Jakobsson et al., 2000) showing the Arctic marine regions. The study area is highlighted by the yellow star.
1.3. CO₂ and DO in the AO

The lack of long-term time-series data from the AO has limited scientists’ understanding of the CO₂ sources and sinks and inorganic carbon cycling in the AO. Sea surface $p$CO₂ data sets from the central AO are scarce with most Arctic studies focused on nearshore and shelf areas (Anderson et al., 2009; Bates et al., 2006, Murata and Takizawa, 2003; Kaltin and Anderson, 2005). Anderson and Kaltin (2001) reported 250-300 μatm $p$CO₂ in the Eurasian Basin surface waters in August 1996. Over the time period of 1991 to 2005, Jutterström and Anderson (2010) found that all waters in the central AO are undersaturated with values typically below 300 μatm. Low $p$CO₂ values of 240-280 μatm have been observed in the Canada Basin adjacent to the Chukchi Sea shelf during summer 2002 (Bates, 2006; Bates et al., 2006), whereas 160-280 μatm $p$CO₂ have been observed along the sea-ice edge of the Chukchi and Beaufort Seas during September (Pipko et al., 2002; Murata and Takizawa, 2003). Fransson et al. (2009) reported lower surface seawater $p$CO₂ values of 150–250 μatm in the Makarov Basin of the Canada Basin from the summer of 2005. Cai et al. (2010) reported 250-365 μatm $p$CO₂ in the Canada Basin during summer 2008. Else et al. (2013) reported ~290-320 μatm $p$CO₂ beneath heavily decayed ice cover in the southeastern Canada Basin of the AO from early September of 2009. In the course of the late-summer 2011, Robbins et al. (2013) measured 322 μatm $p$CO₂ in the surface waters of the Canada Basin. All of these studies are summarized in Table 1.0.
Table 1.0. List of some previous studies of $p$CO$_2$ in or near the Canada Basin of the AO. Our measured $p$CO$_2$ values in open seawater and under multiyear ice coverage in the Canada Basin are typical of previous reports in the region. The list of the studies summarized below proceeds chronologically from top to bottom.

<table>
<thead>
<tr>
<th>Study</th>
<th>Time period</th>
<th>Region</th>
<th>$p$CO$_2$ (µatm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anderson and Kaltin</td>
<td>August 1996</td>
<td>Eurasian Basin</td>
<td>250-300</td>
</tr>
<tr>
<td>(2001)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Jutterström and Anders-</td>
<td>From 1991 to</td>
<td>Central Arctic Ocean</td>
<td>&lt; 300</td>
</tr>
<tr>
<td>son (2010)</td>
<td>2005</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bates (2006)</td>
<td>Summer 2002</td>
<td>Canada Basin (adjacent to the Chukchi Sea</td>
<td>240-280</td>
</tr>
<tr>
<td>Bates et al. (2006)</td>
<td></td>
<td>shelf)</td>
<td></td>
</tr>
<tr>
<td>Pipko et al. (2002)</td>
<td>Summer 2002</td>
<td>Chukchi and Beaufort Seas (along the sea-ice</td>
<td>160-280</td>
</tr>
<tr>
<td>Murata and Takizawa</td>
<td></td>
<td>edge)</td>
<td></td>
</tr>
<tr>
<td>(2003)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Else et al. (2013)</td>
<td>Late summer 2009</td>
<td>Southeastern Canada Basin (beneath heavily</td>
<td>290-320</td>
</tr>
<tr>
<td></td>
<td></td>
<td>decayed ice cover)</td>
<td></td>
</tr>
<tr>
<td>Robbins et al. (2013)</td>
<td>Late-summer 2011</td>
<td>Canada Basin</td>
<td>247-555</td>
</tr>
<tr>
<td><strong>Our study</strong></td>
<td>Late-summer to fall 2012</td>
<td>Canada Basin</td>
<td><strong>270-330</strong></td>
</tr>
</tbody>
</table>
Dissolved $O_2$ (DO) is an important climate parameter and is a key measurement for understanding the processes that control the marine carbon cycle and $CO_2$ variability. DO saturation in the ocean is predicted to decline by 4 to 7% by the end of this century because of climate change (Bopp et al., 2002; Matear et al., 2000; Plattner et al., 2001; Sarmiento et al., 1998). Recent reduction in DO in lower-latitude oceans (e.g., Johnson and Gruber 2007; Mecking et al., 2008; Stramma et al., 2008) may be linked to global climate change, which is also relevant to a warming and freshening AO (Proshutinsky et al., 2009). According to available data, the central AO surface mixed layer is typically 2-3% of atmospheric saturation with DO maxima in the halocline (Falkner et al., 2005). Low concentrations around 90% of saturation have been reported under winter sea ice cover in the southernmost Canada Basin (Sherr and Sherr, 2003). Sherr and Sherr (2003) observed a subsurface DO peak (>400 $\mu$mol kg$^{-1}$) during November 1997 in south-central Canada Basin. Hill and Cota (2005) observed supersaturation of DO in the Canada Basin during the spring and summer of 2002. Timmermans et al. (2010) observed a subsurface DO maximum consistently around 100% saturation between August and December 2008 in the Canada Basin.

Almost all of the $pCO_2$ and DO studies cited above were based on ships collecting data during the low ice periods in the summer. There continue to be significant gaps in our understanding of $CO_2$ and DO dynamics in the AO, especially under ice. To examine $pCO_2$ and DO variability in the AO, we deployed $pCO_2$, DO, temperature, salinity, Chl-$a$, and PAR sensors on ice-tethered profilers (ITPs) as part of the Joint Ocean Ice Study (JOIS) 2012 cruise. In 2012, the AO reached the lowest seasonal ice extent since the satellites began quantifying ice coverage in 1979 (Parkinson et al., 2013; Zhang et al., 2013). The most notable event in 2012 was a very strong storm that swept over the central AO in early August, rapidly enhanc-
ing ice melt and decreasing ice thickness (Parkinson et al., 2013). Our sensors were deployed in late August, a few weeks after the storm and a couple of weeks before the ice extent reached its minimum on September 16. The two instrument packages were deployed in densely and sparsely ice-covered locations, providing a unique opportunity to compare CO₂ and DO variability under these different conditions. My research objective is to better understand the CO₂ and O₂ variability in the region through exploring the following questions:

1. **What is the open-water and under-ice pCO₂ and O₂ variability?**
2. **What controls the variability of pCO₂ and O₂?**
3. **Is the AO a net source or sink for atmospheric CO₂?**
4. **What implications does the changing Arctic have for air-sea CO₂ fluxes, ocean acidification and biological production?**

These questions will be examined in detail by use of correlative relationships, modeling and time-series (bandpass filtering) analysis.
Chapter 2

Methods

2.1. Study Site

Our instrumentation was deployed on two ITPs (ITP-64 and ITP-65) (Toole et al., 2011; Krishfield et al., 2008) (Fig. 2.1 and 2.2) in the Canada Basin as part of the Beaufort Gyre Observing System (BGOS) study during the JOIS 2012 cruise on the Canadian Coast Guard Icebreaker (CCGS) Louis S. St. Laurent. ITP-65 was deployed on a 1.5 m thick ice floe on August 27, 2012 at 80° 53.4 N, 137° 25.8 W and ITP-64 was deployed in open water because of limited ice extent on August 28, 2012 at 78° 46.5 N, 136° 39.8 W (Fig. 2.1). The two drifters were initially 245 km away from each other (Fig. 2.1). The ITPs followed the ice flow, as a result, ITP-64 and ITP-65 traveled 513 and 575 km, respectively, during the period when the sensors transmitted data (Fig. 2.1).

Figure 2.1. Arctic marine regions (left) near the study area (inset) are shown with the ITP drift track overlaid. The study area is magnified on the right figure. Measurement start and end points are highlighted by rectangular boxes labeled with dates. $p$CO$_2$ values at different points over the drift track are shown in color. The top track corresponds to ITP-65 and the bottom track to ITP-64. The sensors were deployed roughly 245 km away from each other. Plotting software is courtesy of Bill Williams (Institute of Ocean Sciences, Canada).
2.2. *In situ* Sensors

2.2.1. The ITP

The ITPs (Krishfield *et al.*, 2008a) consist of 1) a surface buoy that houses an inductive modem, GPS receiver and Iridium satellite phone; 2) a weighted, jacketed, wire rope that extends to the end of the profiling range; and 3) an instrumented profiler (McLane Research Laboratories Inc.) containing a CTD (SBE41CP, Sea-Bird Electronics Inc.), DO sensor (SBE43I, Sea-Bird Electronics Inc.) and bio-optical package (Fig. 2.2).

![ITP schematic](image)

**Figure 2.2.** ITP schematic (left) and SAMI tethered ITP deployment through the ice (right). The SAMI sensor (black cylinder) is located at 6 m depth directly below the urethane casing ~2-4 m below the ice (not shown). The CTD and DO sensors are located above the SAMI (right and left silver housings, respectively). ITP schematic was obtained from WHOI website: http://www.whoi.edu/page.do?pid=23099.
All data collected by the ITPs were transmitted to the surface with inductive modems and telemetered via Iridium satellite. The transmitted data were downloaded, processed, and made available at the Woods Hole Oceanographic Institution (WHOI) website. The ITPs profiled between ~11-800 m depth with 4 profiles per day.

