Associate Editor Initial Decision: Publish subject to minor revisions (Editor review) 2 (04 Jun 2015) by Dr. Silvio Pantoja 3 Comments to the Author: 4 4 June 2015 5 6 Thanks for submitting a revised version of ms bg-2014-406. There are a few issues that 7 need to be taken care of, related to comments expressed in my previous communication 8 (Line # refers to file "bg-2014-406-manuscript-version3.pdf"). 9 10 1. Abstract, Lines 7-9 Production per season. Use per day 11 AR: These values have been converted to production per day. 12 2. Abstract, Page 1, Line 18. Find a synonym for impact: "The impact of deglaciation in 13 14 Glacier Bay has been observed to seasonally impact" 15 AR: The second use of 'impact' has been changed to 'influence'. 16 17 3. Abstract, page 1, Line 23. Add comma: "dissolved inorganic carbon inorganic 18 macronutrients" 19 20 AR: A comma has been added in this spot. 21 22 4. Abstract, page 2, Line 11. Show your data of "substantial spatial and temporal 23 variability" to be able to conclude that "largely reflect glacial influences within the bay". 24 AR: We added "...estimates may reflect..." as to not definitively conclude it is all glacial 25 26 melt, per the assumptions listed in the Caveats section. 27 28 5. Introduction, Page 4, line 22 "Our goal for this study was to better understand carbon 29 cycling in Glacier Bay and how it is impacted by glacial runoff. Additionally, we wish to 30 fill in some gaps in how these processes may influence net community production within 31 a glaciated fjord ecosystem and better understand how continued glacial melt will impact productivity in Glacier Bay, as well as in similar glaciated fjord ecosystems worldwide." 32 33 \*Referee 3 " The justification of the work is (STILL) poorly presented ..." Editor: Is there a scientific question? 34 35 "Justification of the work" is still "poorly presented", as previously mentioned by Editor 36 and Referee 3. In addition, "better understand" (which is not a strong argument to sustain 37 a study) appears twice here. Please rewrite considering those comments (\*) 38 39 AR: 'better understand has been deleted from the justification. Justification text in the 40 Intro was changed to, "Our goal for this study was to estimate the current level of 41 seasonal NCP in Glacier Bay and evaluate how this, along with air-sea CO<sub>2</sub> flux, impact 42 the carbon dynamics in this glaciated fjord. Our findings also contribute to the limited 43 knowledge regarding carbon cycling in Glacier Bay and how it is impacted by glacial

runoff. Our estimates presented are the first to attempt to assess the impact of seasonal

freshwater may influence net community production within a glaciated fjord ecosystem

glacial melt on NCP in Glacier Bay. We wish to fill in some gaps in how glacial

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11 AR: Text was

8. "AR: We have added a "Caveats" section (Section 4.0) that discusses these aspects and how they impact our DIC and NCP values."

The reviewer mentioned "important limitations" that are not reflected neither in the abstract nor in the conclusion sections. I would suggest estimating numerically their impact on your overall conclusions. This has to be reflected in your conclusions.

AR: We have added some additional text that tries to associate numerical values with the error estimates within the Caveats section. However, there is no published data regarding these caveats from regions similar to Glacier Bay for us to accurately attempt to put errors estimates on all physical processes, such as glacial flour, since we do not have a range for magnitude of input of glacial flour or its composition. We have cited a relatively similar study from a Greenland fjord that states their estimates of the amount of NCP they believe to be from biology vs. glacial influences.

9. Conclusions: First two paragraphs do not belong here as written. Please summarize or delete. This is conclusion of your work, based on the scientific question that needs to be clearly expressed at the end of introduction (see #5 above).

AR: Paragraph one was deleted and paragraph two was shortened and edited.

I encourage reviewing those aspects. Thanks again for your interest in Biogeosciences.

1	Assessing Net Community Production in a Graciated Araska Fjord
2	Stacey C. Reisdorph <sup>1*</sup> and Jeremy T. Mathis <sup>1,2</sup>
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15	*Correspondence to: S.C. Reisdorph (screisdorph@alaska.edu)
16	
17	Abstract
18	The impact of deglaciation in Glacier Bay has been observed to seasonally <u>influence</u> the Stacey Reisdorph 6/14/15 8:49 AM  Deleted: impact
19	biogeochemistry of this marine system. The influence from surrounding glaciers,
20	particularly tidewater glaciers, has the potential to <u>effect</u> the efficiency and structure of Deleted: greatly impact
21	the marine food web within Glacier Bay. To assess the magnitude, spatial and temporal
22	variability of net community production in a glaciated fjord, we measured dissolved
23	inorganic carbon, inorganic macronutrients, dissolved oxygen and particulate organic

carbon between July 2011 and July 2012 in Glacier Bay, AK. High net community 1 2 production rates were observed across the bay (~54 to ~81 mmol C m<sup>-2</sup> d<sup>-1</sup>) between the summer and fall of 2011. However, between the fall and winter, as well as between the 3 winter and spring of 2012, air-sea fluxes of carbon dioxide and organic matter respiration 4 5 made net community production rates negative across most of the bay as inorganic 6 carbon and macronutrient concentrations returned to pre-bloom levels. The highest 7 organic carbon production occurred within the west arm between the summer and fall of 2011 with  $\sim 4.5 \times 10^5$  kg C d<sup>-1</sup>. Bay-wide, there was carbon production of  $\sim 9.2 \times 10^5$  g C d<sup>-1</sup> 8 9 between the summer and fall. Respiration and air-sea gas exchange were the dominant 10 drivers of carbon chemistry between the fall and winter of 2012. The substantial spatial

and temporal variability in our net community production estimates may reflect glacial

influences within the bay, as melt-water is depleted in macronutrients relative to marine

waters entering from the Gulf of Alaska in the middle and lower parts of the bay. Further

glacial retreat will likely lead to additional modifications in the carbon biogeochemistry

of Glacier Bay with unknown consequences for the local marine food web, which

includes many species of marine mammals.

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## 1 1.0 Introduction

2	Glacier Bay lies within the Gulf of Alaska (Gulf of Alaska) coastal ocean and is a
3	pristine glacially influenced fjord that is representative of many other estuarine systems
4	that border the Gulf of Alaska (Fig. 1). Glacier Bay is influenced by freshwater input,
5	primarily from many surrounding alpine and tidewater glaciers. The low-nutrient influx
6	of freshwater into Glacier Bay, which is highest (up to ~40% freshwater in surface water
7	during the summer; Reisdorph and Mathis, 2014) along the northern regions of the bay,
8	affects the nutrient loading and, thus, biological production and carbon dioxide (CO <sub>2</sub> )
9	fluxes within the bay. The southern region of the bay is less affected by this runoff due to
10	distance from the glacial influence and is more influenced by marine waters that
11	exchange through a narrow channel with a shallow entrance sill (~25 m).
12	Over the past ~250 years, Glacier Bay has experienced very rapid deglaciation,
13	which has likely impacted the biological structure of the bay. As the climate continues to
14	warm, additional changes to this ecosystem and marine population have the potential to
15	impact net community production (NCP) within the bay, with cascading effects through
16	the food web. To better understand the seasonal dynamics of the underlying
17	biogeochemistry in Glacier Bay, we used the seasonal drawdown of the inorganic
18	constituents of photosynthesis within the mixed layer to estimate regional mass flux of
19	carbon and rates of NCP along with air-sea flux rates of CO <sub>2</sub> . This approach has been
20	used in other high-latitude regions to assess ecosystem functionality (e.g. Mathis et al.,
21	2009; Cross et al, 2012; Mathis and Questel, 2013), including net community production
22	and carbon cycling.
23	Previous studies have shown there is wide-ranging variability in rates of primary