2.2.2. SAMI-CO₂

Two Submersible Autonomous Moored Instruments for CO₂ (SAMI-CO₂, Sunburst Sensors, LLC) (Fig. 2.3) were used to measure pCO₂ in seawater (SAMI-12 and SAMI-14). The SAMI uses pH indicators to quantify pCO₂. The quantification is done through equilibration with a pH indicator solution contained within a gas permeable membrane (DeGrandpre et al., 1995, 1999). The equilibrated solution is pumped into an optical cell where the absorbance of the indicator is measured at specific wavelengths (Fig. 2.3). With the pH indicator solution renewed for each measurement and the optical blanks measured intermittently, the SAMI-CO₂ has no detectable drift (DeGrandpre et al., 1995, 1999). The sensors were calibrated in our lab and tested with the ITP off the WHOI dock prior to deployment. SAMI-CO₂ are free from field calibration and are capable of operating at low temperatures of Arctic waters by adding ethylene glycol to the indicator and the blank solutions. Before deployment, each SAMI was calibrated using standard calibration gases over the pCO₂ range of interest at the expected average in situ temperature (DeGrandpre et al., 1995). Both SAMIs we deployed were connected to the ITP wire at ~6 m depth and recorded the pCO₂ at 2 hour intervals. SAMIs passed all data to the surface controller through an inductive modem interface (SBE UIMM 350 m). For unknown reasons, the SAMI inductive modem communications failed on both ITPs and the time-series for ITP-64 and ITP65 were limited to 49 and 41 days, respectively.
Figure 2.3. The SAMI-CO$_2$ sensor is shown on left. The right picture shows the sensor part of the SAMI (optical cell and membrane equilibrator). Pictures obtained from Sunburst Sensors website.

### 2.2.3. Ancillary Sensors

The SAMIs were interfaced with O$_2$ sensors (Optode 4175, Aanderaa Data Instruments) and a conductivity, temperature, and depth (CTD) sensor (SBE37SI, Sea Bird Electronics Inc.) (Fig. 2.4). The CO$_2$, O$_2$, and CTD sensors were distributed on the wire rope with depths differing by <0.5 m (Fig. 2.2). SAMIs passed all ancillary sensor data to the surface controller through an inductive modem interface (SBE UIMM 350 m). The O$_2$ sensors were calibrated in the lab using both an air-saturated solution and a zero oxygen solution at room temperature. All instruments were tested off the WHOI dock prior to deployment. Like the SAMIs, the profilers were configured with CTD sensors (SBE41CP, Sea-Bird Electronics Inc.) that had an integrated dissolved O$_2$ sensor (SBE43I, Sea-Bird Electronics Inc.) and a full bio-optical system mounted at the very top of the endcap of the profiler. The bio-optical package included an irradiance detector (PAR-LOG, Satlantic Inc.) that recorded the intensity of the photosynthetically active radiation (PAR) and a customized ‘triplet’ fluorometer (ECO FLbb-CD, WETLabs Inc.) that measured chlorophyll fluorescence (Chl-$a$), colored dissolved organic matter fluorescence and optical backscatter (Laney et al., 2014). All bio-optical sensors were calibrated separately as described by Laney et al. (2014). Calibration details of oth-
er sensors on the profilers are summarized in Krishfield et al. (2008b). PAR sensor was not calibrated or had other problems.

Figure 2.4. Sensors deployed with the SAMI or on the profiler: Aanderaa oxygen sensor (on SAMI) (top left), Microcat CTD sensor (on SAMI) (top right), Satlantic PAR sensor (on profiler) (bottom left), and WETLabs Chl-a sensor (on profiler) (bottom right).

2.3. Ancillary Data

A variety of other data were collected from different platforms or Internet sources. The wind speed data near ITP locations were obtained from the European Center for Medium range Weather Forecasting (ECMWF) website: http://apps.ecmwf.int/datasets/data/interim-full-daily. Barometric pressure and air temperature data were obtained from the Ice-Mass Balance (IMB) Buoy website: http://imb.erdc.dren.mil/2012L.htm. Hourly average mole fractions of atmospheric CO₂ (xCO₂) measured at Barrow, Alaska were obtained from the National Oceanic and Atmospheric Administration (NOAA) Earth System Research Laboratory (ESRL) website: http://www.esrl.noaa.gov/gmd/dv/ data/. Ice coverage data were obtained from the National Snow and Ice Data Center (NSIDC): http://nsidc.org/data. The PAR data from the sensors mounted on the ITPs was uncalibrated, so we used surface solar radiation
that corresponded to the ITP locations computed at 7.5 minute intervals. The PAR data were adjusted for light loss and scattering assuming an attenuation of 40% at ~9 m (Penta et al., 2008).

2.4. Data validation

An underway $p$CO$_2$ equilibrator-infrared measurement system (SUPER-CO$_2$, Sunburst Sensors) (Fig. 2.5) provided validation data during the ITP deployment. In order for SUPER-CO$_2$ to be used as a reference for the SAMI, it required correction for warming inside the water line and equilibrator. The SUPER-CO$_2$ values were therefore corrected to the sea surface temperature using the following formula (Dickson et al., 2007):

$$ (p\text{CO}_2)_{T_s,\text{wet}} = (p\text{CO}_2)_{T_e,\text{wet}} \times \exp[0.0423(T_s - T_e)] \quad (2.1) $$

where $T_s$ is the sea surface temperature, $T_e$ is the temperature measured in the SUPER-CO$_2$ equilibrator.

After correction factors were applied on all SUPER $p$CO$_2$ values they were further compared to SAMI $p$CO$_2$ values collected during the time the ship was at the ITP location. SAMI offsets were corrected accordingly. Once the initial shipboard data verifies SAMI accuracy at the time of deployment, there are very few mechanisms that can lead to drift and inaccuracy later on. Few primary known mechanisms include certain types of electronic problems, poor pump flushing, and biofouling. Issues related to electronics and pump flushing can be readily diagnosed via metadata transmitted with each SAMI measurement. Biofouling is not a significant problem with SAMI over the deployment period.
Figure 2.5. The SUPER-CO$_2$ system (Sunburst Sensors, LLC) was used to quality control the SAMI data. It is designed for automated shipboard analysis of seawater and atmospheric $p$CO$_2$. SUPER-CO$_2$ can take measurements at subsecond temporal resolution. Image obtained from Sunburst Sensors website.

Dissolved O$_2$ from both systems (SAMI and profiler) were quality controlled using the following procedure. The surface SAMI and profiler DO values at the beginning of the ITP deployment were corrected for offsets by comparing with surface shipboard DO measurements performed during the deployment (Winkler titration method). After the ship left the ITP location, the surface SAMI DO data were corrected for sensor drift by referencing to the profiler data corrected to the deep-water isopycnal (constant density) values (Timmermans et al., 2010). The QC procedure is based on the assumption that DO variability on deep isopycnals is negligible (Timmermans et al., 2010). For this purpose, the deep DO profiler values were first corrected with deep bottle data from the time of deployment and all subsequent deep values were corrected to these values by applying an offset, then the surface SAMI DO values were compared with the surface profiler values and corrected for offset.

SAMI temperatures were compared with SAMI-CTD temperatures and corrected for offset. Salinity measured by SAMI-CTD and profiler surface CTD agreed well so did not require correction.
2.4.1. Partial Pressure of carbon dioxide (\(p\text{CO}_2\))

Temperature-corrected SUPER \(p\text{CO}_2\) values and SAMI \(p\text{CO}_2\) values at the beginning of the measurement were compared and a difference of 75 µatm and 21 µatm were observed for SAMI-12 and SAMI-14, respectively. In both cases, SAMIs were overestimating \(p\text{CO}_2\) data and the data were corrected. The SAMI-12 had a leaky reagent bag, which was replaced prior to deployment and likely led to the large offset. After offsets were applied on all SAMI \(p\text{CO}_2\) values, the data were further compared to SUPER \(p\text{CO}_2\) values during the time the ship was at the SAMI locations. The mean difference ± standard deviation between the corrected SAMI \(p\text{CO}_2\) and SUPER \(p\text{CO}_2\) values were 1.3 ± 0.1 µatm (n=3) for ITP-64 and 0.3 ± 0.3 µatm (n=5) for ITP-65 (Fig. 2.6).

**Figure 2.6.** Discrete \(p\text{CO}_2\) measured underway by SUPER system was used to quality control the SAMI \(\text{CO}_2\) data. The blue (ITP-64) and black (ITP-65) filled symbols are SAMI \(p\text{CO}_2\) data and the open red symbols are SUPER \(p\text{CO}_2\) data. Corrected SAMI data overlaps SUPER data while the ship was at the ITP deployment location. The ship was moving to other locations between ITP deployments.
2.4.2. Dissolved Oxygen Concentration (DO)

As stated above, the optodes on SAMI-12 and SAMI-14 measured DO at 6 m depth at 2 hour intervals. Profiler-O₂ sensors (O₂ sensors on ITP-64 and ITP-65) took measurement along the water column down to ~800 m depth, but at ~11 m depth the measurements had 6-hour intervals. SAMI DO values at the beginning of the deployment were lower than the Winkler (ship) values by 31 µmol/kg and 23 µmol/kg for SAMI-12 and SAMI-14, respectively. Therefore, offset correction was applied to both data sets. The offset corrected SAMI DO data were then compared with offset and drift corrected ITP surface DO data (see next paragraph) to check for possible drift. A constant linear drift was observed with both SAMI-12 and SAMI-14 DO data (Fig. 2.7).

![Figure 2.7. Profiler surface DO (red) and SAMI DO (black) at the beginning of the measurement were compared to surface shipboard DO measurements (blue) performed at the time of the deployment. The top figure refers to ITP-64 and bottom figure refers to ITP-65. After correcting to the bottle data, SAMI DO was corrected to the profiler data. The SAMI DO data appear noisier than the profiler data because they measured every 2 hours, capturing the short-term variability. The profiler was at the surface only ~4 times per day.](image-url)
Comparison of the profiler DO data with the shipboard data found an offset of 16 μmol/kg for ITP-64 and 2 μmol/kg for ITP-65. In both cases, the profiler DO values were lower than the ship DO values and were corrected accordingly. The offset-corrected profiler surface DO values were then compared with deep DO values, as described above. Based on this analysis, the optode sensors on the ITPs drifted by ~10 μmol/kg or less relative to the profiler data and offsets were applied accordingly (Fig. 2.7).

**Figure 2.8.** ITP-64 (left) and ITP-65 (right) profiler surface DO and SAMI DO correlation plots. Black lines refer to data correlation lines and gray lines refer to 1:1 lines. After offset and drift correction profiler surface DO and SAMI DO had $r^2$ values of 0.44 and 0.60, respectively, and a slope of 0.45 and 0.70, respectively. SAMIs measured DO at 6 m depth at 2 hour intervals and profilers measured DO at ~11 m depth at 6 hour intervals.

After applying offset and drift correction on SAMI and ITP DO data, the data sets better agree with each other (Fig. 2.8). There is considerable scatter primarily because this is very narrow range of DO variability, so any noise in the measurements would translate into scattering of data. Also, at times the SAMI and the profiler O$_2$ sensors might have encountered slightly different masses of water due to local processes at different depths (the two
instruments had <0.5 m depth difference). A correlation plot between SAMI-12 DO and profiler DO values gives $r^2 = 0.44$ and a slope of 0.45, and, that between SAMI-14 DO and profiler DO values gives $r^2 = 0.60$ and a slope of 0.70. After the correction the SAMI and shallowest ITP profile DO data compared to within ±0.8 and ±0.7 μmol kg$^{-1}$ for ITP-64 and ITP-65, respectively.

2.4.3. Temperature

SAMI-CO$_2$, surface CTD, and profiler CTD, all measured temperature. The first two took measurements at 2 hour intervals and the latter at 6 hour intervals at its shallowest depths (~11 m). Both SAMIs gave a constant temperature offset. Their temperatures were always 0.16 °C lower than the temperature measured by the CTDs mounted on them. The SAMI temperatures were corrected accordingly for offset because the CTD temperature was considered more accurate. After offset correction, the temperature data from both sensors agreed well (Fig. 2.9) with the mean difference and the standard deviation being small. A correlation plot between SAMI-12-CO$_2$ temperature and SAMI-12-CTD temperature gives $r^2 = 0.87$ and a slope of 0.90, and that between SAMI-14-CO$_2$ temperature and SAMI-14-CTD temperature gives $r^2 = 0.85$ and a slope of 0.84 (Fig. 2.9). After the correction, the SAMI and CTD data compared to within ±0.009 and ±0.008 °C for ITP-64 and ITP-65, respectively. The SAMI-CO$_2$ temperatures were compared only with the SAMI-CTD temperatures because the CTD temperature was considered more accurate and the profiler CTDs were at different depths than the SAMIs. The profiler CTDs also had less measurement frequency at their shallowest depths compared to the SAMIs.
Figure 2.9. ITP-64 (left) and ITP-65 (right) CTD and SAMI temperature correlation plots. After correcting SAMI temperature for offsets the $r^2$ values were 0.87 and 0.85, respectively, and slopes were 0.90 and 0.84 respectively. Both SAMI and CTD on the SAMI took measurements at 2 hour intervals.

2.4.4. Salinity

SAMI-CTD and profiler-CTD measured salinity at 6 m and ~11 m depth at 2 hour and 6 hour intervals, respectively. The data from the sensors agreed well so did not require any correction (Fig. 2.10). SAMI-CTD and profiler-CTD data compared to within ± 0.006 and ±0.008 units for ITP-64 and ITP-65, respectively. A correlation plot between ITP-64 salinity and SAMI-12 salinity gives $r^2 = 0.62$ and a slope of 0.62 and that between ITP-65 salinity and SAMI-14 salinity gives $r^2 = 0.98$ and a slope of 1.0. The correlation is 0.92 with a slope of 0.95 for ITP-64 and 0.99 with a slope of 1.0 for ITP-65 (Fig. 2.10) until the water approached the freezing temperature.
Figure 2.10. ITP-64 (left) and ITP-65 (right) profiler surface salinity and SAMI salinity correlation plots. The salinity data did not need offset and drift correction. The $r^2$ values until the water reached the freezing temperature are 0.92 and 0.99 for ITP-64 and ITP-65, respectively, and the slopes are 0.95 and 1.0, respectively. SAMI-CTD and profiler-CTD measured salinity at 6 m and ~11 m depth at 2 hour and 6 hour intervals, respectively.

2.4.5. Bio-optical parameters

Chl-$a$ data were obtained by direct conversion of digital counts to concentration according to the manufacturer calibration sheets. Chl-$a$ data from both profilers contained a number of outliers so appropriate ranges of tolerance were applied to both data sets to filter out the outliers. The surface solar radiation corresponding to the ITP locations was scaled using the surface PAR data collected on the ship to account for the nearly constant cloud cover and absorption due to seawater to 9 m.

2.5. Calculations

2.5.1. Water Density

Water density was calculated using *in situ* temperature and salinity data at corresponding depths. First, the density is calculated at one standard atmospheric pressure. Then the secant bulk modulus is calculated to get density under a given condition. The calculation is an
implementation of the International Equation of State of Sea Water, 1980 (IES80) taken from "Introductory Dynamical Oceanography" by Pond and Pickard (Appendix 3, pp 310-311).

2.5.2. Freezing Temperature of Water

In order to observe when ice formation could occur, the freezing point of seawater was calculated using in situ salinity data (Millero and Leung, 1976) according to the formula in Millero and Leung (1976).

2.5.3. Mixed Layer Depth (MLD)

The mixed layer depth (MLD) was calculated from depth-resolved density and temperature, with MLD as the depth where the density difference from the SAMI depth is +0.3 kg m\(^{-3}\) (Fig. 3.2). This value is close to the value of 0.26 kg m\(^{-3}\) found by Timmermans et al. (2010) using the same approach for the central Canada Basin. We used a density difference of 0.3 kg m\(^{-3}\) because it gives computed MLDs that best match in situ MLDs when observed visually based on the water column density profile.

2.5.4. Total Alkalinity (A\(_T\))

Alkalinity was derived from a salinity-total alkalinity (A\(_T\)) relationship for the Chukchi Sea, Beaufort Sea, and Canada Basin (Yamamoto-Kawai et al., 2005; Fig 2.11). The relationship was corrected for the influence of sea ice melting/formation and has a coefficient of 0.99. The corrected relationship was used; however, there are various mechanisms such as variable contributions of river runoff, sea ice meltwater (Ulfsbo et al., 2014), carbonate mineral precipitation or dissolution (Cross et al., 2013) that can introduce significant uncertainties to this conservative relationship.
Figure 2.11. Historical Salinity-$A_T$ (shown as TA in this figure) relationship (right) for the Canada Basin and Beaufort Sea (Yamamoto-Kawai et al., 2005). The left figure refers to the location the salinity and alkalinity data were collected from.

Ulfsbo et al. (2014) found a mean difference between the measured and computed $A_T$ of $4 \pm 23 \mu$mol kg$^{-1}$. The relationship overestimated $A_T$ by $\sim 30 \mu$mol kg$^{-1}$ in the Canada Basin and near the Laptev Sea, where the discrepancies were the largest. Our $A_T$ was computed from ITP-64 and ITP-65 salinity using the following equation based on the historical Salinity-$A_T$ relationship for the Canada Basin and Beaufort Sea by Yamamoto-Kawai et al. (2005):

$$A_T = 62.47 \times \text{Salinity} + 261.34$$  \hspace{1cm} (2.2)

where $A_T$ is in $\mu$mol kg$^{-1}$ and S is in situ salinity (unitless).

2.5.5. Normalized DIC (nDIC)

Normalized DIC reduces the variability in DIC caused by riverine input of alkalinity and advective process. The normalization of DIC to a constant salinity (35) (Fig. 4.5) was done according to following formula:
\[ n\text{DIC} = \text{DIC} \times \left(\frac{35}{\text{Salinity}}\right) \]  

(2.3)

where DIC is \textit{in situ} DIC (in \( \mu \text{mol kg}^{-1} \)) and Salinity is \textit{in situ} Salinity.

**2.5.6. Equilibrium O$_2$ Concentration in Seawater**

Saturated DO is the dissolved amount of O$_2$ in seawater in equilibrium with the atmosphere, but it is the partial pressure that controls the gas exchange. Consequently, the water vapor reduces the partial pressure of O$_2$ and needs to be accounted for in gas exchange calculations. DO saturation values were calculated using \textit{in situ} temperature, salinity and barometric pressure with water vapor pressure accounted for using equations in Gnaiger and Forstner (1983). Salty water vapor pressure was calculated at \textit{in situ} temperature and salinity and O$_2$ solubility was calculated from temperature, salinity and barometric pressure (Gnaiger and Forstner, 1983). Equilibrium O$_2$ concentration was then calculated using the following formula:

\[ \text{O}_2 \text{ (equilibrium O}_2\text{ concentration at \textit{in situ} T, S and P) = O}_2 \text{ (O}_2\text{ saturation at \textit{in situ} T and S) \times P[(1-P_w/P) (1-\Theta P)] / [(1-P_w)(1-\Theta)]} \]  

(2.4)

where, T is temperature (in °C), S is salinity (unitless), P is barometric pressure (in atm), \( P_w \) is the water vapor pressure (in atm), and

\[ \Theta = 0.000975 - (1.426 \times 10^{-5} \times T) + (6.436 \times 10^{-8} \times T^2) \]  

(2.5)

**2.5.7. Atmospheric pCO$_2$ at the Sea Surface**

\( \times CO_2 \) data downloaded from the NOAA-ESRL website were multiplied by local barometric pressure to get pCO$_2$ in dry air. Atmospheric pCO$_2$ in wet air was then calculated by
including water vapor pressure (Dickson et al., 2007) according to the following formula (Dickson et al., 2007):

\[ p(\text{CO}_2 \text{ at sea surface}) = p(\text{CO}_2 \text{ in dry air}) \times (1 - P_w) \]  

(2.6)

where, \( P_w \) is the water vapor pressure over the seawater sample of \textit{in situ} salinity at the temperature of equilibration (Gnaiger and Forstner, 1983).