Previous studies have shown there is wide-ranging variability in rates of primary

1 production within other glaciated fjord systems, though NCP data within these 2 ecosystems are sparse. Fjords within the Central Patagonia region (48°S – 51°S) are 3 strongly influenced by glaciated terrain and freshwater runoff, similar to influences in 4 and around Glacier Bay. A study by Aracena et al. (2011) looked at water column 5 productivity in response to surface sediment export production in various Chilean 6 Patagonia fjords (41-56°S). They calculated primary production rates during the summer between ~35 mmol C m<sup>-2</sup> d<sup>-1</sup> in the more southern regions (52°S - 55°S) and ~488 C m<sup>-2</sup> d<sup>-1</sup> 7 <sup>1</sup> to the north (41°S - ~44°S). In Central Patagonia, Aracena et al. (2011) estimated 8 primary productivity at ~57 mmol C m<sup>-2</sup> d<sup>-1</sup> in the spring, a value comparable to some 9 10 seasonal estimates in Glacier Bay, and found primary production rates comparable to 11 those of Norwegian fjords (~9 to ~360 mmol C m<sup>-2</sup> d<sup>-1</sup>). 12 There have been a number of studies conducted within Glacier Bay, though 13 conclusions of several studies are contradictory. Many of these studies had a short 14 duration and limited coverage, missing much of the spatial, seasonal, and annual 15 variability (Hooge et al, 2003). This lack of data leads to a significant gap in 16 understanding of carbon cycling in Glacier Bay, as well as a lack of predictability of 17 responses to changes in this estuarine system as climate change progresses. To capture 18 some of the seasonal and spatial variability in the bay, we collected and analyzed 19 monthly samples over a two-year period. This sampling regime, along with the variety of 20 samples taken, has provided us with the most robust dataset collected in Glacier Bay and 21 allowed us to elucidate the dynamic nature of NCP in a glaciated fjord. Our goal for this

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study was to estimate the current level of seasonal NCP in Glacier Bay and evaluate how

this, along with air-sea CO2 flux, impact the carbon dynamics in this glaciated fjord. Our

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and how it is impacted by glacial runoff. Our estimates are the first attempt to assess the impact of seasonal glacial melt on NCP in Glacier Bay. We wish to fill in some gaps in how glacial freshwater may influence net community production within a glaciated fjord ecosystem and estimate how continued glacial melt may impact productivity in Glacier Bay.

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## 2.0 Background

Glacier Bay was once covered by one large icefield, the Glacier Bay Icefield, that has been rapidly retreating since the Industrial Revolution, scouring the bay and leaving behind many alpine and tidewater glaciers. Currently, the marine portion of Glacier Bay is roughly 100 km from the entrance sill to the end of the west arm, and reaches depths > 400 m and > 300 m in the east arm and west arm, respectively (Fig. 2).

Seasonal variation in factors such as light availability, turbulent or wind mixing and freshwater input, impact physical conditions that are vital to primary production, including stratification, photic depth, and nutrient availability. These drivers of NCP vary temporally and spatially within Glacier Bay. Glacial runoff, along with glacial stream input, impart freshwater into the marine system, especially along the arms of the bay. Peak runoff has been shown to occur during the fall, though there is fairly constant flow from June to September (Hill, 2009). Low-nutrient glacial runoff is prevalent, and while it aids in stratification, its low macronutrient concentrations dilute available nutrients in the northern regions nearest tidewater outflows. In the lower parts of the bay, glacial influence is lower and macronutrients are more abundant allowing higher levels of

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primary production during spring and summer. Glacier Bay maintains relatively elevated 1 2 phytoplankton concentrations throughout the year compared to levels observed in similar 3 Alaskan fjords (Hooge & Hooge, 2002). However, insufficient research has been done on 4 the biological system within Glacier Bay to understand why this occurs. 5 For this paper, we have calculated seasonal NCP and air-sea carbon flux for the 6 four regions within Glacier Bay in order to better understand ecosystem production in a glacially dominated environment, representative of much of the southern coastal AK 7 8 region. This study has greatly enhanced our understanding of how glacial melt and air-9 sea flux impacts DIC concentrations, and thus NCP, in estuaries, like Glacier Bay, which 10 are numerous along the Gulf of Alaska coast in Alaska, as well as other glaciated fjords 11 worldwide. 12 13 3.0 Methods

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Ten oceanographic cruises took place aboard the National Park Service's R/V Fog Lark between July 2011 and July 2012. Water column samples were collected at six depths (2, 10, 30, 50,100 m and near the bottom) at each station throughout the bay (Fig. 1) with a maximum depth within the west arm of ~430 m (Fig. 2). Sampling depths correspond with those currently being used by the Glacier Bay long-term monitoring program and determined by the USGS in the1990s. Each 'core' station (Fig. 1) was sampled during every oceanographic sampling cruise, while all 22 stations were sampled during the months of July and January. "Surface" water refers to water collected from a depth of 2 m unless otherwise stated. Seasonal data was calculated by averaging each measured parameter at each depth for all cruises during the respective seasons. The

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1	summer season consists of June, July and August, fall includes September and October;
2	winter is comprised of February and March cruises, and the spring season includes the
3	months of April and May. Data has been averaged regionally within each of the four
4	regions of the bay (lower bay, central bay, east arm, and west arm) (Fig. 1). Regional
5	boundaries were selected based on historical and ongoing research in Glacier Bay.
6	Bathymetry data (Fig. 2) was retrieved from the National Geophysical Data Center.
7	Conductivity, temperature and pressure were collected on downcasts with a
8	Seabird 19-plus CTD. Dissolved oxygen (oxygen) was sampled and processed first to  Stacey Reisdorph 6/14/15 8:54 AM  Deleted: DO
9	avoid compromising the samples by atmospheric gas exchange. Samples for oxygen  Stacey Reisdorph 6/14/15 8:54 AM  Deleted: DO
10	analysis were drawn into individual 115 ml Biological Oxygen Demand flasks and rinsed
11	with 4-5 volumes of sample, treated with 1 mL MnCl <sub>2</sub> and 1 mL NaI/NaOH, plugged,
12	and the neck filled with DI water to avoid atmospheric exchange. Dissolved oxygen was
13	sampled and analyzed using the Winkler titrations and the methods of Langdon (2010).
14	Samples were analyzed within 48 hours. Apparent oxygen utilization (AOU) was derived
15	from observed oxygen concentrations using Ocean Data View calculations in version  Stacey Reisdorph 6/14/15 8:54 AM  Deleted: DO
16	4.6.2 (Schlitzer, 2013).
17	DIC and total alkalinity (alkalinity) samples were drawn into 250 mL borosilicate  Stacey Reisdorph 6/14/15 8:57 AM  Deleted: TA
18	bottles. Samples were fixed with a saturated mercuric chloride solution (200 µl), the
19	bottles sealed, and stored until analysis at the Ocean Acidification Research Center at the
20	University of Alaska Fairbanks. High-quality DIC data was attained by using a highly
21	precise (0.02%; 0.4 µmoles kg <sup>-1</sup> ) VINDTA 3C-coulometer system. <u>Alkalinity was</u> Stacey Reisdorph 6/14/15 8:57 AM  Deleted: TA
22	determined by potentiometric titration with a precision of ~1 μmoles kg <sup>-1</sup> . Certified

reference material, prepared and distributed by Scripps Institute of Oceanography,