2.6. Modeling

2.6.1. Partial Pressure of carbon dioxide \((p\text{CO}_2)\)

When carbon dioxide enters seawater it reacts with the water. The reactions can be represented by the following series of equilibria:

\[ \text{CO}_2 (g) \rightarrow \text{H}_2\text{CO}_3^* \text{ (aq)} \]  

(2.7)

\[ \text{H}_2\text{CO}_3^* \text{ (aq)} \leftrightarrow \text{H}^+ \text{ (aq)} + \text{HCO}_3^- \text{ (aq)} \]  

(2.8)

\[ \text{HCO}_3^- \text{ (aq)} \leftrightarrow \text{H}^+ \text{ (aq)} + \text{CO}_3^{2-} \text{ (aq)} \]  

(2.9)

It is difficult to analytically distinguish between \( \text{CO}_2 \text{ (aq)} \) and \( \text{H}_2\text{CO}_3 \text{ (aq)} \) so both are combined and the sum is expressed as the concentration of a hypothetical species, \( \text{H}_2\text{CO}_3^* \text{ (aq)} \).

2.6.2. Dissolved Inorganic Carbon (DIC)

Total dissolved inorganic carbon (DIC), defined as the sum of concentrations of the different carbon species i.e. \( \text{HCO}_3^- \), \( \text{CO}_3^{2-} \), \( \text{H}_2\text{CO}_3^* \) in seawater, was calculated using CO2SYS (Fig. 4.5) developed for \( \text{CO}_2 \) system calculations (van Heuven et al., 2011): http://cdiac.ornl.gov/ftp/co2sys/. In our calculations, CO2SYS takes temperature, salinity, depth, \( p\text{CO}_2 \) and salinity-derived alkalinity \( (A_{\text{Tsain}}) \) (protolytic nutrients e.g. phosphate are assumed to be negligible) as input and returns DIC and other \( \text{CO}_2 \) parameters as output, in-
cluding aragonite saturation ($\Omega_{\text{arag}}$). The saturation state of seawater with respect to aragonite (a form of calcium carbonate found in plankton and invertebrates) can be defined as the product of the concentrations of dissolved calcium ions ($\text{Ca}^{2+}$) and carbonate ions ($\text{CO}_3^{2-}$) in seawater, at the *in situ* temperature, salinity, and pressure, divided by the stoichiometric solubility product ($K_{sp}^*$) for those conditions, according to the following equation:

$$\Omega_{\text{arag}} = \frac{[\text{Ca}^{2+}] \times [\text{CO}_3^{2-}]}{K_{sp}^*}$$  \hspace{1cm} (2.10)

### 2.6.3. Dissolved Oxygen Concentration (DO)

$\text{CO}_2$ and $\text{O}_2$ are intimately related through photosynthesis and respiration. Photosynthesis is the production of organic matter from inorganic carbon and respiration is the destruction of organic matter by reaction with oxygen in the presence of sunlight. The photosynthetic reaction, which represents the average stoichiometry of organic matter production based upon analysis of seawater, is:

$$106 \text{CO}_2 + 16 \text{NO}_3^- + \text{HPO}_4^{2-} + 122 \text{H}_2\text{O} + 18 \text{H}^+ \rightarrow \text{Organic matter} + 138 \text{O}_2$$  \hspace{1cm} (2.11)

In the process of photosynthesis, $\text{O}_2$ is produced while $\text{CO}_2$ is consumed, thus making the $\text{O}_2$ profile potentially a mirror image of the $\text{CO}_2$ profile in seawater. Therefore, DO is a very valuable counterpart to $\text{CO}_2$ as the two can be modeled using similar processes and divergence from model predictions or expected correlations provides additional insight into the controlling processes of the carbon cycle.

### 2.6.4. Temperature, Salinity

Sea surface temperature can be used as an indicator of various phenomena going on in the seawater based on the assumption that the anomalies of sea-surface temperature i.e. devia-
tion from the mean, indicates potential mixing of different water masses and sea surface heating and cooling due to air-sea exchange of heat and absorption of solar radiation. In the AO surface water, however, temperature hardly varies and is not a strong indicator of these processes.

In the AO, salinity differences in seawater tend to cause stronger stratification inhibiting vertical mixing of seawater in the water column. Evaporation (not prominent in the AO) and ice formation can cause an increase in salinity. When the seawater freezes it excludes salts from the ice through a process called “brine rejection” and the surrounding seawater becomes saltier and denser, sinking into the water column causing mixing. Saline water also holds less gas because some of the water molecules are occupied to dissolve the salt and these more concentrated dissolved gases sink down with the brine. Precipitation (rain) and ice melt are other sources of salinity variability causing a decrease in salinity by increasing the proportion of H₂O and diluting the concentrations of salts. Because these effects reduce density, they are more persistent at the air-sea interface and require a mixing mechanism such as wind or ice shear to dissipate.

2.6.5. Model Calculations

2.6.5.1. DIC and DO flux

The flux (changes over time) of DIC and DO potentially originate from 4 different processes in the AO as described in the following mass balance:

\[ H \times \Delta C/\Delta t = F_{\text{gasex}} + F_{\text{NCP}} + F_{\text{mix}} + F_{\text{brine}} \]  

(2.12)

where H is the mixed layer depth, \( \Delta C \) is the difference in DIC or DO values between two measurements over time \( \Delta t \), and \( F_{\text{gasex}}, F_{\text{NCP}}, F_{\text{mix}}, \) and \( F_{\text{brine}} \) are the fluxes due to gas transfer
across the air-water interface, net community production (NCP), vertical mixing and advection, and brine rejection during ice-formation, respectively. We assume CaCO₃ dissolution and formation is not significant. If each of these processes can be modeled using established relationships and physical data, it is possible to quantitatively predict the overall variability in DIC and DO (e.g. DeGrandpre et al., 2004). In this evaluation, we focus on calculation of \( F_{\text{gasex}} \) and \( F_{\text{NCP}} \) and the contribution of the other fluxes, which are more difficult to model, are estimated by examining the residual variability. The \( F_{\text{gasex}} \) of CO₂ and DO across the air-water interface was calculated using the diffusive boundary layer model:

\[
F_{\text{gasex}} = k \Delta C (1-f) \tag{2.13}
\]

where \( k \) is the gas transfer velocity and \( f \) is the fractional ice coverage. \( k \), which accounts for the rate of gas diffusion through the air-sea boundary layer, was estimated using a wind speed relationship and adjusted for different temperatures and gases (i.e., CO₂, DO) using the updated equation in Wanninkhof (2014):

\[
k = 0.251 \langle U^2 \rangle \left( \frac{Sc}{660} \right)^{-1/2} \tag{2.14}
\]

where \( \langle U^2 \rangle \) is average squared wind speed adjusted to 10 m height and \( Sc \) is the Schmidt number which accounts for differences in molecular diffusivity between gases. A negative \( F_{\text{gasex}} \) represents a flux from the atmosphere to the ocean.

Since air-sea gas flux in the Canada Basin is expected to be virtually insignificant for the most of the year due to significant ice cover, annual uptake of CO₂ (in Tg C yr⁻¹) was calculated as the product of late summer-autumn CO₂ flux (in mmol m⁻² d⁻¹) and the corresponding area (in km²). We assumed that gas flux is significant over a 90 day season when open water was significant.
We interpret short-term salinity changes that return to an earlier baseline to be primarily due to different water masses that advected past the non-Lagrangian (i.e. it did not follow the water) drifter (Timmermans et al., 2012). Vertical mixing is assumed negligible due to the strong halocline (salinity stratification), as discussed below. The DIC and O₂ flux from brine formation were not modeled but their importance in these data can be assessed by comparison of the DO and DIC time-series, as discussed below.

We evaluated contributions of NCP in three ways: 1) by fitting the residual signal not accounted for by gas exchange; 2) revealing the diurnal signal by computing bandpass filtered DIC and DO, after removing the contribution of $F_{\text{gasex}}$ and 3) by using a photosynthetic model (explained below). To obtain the bandpass filtered data, 4 hour low pass and 30 hour high pass filters were used to pull out the diurnal signal. The output is the hourly rate of change in DIC and DO. NCP from DO was compared with NCP from DIC by conversion with the photosynthetic quotient 1.4 (see Equation 2.11). Gross primary production (GPP) was then calculated by adding nighttime respiration to daytime NCP. Because influence of non-local advection appear in the NCP data (Fig. 4.4) as different water masses often contain different levels of CO₂ and DO, DIC and DO in seawater can change disparately when there are potential influence from non-local processes (DeGrandpre et al., 1998). This is discussed in more detail below. Data corresponding to those time periods were omitted in the daily mean of NCP and GPP (Fig. 4.4), as discussed below.

**2.6.5.2. Gas Exchange and NCP model**

The mass balance (Equation 2.12) used two biogeochemical models for $p\text{CO}_2$ and DO. The $p\text{CO}_2$ model was generated using an in-house program for carbonate system calculations.
The model takes the initial DIC and $A_{\text{Tsalin}}$ at \textit{in situ} temperature and salinity in the surface seawater as the initial input. DIC and $A_{\text{Tsalin}}$ initial were 1878 and 1975 $\mu$mol kg$^{-1}$, respectively, for ITP-64, and 1904 and 2003 $\mu$mol kg$^{-1}$, respectively, for ITP-65. A constant $A_T$ was then used essentially removing all DIC variability due to water mass changes within the model. The DIC was incremented at each time step for contributions from air-water gas exchange using the calculated $pCO_2$. The new $pCO_2$ was then solved from the $A_T$, \textit{in situ} temperature and salinity, and the new DIC. This calculation gives the modeled variability of $pCO_2$ from gas exchange. The remainder of the variability after contribution from gas exchange was assumed to be NCP. A residual least squares approach was used to best fit the DO data. Using this NCP and adjusting for the photosynthetic quotient (1.4), gave a reasonable fit to the $pCO_2$ data. Since the program calculated the carbonate equilibria and solubility constants as a function of temperature, the effects of heating and cooling on the $pCO_2$ are intrinsic in the model. A similar approach was used in the DO model. Because DO exists in only one dissolved from, the modeling of DO was comparatively simple. Both $pCO_2$ and DO gas exchange models were calculated assuming ice-free conditions as well with the goal to show to what extent AO ice coverage controls $pCO_2$ and DO variability.

2.6.5.3. Photosynthetic model

Net primary production (NPP) was calculated with an exponential model (Platt \textit{et al.}, 1980, Cullen \textit{et al.}, 1992) that takes into account the light (PAR) and Chl-$a$ dependence of photosynthesis:

$$NPP = [\text{Chl}-a \ (P_s^{\text{Chl}-1})(1 - e^{-\alpha})] \ 0_p^{(T-20)}$$

(2.15)

where NPP is in mg C m$^{-3}$ h$^{-1}$ (C is carbon), Chl-$a$ is the concentration of chlorophyll-$a$ in mg m$^{-3}$, $P_s^{\text{Chl}-1}$ is the Chl-$a$–specific maximum rate of photosynthesis in the absence of photoin-
hibition (mg C mg Chl-α⁻¹ hr⁻¹), T is the *in situ* temperature in °C, θ_P is the Arrhenius temperature coefficient (1.036) and

\[ a = (\alpha Q_{PAR} (P_a \text{Chl}^{-1})^{-1}) \]  \hspace{1cm} (2.16)

where α is the Chl-α specific rate of light-limited photosynthesis (in mg C mg Chl-α⁻¹ hr⁻¹ (μmol quanta m⁻² s⁻¹)⁻¹), and Q_{PAR} is the downwelling PAR (in μmol quanta m⁻² s⁻¹).

Community respiration (R₁) was estimated using the formula below:

\[ R_1 = R \theta_R (T-20) \]  \hspace{1cm} (2.17)

where R is the nighttime respiration rate based on DO, θ_R is the Arrhenius temperature coefficient (1.045), and T is the *in situ* temperature in °C. We observed that, with a fixed respiration rate, respiration was significantly overestimated over the time-series as the production decreased with decreasing sunlight. Respiration in the dark Arctic is very small (Sherr and Sherr, 2003), therefore, it was assumed to be proportional to NPP where the daily mean of respiration is equal to the mean of NPP on that day, tightly linking NPP and respiration as expected in nutrient-depleted surface waters (Dugdale and Goering, 1967). The new respiration is:

\[ R_2 = R_1 \times (\text{NPP} \times k) \]  \hspace{1cm} (2.18)

where k is a proportionality constant (0.4) obtained by a residual sum of squares fit. NCP was then calculated as the difference between NPP and R₂.
Chapter 3

Results

The 49-day time-series data for ITP-64 and 41-day time-series data for ITP-65 are shown in Fig. 3.1. Gaps in the data correspond to instrument data communication dropouts between the SAMI and inductive modem. Corresponding physical data are shown in Fig. 3.2. The ice cover near ITP-64 ranged from 0 to 32% with a mean of 10 ± 10%, and ITP-65 ice cover ranged from 38.4 to 89.2% with a mean of 63.2 ± 15.8% (Fig. 3.1e, Table 3.0). Over the deployment time periods, the $pCO_2$ was well below atmospheric saturation and very similar for both locations. ITP-64 $pCO_2$ ranged from 282 to 323 µatm with a mean of 306 ± 9 µatm, and ITP-65 from 272 to 330 µatm with a mean of 304 ± 16 µatm (Fig. 3.1a, Table 3.0). ITP-64 $pCO_2$ steadily increased from ~285 to 310 µatm until around 09/23/12 when it leveled off (Fig. 3.1a). DO showed very little variability at both locations but were above or near saturation for the most of the time period (Fig. 3.1b, saturation is not shown. The saturation level ranged between 100.8 and 105.2% with the mean being 102.8 ± 1% for ITP-64, and between 99.2 and 103.3% with the mean being 101.3 ±1% for ITP-65). ITP-64 DO ranged from 387 to 395 µmol kg$^{-1}$ with a mean of 392 ± 2 µmol kg$^{-1}$ and ITP-65 from 381 to 390.3 µmol kg$^{-1}$ with a mean of 385 ± 2 µmol kg$^{-1}$ (Fig. 3.1b, Table 3.0). DO levels were above or near saturation for the most of the time period. The diurnal cycles are distinct in ITP-64 DO data (Fig. 3.1b) until the daylight period approached zero around 10/06/12 (Fig. 4.3), whereas in ITP-65 DO data they are almost absent (Fig. 3.1b). Diurnal cycles in DIC at ITP-64 are also evident once the long-term trend is removed (see below).
Figure 3.1. Biogeochemical and physical time-series data collected at ~6 m on ITP-64 (black) and ITP-65 (red) (panels a-d). ITP-64 was deployed under little ice whereas ITP-65 was deployed under multi-year ice (panel e). Gaps in the data correspond to instrument communication dropouts. The light black line and light red line in the temperature plot refer to freezing temperature of water at ITP-64 and ITP-65, respectively (panel d). Note that the colorbar scale changes from panel f to g.
ITP-64 temperature was slightly higher than ITP-65 temperature until they converged towards the freezing point (Fig. 3.1c). The freezing point of seawater was ~0.03 °C higher at ITP-65 due to the significantly higher salinity. ITP-65 approached freezing temperature around 09/21/12, a couple of days earlier than ITP-64 (Fig. 3.1c). The difference in salinity between the two ITPs locations was relatively large considering the distance between the locations. ITP-64 salinity had a mean value of 27.5 ± 0.1, whereas ITP-65 had a mean value of 28.0 ± 0.1 (Fig. 3.1d, Table 3.0). Over most of the measurement period, ITP-64 Chl-α was ~3 times higher than ITP-65 Chl-α (Fig. 3.1f, 3.1g; Table 3.0). The atmospheric $p$CO$_2$ measured in Barrow, Alaska ranged from 375.4 to 404.8 μatm with a mean of 385.6 ± 5.8 μatm. The average depth of mixed layer was 21.5 ± 2.0 m at ITP-64 and 22.0 ± 2.5 at ITP-65 (Fig. 3.2).

Table 3.0 The mean ± standard deviation of the difference for different parameters for ITP-64 and ITP-65 at ~6 m depth. The right-most column includes the difference between the mean of the parameters.

<table>
<thead>
<tr>
<th>Parameters</th>
<th>ITP-64</th>
<th>ITP-65</th>
<th>Mean Difference (ITP-64 - ITP-65)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ice coverage (%)</td>
<td>10 ± 10</td>
<td>63.2 ± 15.8</td>
<td>-53.2</td>
</tr>
<tr>
<td>Temperature (°C)</td>
<td>-1.46 ± 0.05</td>
<td>-1.50 ± 0.03</td>
<td>0.04</td>
</tr>
<tr>
<td>Salinity</td>
<td>27.5 ± 0.1</td>
<td>28.0 ± 0.1</td>
<td>-0.5</td>
</tr>
<tr>
<td>$p$CO2 (μatm)</td>
<td>306 ± 9</td>
<td>304 ± 16</td>
<td>2.4</td>
</tr>
<tr>
<td>Δ$p$CO2 (μatm)</td>
<td>-79.5 ± 9.5</td>
<td>-80 ± 15</td>
<td>-0.5</td>
</tr>
<tr>
<td>CO2 flux (mmol m$^{-2} \cdot$ d$^{-1}$)</td>
<td>-7.8 ± 6.8</td>
<td>-2.5 ± 2.6</td>
<td>5.3</td>
</tr>
<tr>
<td>DO (μmol kg$^{-1}$)</td>
<td>392 ± 2</td>
<td>385 ± 2</td>
<td>7.4</td>
</tr>
<tr>
<td>ΔDO (% saturation)</td>
<td>3.0 ± 0.1</td>
<td>1.3 ± 0.1</td>
<td>1.7</td>
</tr>
<tr>
<td>DO flux (mmol m$^{-2} \cdot$ d$^{-1}$)</td>
<td>15.6 ± 12.6</td>
<td>2.2 ± 2.7</td>
<td>13.4</td>
</tr>
<tr>
<td>Chl-α (μg L$^{-1}$)</td>
<td>0.28 ± 0.01</td>
<td>0.100 ± 0.002</td>
<td>0.18</td>
</tr>
<tr>
<td>NCP (mmol m$^{-2} \cdot$ d$^{-1}$)</td>
<td>7.4</td>
<td>~0</td>
<td>7.4</td>
</tr>
<tr>
<td>GPP (mmol m$^{-2} \cdot$ d$^{-1}$)</td>
<td>13</td>
<td>~0</td>
<td>13</td>
</tr>
</tbody>
</table>
Figure 3.2. Depth-resolved density, salinity and temperature contour plots for ITP-64 are shown in panel a, b and c, respectively, and those for ITP-65 are shown in panel d, e and f, respectively. MLDs (red line) are overlapped on the density (panels a and d) and salinity (panels b and e) contour plots. Wind speed is shown in panel e. Wind speed was uniform over both locations.
Chapter 4

Discussion

4.1. Observations based on in situ data and model estimates

The variability of CO₂ and DO was investigated based on various processes that contribute to the variability as described by Equation 2.12. Partially open-water and partially ice-covered in situ biogeochemical data are shown in Fig. 3.1. Biogeochemical models were employed to better explain the potential contribution from the major processes (Fig. 4.1, 4.5 and 4.6). We examine the contribution from gas exchange to the variability of CO₂ and DO in seawater under in situ ice cover and assuming ice-free condition, and then investigate the contribution of NCP to the variability. Other possible mechanisms that may have influence on the variability have also been assessed.