- 1 University of California, San Diego (Dr. Andrew Dickson's Laboratory), were run daily
- 2 before sample analysis to ensure accuracy of sample values. The VINDTA 3C provides
- 3 real-time corrections to DIC and <u>alkalinity</u> values according to in-situ temperature and
- 4 salinity.
- 5 <u>Dissolved macronutrient samples (nitrate, phosphate, silicate) were filtered</u>
- 6 through 0.8 μm Nuclepore filters using in-line polycarbonate filter holders into 25 ml
- 7 HDPE bottles and frozen (-20°C) until analysis at UAF. Samples were filtered to remove
- 8 any particles, such as glacial silt, that had the potential to clog equipment during analysis.
- 9 Samples were analyzed within several weeks of collection using an Alpkem Rapid Flow
- 10 Analyzer 300 and following the protocols of Mordy et al. (2010).
- 11 Particulate organic carbon (POC) samples were collected from Niskins into brown
- 12 1 L Nalgene bottles and stored for filtering within 2 days of collection. Samples were
- 13 collected at 2 m, 50 m and bottom depths. A known volume of samples was filtered
- 14 through muffled and preweighed 13 mm type A/E glass fiber filters using a vacuum
- pump. Muffling involved using tweezers to wrap filters in aluminum foil and heating
- 16 them at 450°F for ~6 hours in a muffling furnace in order to remove any residual organic
- 17 material. Filtered sampled were frozen for transport back to UAF where they were then
- dried and reweighed. Analyses were completed by OARC at UAF and were run using the
- methods outlined in Goñi et al. (2001).
- The partial pressure of  $CO_2$  ( $pCO_2$ ) was calculated using CO2SYS (version 2.0), a
- 21 program that employs thermodynamic models of Lewis and Wallace (1995) to calculate
- 22 marine carbonate system parameters. Seasonally averaged atmospheric pCO<sub>2</sub> values
- 23 (µatm) were used (388.4, 388.9, 393.4, 393.8 and 391.8 for summer 2011 through

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- 1 summer 2012, respectively and were averaged from the monthly averaged Mauna Loa
- 2 archive found at www.esrl.noaa.gov. For seawater pCO<sub>2</sub> calculations in CO2SYS we
- 3 used K<sub>1</sub> and K<sub>2</sub> constants from Mehrback et al., 1973 and refit by Dickson and Millero
- 4 (1987), KHSO<sub>2</sub> values from Dickson, the seawater pH scale, and [B]<sub>T</sub> value from
- 5 Uppström (1974).
- 6 CO<sub>2</sub> fluxes were calculated using seasonally averaged seawater temperature, wind
- 7 speed, and seawater and atmospheric  $pCO_2$  data using the equation,

Flux = 
$$L * (\Delta pCO_2) * k$$
 (Eq. 1)

- 9 where L is the solubility of CO<sub>2</sub> at a specified seawater temperature in mmol m<sup>-3</sup> atm<sup>-1</sup>
- and  $\Delta p CO_2$  represents the difference between seawater and atmospheric  $p CO_2$  in  $\mu$ atm. k
- 11 is the steady/short-term wind parameterization in cm hr<sup>-1</sup> at a specified wind speed and
- 12 follows the equation,

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$$k = 0.0283 * U^3 * (Sc/660)^{(-1/2)}$$
 (Eq. 2)

- where U is wind speed in m s<sup>-1</sup>, Sc is Schmidt number, or the kinematic velocity of the
- water divided by the molecular diffusivity of a gas in water, and was normalized to 660
- 16 cm hr<sup>-1</sup>, equivalent to the *Sc* for CO<sub>2</sub> in 20°C seawater (Wanninkhof and McGillis, 1999).
- 17 Wind speeds were cubed using the methods of Wanninkhof and McGillis (1999) in an
- 18 attempt to account for the retardation of gas transfer at low to moderate wind speeds by
- 19 surfactants and the bubble-enhanced gas transfer that occurs at higher wind speeds.
- 20 Seawater temperatures for flux calculations were taken from surface bottle CTD
- 21 data. Wind speeds were obtained from a Bartlett Cove, AK weather station (Station
- 22 BLTA2) located in Glacier Bay and maintained by the National Weather Service Alaska
- 23 Region.

1 NCP calculations were made using the seasonal drawdown of photosynthetic 2 reactant DIC within the mixed layer (upper 30 m) and were normalized to a salinity of 3 35. NCP production was calculated between each season from the summer of 2011 to the 4 summer of 2012 (i.e. the change in concentrations between each consecutive season) 5 according to the equation (Williams, 1993), 6  $NCP = DIC_{season2} - DIC_{season1}$ (Eq. 3)7 =  $\Delta$ DIC (moles C per unit volume area) 8 The influx of high-DIC waters (e.g., river discharge) can cause a dampening of the NCP 9 signal. This effect can be accounted for by normalizing DIC to a constant deep-water 10 reference salinity (S=35; Millero, 2008). Since this equation only reflects the effects of 11 DIC, freshwater influences on alkalinity were accounted for by correction of the seasonal 12 changes in alkalinity (Lee, 2001) using the equation, 13  $\Delta DIC_{Alk} = 0.5*(\Delta Alk + \Delta NO_3^-)$ (Eq. 4)14 and subtracting this value from the seasonal change in salinity-normalized DIC (nDIC), 15 thus providing an NCP in which the significant process influencing seasonal changes to 16 DIC concentrations is biological productivity (Bates et al., 2005; Mathis et al., 2009; 17 Cross et al., 2012). Error imparted in calculating parameters, including DIC analysis and averaging of nutrient concentrations within the mixed layer, are propagated through our 18 19 NCP estimates at  $\sim \pm 5\%$  of the final NCP calculation. Error propagated through each

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NCP estimate is listed with the NCP calculations in Table 1.