4.1.1. ITP-64

Results from biogeochemical models are compared with in situ pCO₂ and DO data in Fig. 4.1. The models reproduced the major features and overall long-term trends in pCO₂ and DO variability reasonably well. The model pCO₂ trend due to gas exchange initially matches the in situ data (r²=0.92), but overshoots after 9/23/12 (Fig. 4.1a). In situ pCO₂ maintained significant correlation with CO₂ gas flux (r²=0.75), which dropped dramatically after 09/23/12 (r²=0.08). The modeled gas exchange with no ice is very similar (Fig. 4.1) because there is very little ice coverage at this location (Fig. 3.1e). The DO model from gas exchange should in principle follow the DO data until 09/23/12 as with pCO₂, which it did not (Fig 4.1b). The DO maintained very weak correlation with pCO₂ until 09/23/12 (r²=0.001). This may be attributed to the relative saturation levels between pCO₂ and DO (26% pCO₂ and 3% DO) that
make the CO₂ gas exchange more dominant, and makes DO not correlate strongly with gas exchange. The correlation between pCO₂ and DO becomes stronger after 09/23/12 (r²=0.60) as phytoplankton production, though small later, continued to add DO to seawater and gradual increase in sea ice cover (Fig. 3.1e) resulted in accumulation of DO in water due to reduced sea to air flux of DO. Gas exchange drives modeled DO quickly to saturation and under predicts observed DO levels (Fig. 4.1b). These deviations from the in situ data indicate that there are some other process decreasing pCO₂ and increasing DO, counteracting the air-sea exchange.

**Figure 4.1.** ITP-64 pCO₂ and DO data (black) and model time-series data calculated from modeled gas exchange using the ice cover (blue solid) shown in Fig. 3.1 and without ice cover (blue dashed). Modeled gas exchange plus NCP assuming the residual signal is all NCP are shown (red). The green line corresponds to the equilibrium concentration of O₂ at in situ temperature, salinity and atmospheric pressure.
If the remainder of the variability is assumed to be NCP, a residual least squares approach finds an NCP of 8.6 mmol DO m$^{-2}$ d$^{-1}$ to best fit the ITP-64 DO data ($r^2=0.60$) (Fig. 4.1b, red curve). Using this NCP and adjusting for the photosynthetic quotient (1.4) gives a reasonable fit to the $p$CO$_2$ data ($r^2=0.55$) (Fig. 4.1a). These results suggest that biological production is an important contributor to $p$CO$_2$ and DO variability in this low ice-coverage gas record. This speculation is supported by the significant levels of Chl-a (Fig. 3.1f) and that there were detectable nutrients. The nitrate concentration in the ITP-64 mixed layer varied between 0.2 and 2 µM (Fig. 4.2). If all of these nutrients were utilized by photosynthesis, DIC at ITP-64 would go down by ~7 µmol kg$^{-1}$ and DO would go up by ~10 µmol kg$^{-1}$ based on Redfield stoichiometric calculations (Redfield, 1934) (Equation 4.1), manifesting the significant influence of nutrient-fueled biological activity on the variability of $p$CO$_2$ and DO at that location. According to the residual fit model, DIC at ITP-64 went down by 19 µmol kg$^{-1}$ and DO went up by 18 µmol kg$^{-1}$ due to NCP. The Redfield stoichiometric equation is:

$$C:N:P = 106:16:1 \quad (4.1)$$

where C is carbon, N is nitrogen, and P is phosphorous.

The NCP derived by the best-fit approach is close to the daily mean NCP, 7.4 mmol DO m$^2$ d$^{-1}$ and 7.0 mmol C m$^2$ d$^{-1}$ (C = CO$_2$), calculated from the bandpass filtered DO and DIC data (Fig. 4.4b). As can be seen in Figure 4.3, the filtered data more clearly show the diurnal pattern in the DO and CO$_2$ data. The NCP from DIC matches with the NCP from DO during some periods (e.g. $r^2=0.72$ from 09/04/12 to 09/07/12), however, at times they are poorly correlated. These times of poor correlation correspond to periods when salinity rapidly changed (marked by arrows in Fig. 4.4a) suggesting lateral advection of different water masses. The decoupling (non-Redfield variability) of CO$_2$ and DO may be due to advection of a
residual production signal where, for example, a bloom has occurred in the past. The DO more rapidly equilibrates with the surface ocean leaving behind a residual CO$_2$ signal (DeGrandpre et al., 1998). When water masses moved by the sensors, the DO did not vary as much as the pCO$_2$ (or DIC). These appear as large oscillations in the NCP calculated from DIC (Fig. 4.4b).

![Figure 4.2](image-url)

**Figure 4.2.** Nitrate concentration (red dot) in the mixed layer sampled at station CBN2 near ITP-64. The date and time for the first sampling are included on top of the figure. The samplings were done at 3 minute intervals. The average MLD at ITP-64 is 21.5 ± 2 m.

The photosynthetic model that calculated NCP using Chl-$a$ and PAR data (Equation 2.15, Fig. 4.3) found a mean NCP of 7.2 mmol C m$^{-2}$ d$^{-1}$. The modeled data was compared with bandpass filtered NCP data calculated from *in situ* DO and it reproduced the rate of change in DO in seawater considerably well ($r^2$=0.70) (Fig. 4.3), suggesting that the daily rate of change in DO in seawater was for the most part driven by biological activity i.e. photosynthesis. The model used an $\alpha$ value of 0.04 mg C mg Chl-$a^{-1}$ hr$^{-1}$ (µmol quanta m$^{-2}$ s$^{-1}$)$^{-1}$, which is the initial slope of the photosynthesis rate per unit water column and the instantaneous PAR intensity relationship. The $P_s$Chl value used in the model was 33 mg C mg Chl-$a^{-1}$ hr$^{-1}$, based on residual fit. Average nighttime respiration based on the rate of change in DO was 34 mmol m$^{-2}$ d$^{-1}$. 
Figure 4.3. ITP-64 modeled NCP calculated from Chl-a and PAR (red), and NCP calculated from *in situ* bandpass filtered DO (black, same data as shown in Figure 4.3). The model reproduces photosynthesis-driven rate of change in DO in seawater reasonably well ($r^2=0.70$). After 09/26/12, NCP from DO was significantly influenced by other (mostly advection) processes and modeled NCP became smaller and smaller as the length of daytime approached zero (PAR, lower figure). This model was not employed for ITP-65, as NCP was negligible based on the DO model results (see text). The PAR data shown here corresponds to solar radiation at ~8 m depth, which was initially scaled using the surface PAR data collected on the ship.

Although air-sea gas exchange and NCP explained $p$CO$_2$ and DO variability reasonably well and advection seemed to have significantly influenced the variability, there is some variability in the data that could not be explained, especially in the CO$_2$ record (Fig. 4.1). This variability may be linked to water movement. Timmermans *et al.* (2012) found that the ocean surface layer beneath sea ice in the Canada Basin is characterized by significant horizontal density gradients and concluded that submesoscale (a scale of intermediate size) processes significantly influence lateral density variability and the properties of the surface-layer. The surface waters in the central basin have distinctly different physical and chemical properties compared to the subsurface halocline, influencing the inventory of DIC and DO in the mixed layer. This is what likely created the large swings in the calculated NCP, as explained above.
Figure 4.4. ITP-64 salinity and biogeochemical data are shown in panel a-c and ITP-65 data are shown in panel d-f where NCP from DIC and DO are shown in red black, respectively. Abrupt changes in salinity data highlighted by downward arrow in panel a and d indicate potential advection of waters with different levels of $p$CO$_2$ and DO, thereby this short-term variability appears in the NCP calculated from the rate of change in DO (black) and DIC (red), shown in in panel b and e. NCP from DO is compared with NCP from DIC after conversion with photosynthetic quotient (1.4). Daily mean (circle) ± standard deviation (vertical bars) of NCP calculated from DO are shown in panel c for ITP-64 and f for ITP-65. The mean was calculated over a 24 hour period starting at 8 a.m.
Figure 4.5. Comparison between *in situ* DIC (black) and normalized DIC (red) for ITP-64 and ITP-65 are included in panel a and panel d, respectively. NCP calculated from DIC (black) are compared with NCP calculated from normalized DIC (red) in panel b and panel d, which correspond to ITP-64 and ITP-65, respectively. Normalized data differ considerably from the parent data suggesting that the influence of local and non-local process is significant at both ITP locations.
Others have observed that brine formation can be a significant contributor to mixed-layer and pycnocline DIC and DO levels (Nomura et al., 2006; Rysgaard et al., 2007). Ice formation could have occurred after ~09/23/12 based on the temperature. However, if $F_{\text{brine}}$ was significant DO and $pCO_2$ would simultaneously go up as the gases are rejected from the ice, which they did not (Fig. 4.1), indicating that brine formation was not a significant factor. All of these observations point to the conclusion that, apart from gas exchange and NCP, horizontal gradients was the only other process to significantly contribute to the $pCO_2$ variability, whereas DO variability for the most part was dominated by gas exchange and NCP.

### 4.1.2. ITP-65

We compared *in situ* $pCO_2$ and DO to modeled $pCO_2$ and DO calculated from gas exchange (Fig. 4.6) for data from ITP-65. The model initially fits the $pCO_2$ trend reasonably well ($r^2=0.60$ until 09/09/12) but does not fit well where there are humps in the *in situ* $pCO_2$ data (e.g. around 09/10/12, 09/26/12, 10/02/12). $pCO_2$ increased until 09/15/12 (Fig. 4.6a), then leveled off until 09/22/12, after when it varied up and down around a mean of ~315 $\mu$atm until the end of the measurement period. On the other hand, gas exchange explains almost all of the DO variability (Fig. 4.6b). The correlation coefficient between *in situ* DO and DO modeled from gas exchange is 0.53, which becomes 0.82 when the sudden changes around 09/08/12, 09/26/12 and 10/02/12 are ignored. DO maintained a mean of ~388 $\mu$mol kg$^{-1}$ until 09/06/12 (Fig. 4.6b), after this date it steadily went down to ~384 $\mu$mol kg$^{-1}$ and maintained this mean value until 09/23/12. After 09/23/12, DO varied up and down around the mean of ~384 $\mu$mol kg$^{-1}$ until the end of the measurement period. Based on the model, gas exchange keeps the DO near saturation. However, the contribution from gas exchange to the variability of $pCO_2$ and DO under heavy ice cover is small relative to that if no ice is present (Fig. 4.6).
Rutgers van der Loeff et al. (2014) found that gas transfer velocity in the ice-covered central AO regions is small (<0.42 cm hr\(^{-1}\)) when a residence time of at least 200 days is given. Therefore, CO\(_2\) and O\(_2\) exchange through ~63% ice cover at ITP-65 can be assumed small, which it is. \(p\)CO\(_2\) levels in seawater increase by 21.13 \(\mu\)atm, whereas DO decreases by 3.08 \(\mu\)mol kg\(^{-1}\) when ice-free condition is assumed (Fig. 4.6), manifesting that ice cover significantly inhibits gas exchange.

**Figure 4.6.** ITP-65 \(p\)CO\(_2\) and DO data (black) and model time-series data calculated from modeled gas exchange using the ice cover (blue solid) shown in Fig. 3.1 and without ice cover (blue dashed). Modeled gas exchange plus NCP assuming the residual signal is all NCP are shown (red). Green lines correspond to the equilibrium concentration of O\(_2\) at *in situ* temperature, salinity and atmospheric pressure.

The DO results based on gas exchange support that other sources of variability such as NCP, brine formation and vertical mixing are negligible. NCP is low as light under multiyear ice is too small and nutrients are too depleted (Fig. 4.7) to sustain significant production. NCP is low as it is primarily light-limited and availability of light under multiyear ice is too small to sustain significant production. There was no detectable nitrate in the ITP-65 mixed layer and Chl-\(a\) was small, almost 3 times smaller than ITP-64 Chl-\(a\). Although DO variabili-
ity for the most part can be explained by gas exchange, there are some big changes in $pCO_2$ (around 09/10/12, 09/26/12, 10/02/12; Fig. 4.6a) that gas exchange, NCP (NCP ~0), $F_{brine}$ could not explain. These changes are potentially linked to horizontal advection, as discussed in the case of ITP-64. These influences can be observed in the salinity data (highlighted by downward arrow, Fig. 4.4d), which suggests episodic mixing of water masses with different salinities (e.g. around 09/08/12, 09/26/12, 10/02/12). These observations indicate that horizontal gradients significantly dominated the $pCO_2$ variability under multiyear ice, whereas DO variability for the most part was dominated by gas exchange.

![Figure 4.7: Nitrate concentration in the ITP-65 mixed layer. The date and time for the first sampling are included on top of the figure. Samplings were done at 3 minute intervals. The average MLD at ITP-65 is 22.0 ± 2.5 m.](image_url)

### 4.2. Air-sea fluxes of CO$_2$

We compared our CO$_2$ flux estimates with previous estimates from the Canada Basin and nearby regions. Our estimated CO$_2$ flux during late summer-autumn is -7.8 ± 6.8 mmol m$^{-2}$ d$^{-1}$ for ITP-64 and -2.5 ± 2.6 mmol m$^{-2}$ d$^{-1}$ for ITP-65 (Fig. 4.8, Table 3.0). The range of variability at each site is primarily dominated by the gas transfer velocity (Equation 2.14) because of the highly variable wind speed (Fig. 4.8). Murata and Takizawa (2003) found that the
CO₂ flux in the slope waters of western AO during the summers of 1998-2000 ranged between −16.9 and −0.1 mmol m⁻² d⁻¹ and was −12 mmol m⁻² d⁻¹ in the western Beaufort Sea. In the spring of 2002 and 2004, Bates et al. (2006) estimated <−3 mmol m⁻² d⁻¹ CO₂ flux in the Canada Basin, when sea-ice cover ranged from 95% to 100%. During summertime, the fluxes in the Canada Basin were higher at ∼−55 mmol m⁻² d⁻¹, when sea-ice cover ranged from 0% to 90%. Our CO₂ flux estimates over the entire summer period (assuming 90 days of period when ice coverage is less) are −0.70 mol m⁻² for ITP-64 and −0.23 mol m⁻² for ITP-65. Since air-sea gas flux in the Canada Basin is expected to be virtually negligible for the most of the year due to significant ice-cover, we assume that our late summer-autumn flux values are reasonable estimates of annual CO₂ flux in the region. When the flux estimates are extrapolated over the entire Canada Basin with an area of 4,489,000 km² (Macdonald et al., 2009), the CO₂ flux is estimated to be −21.5 Tg C yr⁻¹ (Tg = teragrams = 10¹² grams) under ITP-64 condition (10 ± 10% ice cover) and −6.5 Tg C yr⁻¹ under ITP-65 condition (63 ± 16% ice cover). Bates et al. (2006) estimated the annual CO₂ flux in the Canada basin to be in the range of -6 to -19 Tg C yr⁻¹, appreciably close to our range. When the entire AO area of 10,700,000 km² (Macdonald et al., 2009) is considered, our flux estimates are -51.4 Tg C yr⁻¹ and -15.4 Tg C yr⁻¹ under ITP-64 and ITP-65 conditions, respectively. In comparison, annual flux estimates over the AO as reported by Anderson et al. (1994) (mass balance assessment), Lundberg and Haugen (1996) (mass balance assessment, includes Norwegian Sea), Anderson et al. (1998) (mass balance assessment), Anderson et al. (1998) (mass balance assessment, includes river contribution), Kaltin and Anderson (2005) (mass balance assessment), Bates (2006) are −70 ± 65, -110, -24 ± 17, −41 ± 18, -31, and -66 Tg C yr⁻¹, respectively. Our fluxes at ITP-64 are similar to those estimated by Evans et al. (2015) from shipboard pCO₂ data for
the western Arctic coastal ocean calculated from flux climatologies with the 2003 and 2014 sea ice, and 2003-2014 sea ice climatology.

Figure 4.8. CO₂ and O₂ gas flux across air-sea interface are shown for ITP-64 (black) and ITP-65 (red). Negative flux means air to sea gas transfer and vice versa. Fluxes for ITP-65 are considerably smaller than ITP-64 due to heavy ice coverage. The wind speed was uniform over both locations.
Our estimates and all these data show that CO₂ flux in the central AO is considerably variable in space and time. As a strong storm hit the Arctic in the early August of 2012, we compared the wind speed during our study to previous data to see how it compares to typical wind at the Canada Basin and if the post-storm period had a significant influence on our CO₂ gas flux estimate. During our measurement period, the wind speed ranged from 0.8 to 13.3 m s⁻¹ with the mean being 6.2 ± 2.7 m s⁻¹ (Fig. 4.8). According to NCEP reanalysis data, the average wind speed during 1992-2009 at the Canada Basin stayed fairly constant at about 5 m s⁻¹ during winter months October to January (JRA, ERA-Interim, and NCEP-2 ≥6 m s⁻¹), which reduced towards spring and summer (Spreen et al., 2011). During 2005-2006 (a winter storm hit in the winter of 2005-06) and 2007-2008, the strongest wind speed had a maximum daily average of over 14 m s⁻¹ and 10 m s⁻¹, respectively, with a daily average of about 10 m s⁻¹ during 2005-2010 time period (Jackson et al., 2012). The upper limit of the range of the wind speed during our measurement resembling that in 2005-2008 strong wind periods suggests that the wind during our study was strong and likely an aftermath of the storm that hit in the early August, and may have had significant influence on CO₂ gas flux, making the conditions different from many previous studies.

4.3. NCP

The annual or seasonal average NCP in the deep central AO basins are generally low compared to the adjacent shelf seas (Anderson et al., 2003; Codispoti et al., 2013; Sakshaug, 2004) or other open water regions, due primarily to insufficient availability of light and nutrients (Popova et al., 2010; Sakshaug, 2004; Vancoppenolle et al., 2013). This situation is unlikely to change much unless nutrients from 50 m or greater depths can be made available for phytoplankton growth (Codispoti et al., 2013). Even in a more open-water AO, where produc-
tion is expected to increase due to increased amount of area exposed to light, nutrients may be the more limiting factor for production (Tremblay and Gagnon, 2009). Low NCP and GPP values in the AO basin were found in many previous studies, though few estimates from the deep AO during late summer/early fall have been made so far with most studies focused on the productive marginal seas (e.g., Cross et al., 2012; Shadwick et al., 2011; Wassmann et al., 2006).

We have estimated 7.4 mmol m\(^{-2}\) d\(^{-1}\) of NCP in the partially ice-covered sea surface water (corresponding to ITP-64) and \(~0\) mmol m\(^{-2}\) d\(^{-1}\) under multiyear ice-covered sea surface water (corresponding to ITP-65). Olli et al. (2007) estimated primary production in the Amundsen basin (89–88° N) of the central AO during the peak of the productive season (August 2001) to be 4.2-12.5 mmol m\(^{-2}\) d\(^{-1}\) in a \(~15\%\) ice-free surface. In 2002, Bates et al. (2005) measured NCP during spring (5 May-15 June) and summer (15 July-25 August) in the Canada Basin to be \(<1.25\)-2.08 mmol m\(^{-2}\) d\(^{-1}\), with NCP in shelf regions being significantly higher (83-238 mmol m\(^{-2}\) d\(^{-1}\)). Hill et al. (2005) estimated 30 mmol m\(^{-2}\) d\(^{-1}\) productivity in the euphotic zone on the edge of the Canada Basin during summer 2002. Ulfsbo et al. (2014) reported low NCP values (<1 mol m\(^{-2}\) over 63 days) in the ice-covered deep AO basins during late summer of 2011 (5 August - 7 October), with a strong spatial variability, but they sampled significantly larger area than ours. Previous studies have reported productivity in the range of 3.75-10.25 mmol m\(^{-2}\) d\(^{-1}\) in the deep AO (Cota et al., 1996; Wheeler et al., 1996; Gosselin et al., 1997; Chen et al., 2002).