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4.0 Caveats

While seasonal water column DIC concentration changes can be a good
approximation to determine seasonal NCP, there are several estuarine processes that we
were unable to constrain that likely influenced our NCP estimates and act as additional
sources of uncertainty. Some other sources of uncertainty, such as the influence of glacial
flour, was reduced through averaging of spatial and regional parameters as stations were
reoccupied within ~30 days of one another.
Glacial flour can enhance DIC concentrations in seawater. Therefore, there is the
possibility that the inclusion of glacial flour may have increased our DIC concentrations

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with respect to DIC drawdown from primary production. In this case, our estimates may underestimate NCP. However, we were not able to quantify the amount of glacial flour deposited in Glacier Bay or analyze its composition for this study. In Glacier Bay, the influence of glacial flour is limited to the northern regions (i.e. east and west arms) that are directly influence by glacial outflow, many of which enter the bay along inlets and not the main arms of the bay, possibly reducing the impact of glacial flour at many stations, Unfortunately, there is insufficient data to quantitatively estimate the amount and makeup of glacial flour or what error it imparts into our NCP calculations, but we

Freshwater runoff that enters the bay via glacial streams flows over streambeds and can leach minerals and nutrients from bedrock, enhancing these concentrations in the surface waters of Glacier Bay. While stream water runoff in Glacier Bay was not analyzed for this study, studies of glacial runoff in southeast Alaska have shown allochthonous stream water dissolved organic carbon (DOC) to be negatively correlated with glacial coverage (Hood, et al., 2009). Examining watersheds along the Gulf of

assume for the sake of our analysis that it is relatively small.

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1	Alaska, Hood et al. (2009) also found that the most heavily glaciated watersheds were a
2	source of the oldest, most labile (66% bioavailable) dissolved organic matter (DOM) and
3	that increased input of glacial melt was associated with increased proportions of DOM
4	from microbial sources. As we were unable to chemically analyze glacial runoff in
5	Glacier Bay, our NCP calculations using only changes in DIC concentrations
6	underestimate NCP in the bay, though freshwater input is corrected to some degree by
7	salinity normalized DIC concentrations. The quantification of freshwater input into the
8	bay is also hindered by the lack of any active gauging stations within the bay (Hill et al.,
9	2009). Glacially-derived DOC has been shown to be highly bioavailable, though
10	inversely correlated with glacial coverage (Hood et al., 2009; Hood et al., 2015). While
11	the remineralization of highly labile DOC between station occupations could have added
12	DIC back into mixed layer and decreased the signal of seasonal drawdown, any
13	significant contribution of DIC from remineralization in the mixed layer seems unlikely
14	given the slow remineralization rates and the short time periods (~30 days) between
15	station occupations.
16	Additionally, while glacial freshwater input has been shown to have some impact
17	on NCP estimates in Greenland fjords, Meire et al. (2015) found biological processes to
18	be the main driver of carbon dynamics. In a study similar to ours in Glacier Bay, AK,
19	Meire and his team estimated air-sea CO <sub>2</sub> fluxes and NCP in the Godthåbsfjord system in
20	western Greenland, as well as the impact of freshwater on these estimates. They
21	identified highging processes as the most important driver of carbon dynamics

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accounting for 65 to 70% of the total CO<sub>2</sub> uptake by the fjord system (Meire et al., 2015).

1	Some literature suggests that internal waves may form within the lower bay in an	
2	area of station 02, known as Sitakaday Narrows. This is an area of constriction with	
3	accelerated currents that can produce hydraulic instabilities, potentially causing internal	
4	waves that may influence mixing at depth as well as at a distance from this region (Hooge	
5	& Hooge, 2002). These internal waves may affect nutrient replenishment to surface	
6	waters, as well as mixing of DIC across the mixed layer. This addition of high-DIC	
7	waters from depth may also lead to an underestimation of NCP. However, we cannot	
8	make an estimation of how this affects our NCP estimations, as there is debate about how	Jeremy Mathis 6/25/15 2:13 PM <b>Deleted:</b> currently still
9	often internal waves form in Glacier Bay.	Jeremy Mathis 6/25/15 2:12 PM  Deleted: the whether, and
10		Jeremy Mathis 6/25/15 2:12 PM  Deleted: ,
11	5.0 Results	
12	5.1 Spatial and seasonal salinity distributions	
13	Salinity distributions throughout the bay were generally the result of the influence	
14	of glacial runoff. During this summer season salinity ranged from 22.9 in surface waters	
15	at station 20 to 32.5 in the bottom waters of station 24 in Cross Sound. Isohalines were	
16	horizontal down to $\sim$ 50 m from the upper arms through the upper portion of the lower	
17	bay then became vertical in the lower bay, intersecting the surface just north of station 01	
18	(Fig. 3).	
19	Salinity was more constrained during the fall, with a full water column range	
20	between 25.3 in the surface waters at station 07 and 31.4 at depth (~130 m) at station 13.	
21	Similar to the previous summer, isohalines remained horizontal from the upper arms to	

the mid-lower bay near station 01 where they become vertical and intersected the surface.

Salinities in the lower bay near were between ~30 and 31, with the higher salinities at depth in Cross Sound.

During the winter salinity had a narrow range 29.6 and 31.6. The highest salinities were observed in the bottom waters at station 24, though salinity was similar at all depth at this station (~31.4). The lowest salinities (~30) were within the top 10 m of station 12 with similar surface salinities throughout both arms. In the spring, salinity continued to have a narrow range, with bay-wide salinities between ~28.9 at the surface of station 12 and 31.7 in the bottom water of station 24. Salinities below a depth of 50 m were relatively homogenous at ~31 (Fig. 3).

Returning to summer conditions in 2012, a strong salinity gradient was observed in the upper 50 m along the east and west arms. Salinities across the bay ranged from 24.1 in the surface waters of station 12 to 32.2, at depth at station 24. The lowest salinities were observed in the surface waters at the head of both arms, with this low salinity signal stretching south through the through the central bay. Stations within the lower bay had the highest salinities having salinities between ~31 and 32 at all depths.

## 5.2 Spatial and seasonal distributions of DIC and nitrate

DIC and nitrate are important inorganic components that are consumed during photosynthesis at various rates throughout the year in Glacier Bay. DIC concentrations during the summer of 2011 ranged from  $\sim$ 1400 to 2100  $\mu$ mol kg<sup>-1</sup>, with the lowest concentrations in the arms and upper-central bay. Nitrate concentrations throughout the water column ranged from  $\sim$ 2.5 to  $\sim$ 37  $\mu$ mol kg<sup>-1</sup>, with slightly less variability in the surface layer ( $\sim$ 2.5 and 24  $\mu$ mol kg<sup>-1</sup>). Surface nitrate concentrations were low, but

1	remained >5 μmol kg <sup>-1</sup> at all stations. While there was a large drawdown of nitrate,
2	particularly in spring and summer (as much as 20 µmol kg <sup>-1</sup> when compared to winter
3	concentrations), surface waters were not depleted at any of the observed stations.
4	In the fall of 2011, DIC and nitrate concentrations increased in the surface waters
5	with DIC ranging from ${\sim}1700~\mu mol~kg^{1}$ to 2040 $\mu mol~kg^{1},$ while below the surface
6	concentrations reached $\sim 2075~\mu mol~kg^{-1}$ . Water column nitrate concentrations were
7	between ${\sim}12~\mu\text{mol}~kg^{}$ and 32 $\mu\text{mol}~kg^{}$ with similar concentrations within surface
8	waters (11 $\mu$ mol kg <sup>-1</sup> to 30 $\mu$ mol kg <sup>-1</sup> ) and the lowest concentrations observed in the arms
9	DIC concentrations were much more constrained during the winter (~1920 $\mu mol\ kg^{\text{-}1}$ to
10	2075 $\mu mol~kg^{1})$ than during previous seasons. Nitrate concentrations ranged from ${\sim}12$
11	$\mu mol~kg^{-1}$ to 33 $\mu mol~kg^{-1}$ .
12	During the spring of 2012 DIC and nitrate had reduced concentrations in surface
13	waters across the bay. Surface DIC concentrations were between ${\sim}1750~\mu\text{mol}~kg^{}$ and
14	2025 $\mu mol~kg^{1},$ with water column concentrations reaching ~2075 $\mu mol~kg^{1}(Fig.~4).$
15	Nitrate concentrations ranged from $\sim 7~\mu mol~kg^{-1}$ to $\sim 31~\mu mol~kg^{-1}$ , with an observed
16	surface water maximum of $\sim 20~\mu mol~kg^{-1}$ . Further drawdown of DIC and nitrate in
17	surface waters was observed during the summer of 2012. However, concentrations did
18	not drop as low as was observed during the previous summer. DIC concentrations ranged
19	from ~1545 to 2066 $\mu$ mol kg <sup>-1</sup> . Nitrate concentrations varied from ~13 to 33 $\mu$ mol kg <sup>-1</sup> ,
20	with surface concentrations between ${\sim}17$ and 31 $\mu mol\ kg^{\text{-1}}.$ The stations with the lowest
21	DIC and nitrate concentrations were those within the east arm and west arm (Fig. 4).