As stated earlier, insufficient availability of light and nutrients limits production in the AO. The nitrate concentration in the ITP-65 mixed layer under heavy ice cover was virtually zero, whereas in the ITP-64 mixed layer it was in a detectable amount (Fig. 4.2, 4.6). The nu-
trients in the ITP-64 mixed layer were potentially a result of previous mixing that occurred due to winds when the surface water was open to the atmosphere. According to the wind speed record, the wind was strong prior to when the nutrients were measured during the ITP deployment (Fig. 4.8). In addition, advective mixing and tidal mixing can also introduce nutrients into the photic zone (Codispoti et al., 2013). The vertical profile of DO and DIC in the mixed layer (Fig. 4.9 and 4.10) may provide information about mixing events. If biological activity dominates the DO and DIC variability in the mixed layer they would maintain a negative relationship. A positive relationship between DO and DIC suggests mixing that brought up both DO and DIC from the deep water, which seems to have happened weakly at ITP-64. Our NCP estimates show that even with open water conditions NCP remained very low in the AO basin (Fig. 4.4b), which is typical of oligotrophic oceans (see Polovina et al., 2008). Although the NCP is considerably low even in the partially open seawater (at ITP-64) it is significantly higher than that in the ice-covered water (at ITP-65) (Fig. 4.4b and 4.4e), suggesting open water sustains higher production compared to ice-covered water. The gross primary production (GPP), calculated as NCP plus respiration, was also estimated and they are 13 mmol m⁻² d⁻¹ at ITP-64 and ~0 mmol m⁻² d⁻¹ at ITP-65.

**Figure 4.9.** DIC (red dot) in the ITP-64 mixed layer. The date and time for the first sampling are included on top of the figure. Samplings were done at ~15 minute intervals. The average MLD at ITP-64 is 21.5 ± 2 m.
Figure 4.10. Depth-resolved time-series DO data. Top figure refers to ITP-64, bottom figure refers to ITP-65. The DO values are small in the mixed layer and larger at depths. The short-term variability in DO in the mixed layers presumably originated from the combination of large concentration gradients around the sensors and water movement driven by large-scale processes. At both ITPs, surface seawater was always supersaturated with respect to DO.

Since NCP in the Canada Basin is expected to be negligible except during summer due to significant ice-cover, we compared our estimates with other annual estimates in the region, though scarcity of data from the deep AO during late summer-autumn constrained our scope to make any direct comparison. When considered a 90-day open water period, the NCP is estimated to be 0.66 mol m$^{-2}$ yr$^{-1}$ at ITP-64 and ~0 mol m$^{-2}$ yr$^{-1}$ at ITP-65 and the GPP is 1.2 mol m$^{-2}$ yr$^{-1}$ and ~0 mol m$^{-2}$ yr$^{-1}$, respectively. Subba Rao and Platt (1984) and Legendre et al. (1992) estimated phytoplankton production in offshore (depth > 200 m) central Arctic waters north of 65°N to be 0.75 mol m$^{-2}$ yr$^{-1}$. Gosselin et al. (1997) reported an annual primary production of about 1.25 mol m$^{-2}$ yr$^{-1}$ in the central basin. Moran et al. (1997) found that the distribution of total primary production within the interior Arctic increases with latitude from ~ 1 mmol m$^{-2}$ d$^{-1}$ at ~ 82°N to ~ 5-28 mmol m$^{-2}$ d$^{-1}$ from 84-90°N. Bates et al. (2005) estimated the rate of NCP in the AO basin stations to be 0.18-0.54 mol m$^{-2}$ yr$^{-1}$, when a growing season of 120 day was assumed. Anderson et al. (2003) reported annual primary production in the
AO to be on an average less than 0.04 mol m\(^{-2}\) yr\(^{-1}\). All these estimates with our more recent estimates indicate that production in the Canada Basin is considerably low compared to shelf regions and other low-latitude oceans. Our estimates using ITP-64 and ITP-65 data also show that NCP in partially ice-covered water is significantly higher than that under multiyear sea ice, which corroborates earlier observations that leads of open water can enhance productivity (Gosselin et al. 1997; Sakshaug, 2004).

4.4. Gas Flux and NCP Comparison

The CO\(_2\) gas flux rate and NCP at ITP-64 are very similar, 7.8 mmol m\(^{-2}\) d\(^{-1}\) and 7.4 mmol m\(^{-2}\) d\(^{-1}\), suggesting that NCP counterbalanced gas exchange over the period of the study. This is probably why ITP-64 \(pCO_2\) virtually stopped increasing half way through the measurement period. After 09/26/12, when no PAR is available and nutrients are likely depleted (Fig 4.2) and the ice coverage is still not heavy (~20%) (Fig. 3.1e), this counterbalanced coupling between CO\(_2\) gas flux and NCP would, in theory, gradually wane as the upward trend in CO\(_2\) would resume and DO would head towards saturation due to absence of biological activity. The increasing ice coverage toward the end of the measurement period is expected to continue which would gradually stop all significant gas exchange and NCP, at which point ITP-64 CO\(_2\) and DO would be similar to ITP-65 \(pCO_2\) and DO.
Chapter 5

Summary

Biogeochemical data collected from two ITPs provided insights into the inorganic carbon cycle and oxygen variability in the region. Our \( p\text{CO}_2 \) data show similar levels of \( p\text{CO}_2 \) in the Canada Basin compared to many previous data. In the case of both ITPs, \( p\text{CO}_2 \) and DO were undersaturated and slightly supersaturated, respectively, suggesting that DO in the shallow Arctic mixed layer equilibrates with the atmosphere relatively rapidly. ITP-64 data collected under sparse ice and ITP-65 data collected under multiyear ice were evaluated separately and comparatively with the help of biogeochemical models to assess the variability of \( p\text{CO}_2 \) and DO in seawater. ITP-64 \( p\text{CO}_2 \) and DO variability for the most part was linked to gas exchange and biological activity, though some of the variability was linked to salinity which was related to horizontal advection. Temperature had little overall effect, as there was not much warming and brine rejection did not appear to be important even under little ice cover where ice formation would presumably be significant as water reached freezing point. Horizontal gradients dominated \( p\text{CO}_2 \) variability in the densely ice-covered region with small gas exchange and negligible NCP in the surface. One interesting feature this study has revealed is that \( p\text{CO}_2 \) variability under heavy ice and little ice can be strikingly similar even when the contributions from gas exchange and photosynthesis are very different. Our CO\textsubscript{2} gas flux and NCP estimates in comparison with the wide range of previous estimates show that gas flux and NCP estimates are highly variable in space and time over the AO. Previous studies reported increases in primary production in the AO resulting from decreased sea ice extent and a longer phytoplankton growing season, though envisioned that future increases in production will slow down as surface nutrient inventories become depleted (Pabi et al., 2008,
Arrigo et al., 2008). Our estimates compared to previous reports found medium production in the partially ice-covered AO with nutrients in the mixed layer that is typically found in the subsurface Arctic waters (Arrigo et al., 2008), potentially as a result of wind-driven shelf-break upwelling (Carmack and Chapman, 2003).

It is likely that the $p$CO$_2$ levels in the surface water of the Canada Basin will move toward atmospheric saturation if the ice-free period continues to increase. We calculated $\Omega_{\text{arag}}$ at present $p$CO$_2$ levels and at atmospheric saturation (assuming atmospheric $p$CO$_2$ at 400 $\mu$atm) using CO2SYS. Under current conditions, $\Omega_{\text{arag}}$ at ITP-64 ranges from 1.03 to 1.16 with a mean of 1.08 ± 0.03. Under equilibrium conditions, ITP-64 $\Omega_{\text{arag}}$ ranges from 0.84 to 0.88 with a mean of 0.86 ± 0.01. Clearly, with sustained open water condition, equilibrium of surface seawater with the atmosphere would bring $\Omega_{\text{arag}}$ below 1 (Fig. 5.0), favoring aragonite dissolution. Undersaturation of the surface waters of the Canada Basin with respect to aragonite has already been observed (Yamamoto-Kawai et al., 2009b). Since the preindustrial period, the increase in atmospheric CO$_2$, surface water warming, and melting of sea ice has lowered surface calcium carbonate saturation state in the Canada Basin by ~0.3, 0.1 and 0.4 units, respectively (Yamamoto-Kawai et al., 2011). Under such conditions, the calcium carbonate shells of many marine organisms that support the fundamental food web in the ocean dissolve in seawater, jeopardizing the entire food chain in the ocean as well as on land.
Figure 5.0 Aragonite saturation state ($\Omega_{\text{arag}}$) at ITP-64 is shown on the top figure (black) and at ITP-65 on the bottom figure (black). Red lines refer to $\Omega_{\text{arag}}$ when assumed that seawater is in equilibrium with the atmosphere with respect to $p\text{CO}_2$. Equilibrium between seawater and atmosphere brings $\Omega_{\text{arag}}$ below 1, favoring aragonite dissolution.

To our knowledge, no under ice and partially open water Arctic $p\text{CO}_2$ and $O_2$ data have been compared before, though an appropriate comparison between under ice and open water or nearly open water biogeochemical data sets from the AO is challenging due to heterogeneous distribution of sea ice cover. The dynamic nature of sea ice (e.g. drifting, melting, melt ponds) further contributes to this heterogeneity, making the biogeochemical environment even more complex. Therefore, more open water and under ice studies are needed to gain confidence in our understanding of processes and controls that regulate $p\text{CO}_2$ and $O_2$ variability in the AO.
Chapter 6

Conclusions

The major findings and implications of this study are:

(1) Gas exchange, NCP and horizontal gradients are the main sources of the CO$_2$ and O$_2$ variability in the partially ice-covered Arctic Ocean.

(2) $p$CO$_2$ and O$_2$ differed significantly in contributions from gas exchange and NCP under sparse ice and multiyear ice cover.

(3) The $p$CO$_2$ under heavy ice and little ice were strikingly similar even though contributions from gas exchange and photosynthesis were quite different.

(4) Horizontal gradients dominated the more densely ice-covered regions, with no significant NCP in the surface.

(5) Equilibrium of Arctic seawater CO$_2$ with the atmosphere would bring aragonite saturation state below 1.
References:


Apollonio, S. (1959), Hydrobiological measurements on IGY Drifting Station Bravo. Transactions, American Geophysical Union, 40,316-3 19.


Borges, A. V. (2005), Do we have enough pieces of the jigsaw to integrate CO₂ fluxes in the coastal ocean? Estuaries, 28(1), 3–27.


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