**5.3** Rates and Masses of NCP

```
1
                The seasonal transition between the summer and fall of 2011 had the largest rates
 2
       of NCP observed during the year of study. Rates of NCP were positive in all regions of
       the bay and were highest within the east and west arms of the bay at 70.3 \pm 3.5 and 81.3
 3
       \pm 4.1 mmol C m<sup>-2</sup> d<sup>-1</sup>, respectively. A similar NCP rate of 68.9 \pm 3.4 mmol C m<sup>-2</sup> d<sup>-1</sup> was
 4
 5
       observed within the lower bay, while the central bay had the lowest rate between of 53.6
 6
       ± 2.7 mmol C m<sup>-2</sup> d<sup>-1</sup> (Table 1).
 7
                Calculated rates of NCP became negative between fall and winter, as well as from
 8
       winter to spring. Between fall and winter, the lower bay had a rate of -14.2 \pm 0.7 mmol C
       m^{-2} d^{-1} followed by the central bay at -11.5 \pm 0.6 mmol C m^{-2} d^{-1}. Rates of NCP were
 9
10
       negative in the east and west arms (-0.5 \pm 0.03 \text{ and } -1.3 \pm 0.1 \text{ mmol C m}^{-2} \text{ d}^{-1}),
11
       respectively. Between the winter and spring of 2012, rates of NCP remained negative
       within the east and west arms (-36.4 \pm 1.8 mmol C m<sup>-2</sup> d<sup>-1</sup> and -26.6 \pm 1.3 mmol C m<sup>-2</sup> d<sup>-1</sup>
12
       respectively), and to a lesser degree in central bay (-17.5 \pm 0.9 \text{ mmol C m}^{-2} \text{ d}^{-1}). Positive
13
14
       NCP rate was estimated for the lower bay of 17.6 \pm 0.9 mmol C m<sup>-2</sup> d<sup>-1</sup>. Between the
15
       spring and summer of 2012 NCP rates were positive across the bay, with the highest rate
       in lower bay (19.4 \pm 1.0 \text{ mmol C m}^{-2} \text{ d}^{-1}). The central bay and the east arm had rates of
16
17
       17.2 \pm 0.9 and 15.7 \pm 0.8 mmol C m<sup>-2</sup> d<sup>-1</sup>, respectively, while the west arm had a lower
18
       rate at 6.0 \pm 0.3 \text{ mmol C m}^{-2} \text{ d}^{-1}.
19
                The total mass (kg C d<sup>-1</sup>) of carbon produced from NCP was also estimated
20
       between each season (Table 1). Production occurred between the summer and fall of
       2011, with the greatest production in the lower bay (4.5 \times 10^5 \pm 1.3 \times 10^4 \text{ kg C d}^{-1}). The
21
       central bay had a large amount of production (2.2 \times 10^5 \pm 1.1 \times 10^4 \text{ kg C d}^{-1}), followed by
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the west and east arms  $(1.8 \times 10^5 \pm 8.8 \times 10^3)$  and  $(1.8 \times 10^4 \pm 3.8 \times 10^3)$  kg C d<sup>-1</sup> respectively).

1	Between the fall and winter the lower bay had carbon production of $-9.3 \times 10^4 \pm$
2	$4.6 \times 10^3$ kg C d <sup>-1</sup> , while the east arm had a lowest degree of production at $-5.2 \times 10^2 \pm 2.6$
3	kg C d <sup>-1</sup> . NCP masses in central bay and west arm were also negative ( $-4.7x10^4 \pm$
4	$2.3 \times 10^4$ and $-2.7 \times 10^3 \pm 1.4 \times 10^2$ kg C d <sup>-1</sup> , respectively). Between the winter and spring of
5	2012 masses in the east and west arms were estimated at $-3.9 \times 10^4 \pm 2.0 \times 10^3$ kg C d <sup>-1</sup> and -
6	$5.8 \times 10^4 \pm 2.9 \times 10^3$ kg C d <sup>-1</sup> , respectively while the central bay had a value of -7.1 x 10 <sup>4</sup> ±
7	$3.6 \times 10^3$ kg C d <sup>-1</sup> . The lower bay was the only region to have a positive NCP of $1.1 \times 10^5$ ±
8	$5.7 \times 10^3 \text{ kg C d}^{-1}$ .
9	Transitioning from the spring to summer the lower bay had the greatest
10	production $(1.3 \times 10^5 \pm 6.3 \times 10^3 \text{ kg C d}^{-1})$ , followed by the central bay $(7.0 \times 10^4 \pm 3.5 \times 10^3)$
11	kg C d <sup>-1</sup> ). The arms exhibited the lowest biomass production, with an NCP in the west
12	arm of $1.3 \times 10^4 \pm 6.5 \times 10^2$ kg C d <sup>-1</sup> and $1.7 \times 10^4 \pm 8.5 \times 10^2$ kg C d <sup>-1</sup> in the east arm.
13	
14	5.4 Spatial and seasonal distribution of POC
15	During the summer of 2011 surface POC concentrations were between ~12 and
16	$\sim$ 55 $\mu$ mol kg $^{-1}$ . Station 20 had the highest POC concentration at all sampled depths ( $\sim$ 46
17	$\mu mol~kg^{1}, \sim\!\!30,$ and $\sim\!42~\mu mol~kg^{1},$ surface to bottom), while the west arm had the
18	highest POC concentrations below the surface ( $\sim 33~\mu mol~kg^{-1}$ at 50 m and depth). The
19	west and east arms exhibited negative AOU ( $\sim$ -80 and $\sim$ -64 $\mu mol~kg^{\text{-1}},$ respectively).
20	Below the surface concentrations were similar (~9 μmol kg <sup>-1</sup> ), while surface waters had a
21	POC concentration of $\sim$ 28 $\mu$ mol kg <sup>-1</sup> . Lower bay had relatively lower POC concentrations
22	( $\sim$ 15 µmol kg <sup>-1</sup> at all depths).

POC concentrations decreased, especially within surface waters during the fall. A

maximum regional POC concentration (~13 µmol kg<sup>-1</sup>) was observed in surface waters of 1 2 the west arm. Below the surface layer POC concentrations were low, between  $\sim$ 5 and  $\sim$ 8 μmol kg<sup>-1</sup>. A maximum regional surface AOU (~82 μmol kg<sup>-1</sup>) was estimated for the 3 lower bay and a minimum (~2 µmol kg<sup>-1</sup>) in the surface waters of the central bay (Fig. 5). 4 5 In the winter of 2012 surface water POC concentrations were not found to exceed 20  $\mu$ mol kg<sup>-1</sup> and AOU across the bay were on the order of ~70  $\mu$ mol kg<sup>-1</sup>. Surface POC 6 concentrations ranged from ~2 to ~15 µmol kg<sup>-1</sup>, while POC concentrations at depth 7 varied between ~3 and 16 μmol kg<sup>-1</sup>. The regional maximum in POC was in the surface 8 9 waters in the west arm (~11 μmol kg<sup>-1</sup>). The east arm and lower bay both had maximum 10 POC concentrations in the bottom waters ( $\sim$ 14 and  $\sim$ 9 µmol kg<sup>-1</sup>, respectively). 11 POC concentration in the surface waters increased during the spring of 2012, 12 primarily within northern regions of the bay. The east arm had the greatest increase in 13 surface POC (~62 μmol kg<sup>-1</sup>) with concentrations decreasing in the surface water to the 14 south. The west arm and central bay had similar surface POC concentrations of ~35 μmol kg<sup>-1</sup>, and ~30 μmol kg<sup>-1</sup>, respectively. The lower bay had the lowest surface POC 15 concentrations with ~13 µmol kg<sup>-1</sup>, while having the highest rate of NCP and AOU (~93 16 17 μmol kg<sup>-1</sup>). The lower bay subsurface and deepwater AOU values were positive and POC 18 concentrations, ~9 µmol kg<sup>-1</sup> each, were the highest among the regions. 19 AOU values decreased in surface waters across the bay, while rates of NCP were 20 elevated within these waters during the summer of 2012. Surface POC concentrations were highest in the east arm (~50 µmol kg<sup>-1</sup>), while below the surface layer, POC 21 concentrations decreased, ranging from  $\sim$ 4.5 to  $\sim$ 7  $\mu$ mol kg<sup>-1</sup> at 50 m and  $\sim$ 5 to  $\sim$ 8  $\mu$ mol 22 kg-1 at depth. The west arm and central bay regions had surface POC concentrations of 23

2  $\sim 13 \mu mol kg^{-1}$ .. 3 5.5 Relationship between DIC and Oxygen Stacey Reisdorph 6/14/15 8:55 AM 4 Deleted: DO Stacey Reisdorph 6/14/15 8:55 AM 5 During the summer of 2011, oxygen concentrations ranged from ~190 to ~400 Deleted: DO 6 μmol kg<sup>-1</sup>. All samples below the surface layer, as well as surface samples within the 7 lower bay followed the Redfield ratio, with concentrations at depth between ~190 and 280 µmol kg<sup>-1</sup> (Fig. 6). Surface samples of stations within the arms and central bay had 8 Stacey Reisdorph 6/14/15 8:55 AM 9 high oxygen concentrations and low DIC. Surface oxygen was higher than that at depth, Deleted: DO Stacey Reisdorph 6/14/15 8:55 AM 10 ranging between ~230 and 400 μmol kg<sup>-1</sup>. However, in the lower bay DIC concentrations Deleted: DO Stacey Reisdorph 6/14/15 8:55 AM remained elevated (~2030 µmol kg<sup>-1</sup>) and oxygen concentrations were low (~240 µmol 11 Deleted: DO 12 kg<sup>-1</sup>). During the fall, surface samples within the arms and central bay continued to 13 deviate from Redfield. Surface oxygen concentrations ranged from ~210 to ~330 µmol Deleted: DO Stacey Reisdorph 6/14/15 8:55 AM 14 kg<sup>-1</sup> and corresponded with reduced surface DIC concentrations. At depth, oxygen Deleted: DO concentrations varied between ~200 and 280 µmol kg<sup>-1</sup> with C:O ratios close to Redfield. 15 16 All samples, at the surface and at depth, followed Redfield closely with surface Stacey Reisdorph 6/14/15 8:55 AM 17 waters having slightly higher oxygen and lower DIC concentrations than those at depth Deleted: DO Stacey Reisdorph 6/14/15 8:55 AM 18 during the winter of 2012. Surface water oxygen concentrations were between 250 and Deleted: DO 19 ~280 µmol kg<sup>-1</sup>, while deeper waters ranged from ~230 to 255 µmol kg<sup>-1</sup>. Stacey Reisdorph 6/14/15 8:55 AM 20 In the spring, DIC was drawn down and oxygen concentrations increased, having Deleted: DO Stacey Reisdorph 6/14/15 8:55 AM a range between ~270 and 410 µmol kg<sup>-1</sup>. Oxygen concentrations were amplified while 21 Deleted: DO

DIC was reduced at stations in the northern-most regions of both arms. These samples

deviated the most from Redfield, while the remaining samples adhered to the Redfield

~23 µmol kg<sup>-1</sup> and the lower bay exhibited the lowest surface POC concentration with

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1	ratio. Below the surface layer, oxygen concentration throughout the bay ranged from	
2	$\sim$ 250 to 280 $\mu$ mol kg <sup>-1</sup>	
3	During the summer of 2012, the surface waters within the two arms and central	
4	bay continued to diverge from Redfield. DIC concentrations within the more northern	
5	regions of the bay (east arm, west arm, and central bay) were increasingly drawn down,	
6	while <u>oxygen</u> concentrations remained elevated. Surface <u>oxygen</u> concentrations ranged	
7	from ~260 to ~410 μmol kg <sup>-1</sup> , with lower <u>oxygen</u> concentrations at depth, varying from	
8	200 - ~270 μmol kg <sup>-1</sup> .	
9		
10	5.6 Air-Sea gas flux	
11	During the summer of 2011 winds were relatively low, at ~1.6 m s <sup>-1</sup> , with surface	
12	waters of the central bay and the west arm were undersaturated with respect to	
13	atmospheric $CO_2$ with $pCO_2$ values of ~250 $\mu$ atms. The central bay and the west arm	
14	acted as minor sinks ( $\sim$ -0.3 $\pm$ 0.02 mmol C m <sup>-2</sup> d <sup>-1</sup> each). The lower bay and east arm had	
15	much higher seawater $pCO_2$ values of ~488 $\mu$ atms and ~463 $\mu$ atms and acted as sources	
16	for atmospheric $CO_2$ of $\sim 0.2 \pm 0.01$ mmol C m <sup>-2</sup> d <sup>-1</sup> for each region (Fig. 7).	
17	During the fall of 2011, winds increased slightly to ~2.0 m s <sup>-1</sup> and surface waters	
18	in all regions of the bay were oversaturated with respect to the atmospheric CO2. The	
19	lower bay experienced the highest $pCO_2$ at ~670 $\mu$ atms and acted as the largest source for	
20	atmospheric $CO_2$ with a flux of ~1.1 ± 0.06 mmol C m <sup>-2</sup> d <sup>-1</sup> . The central bay also had	
21	elevated $pCO_2$ with ~510 $\mu$ atms leading to outgassing of ~0.5 $\pm$ 0.03 mmol C m <sup>-2</sup> d <sup>-1</sup> . The	
22	east arm had a $pCO_2$ and flux values similar to that of the central bay ( $pCO_2 = \sim 514$	
23	$\mu$ atms; flux = $\sim$ 0.5 mmol $\pm$ 0.03 C m <sup>-2</sup> d <sup>-1</sup> ). Air-sea CO <sub>2</sub> flux in the west arm was $\sim$ 0.3 $\pm$	

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- 1 0.02 mmol C m<sup>-2</sup> d<sup>-1</sup>, similar to the east arm and central bay, but had a slightly lower
- 2  $pCO_2$  of ~482 µatms (Fig. 7).
- 3 Surface waters during the winter of 2012 were oversaturated in CO<sub>2</sub> with respect
- 4 to the atmosphere and all regions experienced outgassing, with average wind speeds of
- 5  $\sim$  2.1 m s<sup>-1</sup>. Regional pCO<sub>2</sub> values were more constrained, especially within the arms and
- 6 central bay, ranging from ~400 μatms in the west arm and central bay to ~432 μatms in
- 7 the east arm. Similar  $pCO_2$  values and seawater temperatures (~3.5°C), led the west arm
- 8 and central bay to experience comparable CO<sub>2</sub> fluxes of  $\sim 0.03 \pm 0.002$  and  $0.06 \pm 0.003$
- 9 mmol C m<sup>-2</sup> d<sup>-1</sup>. The east arm had a slightly higher surface temperature (~4.1°C) and flux,
- with  $\sim 0.18 \pm 0.01$  mmol C m<sup>-2</sup> d<sup>-1</sup>, while the lower bay had a slightly higher CO<sub>2</sub> flux of
- 11  $\sim 0.76 \pm 0.04 \text{ mmol C m}^{-2} \text{ d}^{-1}$ .
- 12 In the spring, seawater temperatures increased slightly to ~5°C across the bay
- 13 while salinity remained similar to winter values (~29 to 31). However, all regions except
- 14 for the lower bay transitioned to sinks for atmospheric CO<sub>2</sub>. pCO<sub>2</sub> in the lower bay
- 15 remained oversaturated with respect to  $CO_2$  at ~423 µatms and had a flux of ~0.11 ± 0.01
- 16 mmol C m<sup>-2</sup> d<sup>-1</sup>. Within the other three regions of the bay, surface water temperatures
- increased by just over 1°C. However, pCO<sub>2</sub> decreased in the surface waters and these
- 18 regions acted as sinks for atmospheric CO<sub>2</sub>. The east arm had the greatest decrease in
- 19 pCO<sub>2</sub>, dropping from ~432 μatms to ~167 μatms and exhibiting seasonal outgassing of ~
- $20 -0.87 \pm 0.04$  mmol C m<sup>-2</sup> d<sup>-1</sup>. The central bay and west arm regions were also seasonal
- sinks for CO<sub>2</sub>, taking up  $\sim -0.39 \pm 0.02$  mmol C m<sup>-2</sup> d<sup>-1</sup> in the central bay and  $\sim -0.60 \pm$
- 22 0.03 mmol C m<sup>-2</sup> d<sup>-1</sup> in the west arm.

1 During the summer of 2012 pCO<sub>2</sub> in the east arm increased to  $\sim$ 337 µatms with  $\sim$  -2  $0.13 \pm 0.01$  mmol C m<sup>-2</sup> d<sup>-1</sup> of ingassing. The central bay had a pCO<sub>2</sub> of ~200 µatms and a flux of  $\sim -0.44 \pm 0.02$  mmol C m<sup>-2</sup> d<sup>-1</sup>. The lower bay and west arm, acted as sources for 3 atmospheric CO<sub>2</sub>, having pCO<sub>2</sub> values of ~411 µatms and ~507 µatms, respectively, while 4 5 the lower bay experienced a near-neutral flux of  $\sim 0.04 \pm 0.002$  mmol C m<sup>-2</sup> d<sup>-1</sup>. The west arm was oversaturated with respect to atmospheric CO<sub>2</sub> with a pCO<sub>2</sub> of ~507 μatms and a 6 7 flux of  $\sim 0.26 \pm 0.01$  mmol C m<sup>-2</sup> d<sup>-1</sup>. 8 6.0 Discussion 9 10 6.1 Relationships of DIC, Nitrate, and Dissolved Oxygen During the summer of 2011 surface waters in the arms and upper-central bay 11

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**Deleted:** DIC, nitrate and DO are important indicators of biological production in a marine ecosystem. One way they can be used as biological production indicators is through Redfield ratios. Carbon and oxygen have a C:O Redfield ratio of 106:-170 (Anderson et al., 1994) and the carbon to nitrate Redfield ratio is 106:16.

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**Deleted:** variability in DIC, nitrate and dissolved oxygen concentrations within the surface waters were a result of primary production, dilution from glacial discharge, or a combination of both processes. S

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deviated from Redfield ratios for C:O and C:N (Figs. 6 and 8) Waters below this surface layer followed the Redfield ratios. Nitrate and phosphate concentrations in the surface waters were not observed to reach depletion during the summer, indicating that they were being continuously supplied to the surface layer and that phosphate (data not shown) was not limiting. Sustained nutrient concentrations and nutrient replenishment may be the result of physical interactions within the bay, including wind, tidal and internal wave mixing, and mixing across sills.

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Increases in oxygen and the reduction in macronutrient concentrations, including DIC, within the more northern arms of the bay was due to primary production coupled with the influence of glacier runoff and salinity-driven stratification limiting mixing and nutrient replenishment in the mixed layer. In the fall of 2011, DIC and nitrate concentrations increased while oxygen decreased in the surface waters as primary production slowed and wind mixing increased. Due to decreasing primary production

- 1 nutrient concentrations were similar within surface waters with the lowest concentrations
- 2 observed in the arms where glacial runoff was still impacting surface waters. Surface
- 3 water ratios for C:O and C:N deviated from the Redfield ratios, but less so than observed
- 4 during summer as primary production began to decrease during the fall (Figs. 6 and 8).
- 5 During the winter of 2012, increased wind mixing and the reduction of glacial input led
- 6 to deeper water column mixing, with much more constrained DIC and nitrate
- 7 concentrations. During the winter nitrate and DIC concentrations continued to increase,
- 8 with C:O and C:N Redfield ratios indicated a decrease in primary production and
- 9 increase in mixing (Figs. 6 and 8). While DIC and nitrate concentrations fell near the
- 10 Redfield ratio, they deviated slightly from Redfield at the highest nitrate concentrations
- 11 (Fig. 4). This may have been due to nitrification of ammonium by bacteria leading to an
- 12 increase the nitrate concentration. Another possibility is 'carbon overconsumption', the
- process in which more DIC is taken up than that inferred from the C:N Redfield ratio
- 14 (Voss et al., 2011). Explanations for carbon overconsumption include the preferential
- 15 remineralization of organic nitrogen (Thomas and Schneider, 1999) or an increased
- release of dissolved organic carbon (Engel, et al., 2002; Schartau et al., 2007).
- 17 As temperatures began to warm in the spring of 2012, the onset of glacial melt
- 18 and primary production reduced DIC and nitrate, while increasing oxygen concentrations
- 19 in surface waters across the bay. DIC and nitrate correlated closely with the Redfield
- 20 ratio except for two surface samples located at the northernmost ends of each arm (Fig.
- 21 8). This deviation may be explained by the fact that these stations were the first to be
- 22 influenced by glacial runoff during the onset of the glacial melt season.
- 23 Further reduction in DIC and nitrate concentrations in surface waters was

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1 observed during the summer of 2012 as primary production intensified, increasing

2 oxygen concentrations. Low nutrient glacial runoff was highest at this time of year,

3 affecting surface water DIC and nitrate concentrations within the arms. However,

4 concentrations did not drop as low as was observed during the previous summer.

5 Macronutrients did not reach depletion during the summer of 2012, implying they were

6 not the limiting primary productivity, possibly due to nutrient replenishment via tidal

7 pumping. Surface nitrate concentration continued to deviate from the C:N Redfield ratio

as these macronutrients were increasingly drawn down by primary productivity and

9 diluted by glacier runoff (Fig. 8). Surface waters in several regions also deviated from the

10 C:O Redfield ratio (Fig. 6) and those most affected were within the east arm and west

arm, as well as upper central bay, where freshwater influence was greatest. Mixing of

12 nutrient-rich marine waters from the Gulf of Alaska likely offset much of the drawdown

from primary production and allowed these surface waters within the lower bay to fall

14 closer to the Redfield ratio.

16 **6.2 NCP** 

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17 The seasonal transition between the summer and fall of 2011 had the largest rates

18 of NCP observed during the year of study. During this time all NCP rates were positive,

signifying enhanced primary productivity in the mixed layer. Rates of NCP became

negative during the seasonal transitions from fall to winter, as well as from winter to

spring. These negative NCP values indicate that air-sea fluxes (discussed in Section 5.6)

and organic matter respiration were prominent, increasing CO<sub>2</sub> (DIC) concentrations in

23 the surface waters and overwhelming any weaker signal from primary production.

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1	Between the fall and winter, the lower bay experienced the highest degree of $CO_2$ flux
2	when compared to biological production. The biological production was overwhelmed by
3	CO <sub>2</sub> influx in the east and west arms, but to a less degree than in regions to the south.
4	Between the winter and spring of 2012 the lower bay was the only region where
5	biological production dominated the CO <sub>2</sub> flux with a positive NCP rate, reflecting the
6	region's nutrient-rich marine influence from the Gulf of Alaska. The CO 2 flux signal
7	exceeded NCP within the east and west arms of the bay and, to a lesser extent, the central
8	bay. Transition from the spring to summer of 2012, primary production was evident in
9	the NCP rates. The west arm experienced a lower rate of NCP, possibly the result of the
10	strong low-macronutrient glacial influences along the arm, which may work to hinder
11	production. Additionally, large volumes of glacial flour imparted into the surface waters
12	from runoff during summer may have limited the photic depth and thus impeded some
13	productivity in the upper arms of the bay.
14	The total mass of carbon produced between seasons via NCP was also estimated
15	(Table 1). Between the summer and fall of 2011, we observed the greatest production of
16	organic carbon of any seasonal transition, with the largest production signal in the lower
17	bay and decreasing to the north as glacial influence increased. Elevated production
18	estimates within the lower could be due to continued nutrient replenishment to surface
19	waters as a result of mixing with the more marine waters outside of the bay.
20	Despite all regions of the bay being dominated by air-sea CO <sub>2</sub> flux during the fall
21	and winter seasons (Table 1) there was a substantial contrast in magnitudes of estimates  Deleted: between  Jeremy Mathis 6/25/15 2:22 PM  Deleted:

between the marine-dominated lower bay and the glacially-influenced east arm. These

differences in magnitude were likely the result of a higher degree of wind and tidal

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mixing at stations outside of and near the mouth of the bay, allowing this region to have elevated air-sea flux when compared to the east and west arms (Fig. 7).

The production signal within the arms and central regions of the bay continued to be overwhelmed by air-sea flux between the winter and spring of 2012 (Table 1). While production estimates remained negative in the northern regions of the bay, the lower bay had a positive NCP mass signifying increased primary production and a decrease in air-sea flux in this region. This increase in NCP in the lower bay may be been the result of earlier nutrient replenishment via the more marine waters outside of the bay. Between the spring and summer there was increased production across the bay as stratification strengthen and the hours of daylight increased, with the largest production estimates in the lower bay. The east and west arms exhibited the lowest biomass production, likely hindered by the inundation of low-nutrient glacial runoff that formed a fresh surface layer and imparted glacial flour into the surface waters in these regions.

#### 6.3 Air-Sea Flux

Aside from primary production, air-sea carbon dioxide (CO<sub>2</sub>) flux also impacts carbon concentrations within surface waters. In Glacier Bay, air-sea fluxes varied regionally and seasonally between the summer of 2011 and the summer of 2012. During the summer of 2011 winds were relatively low, reducing turbulent mixing, allowing for stratification and, thus, primary production. Surface waters in the lower bay and east arm acted at sources for atmospheric CO<sub>2</sub>, while the central bay and the west arm acted as sinks (Fig. 7). Drawdown of CO<sub>2</sub> in the west arm may be attributed to primary production, as well as the influx of low nutrient glacial melt. The central bay has been noted to have elevated production levels (Hooge and Hooge, 2002) that may account for

- 1 the drawdown of DIC and the region's sink status. Within the east arm seawater
- 2 temperatures were high, increasing the  $pCO_2$  of these waters and, combined with
- 3 influence of the reduced <u>alkalinity</u> concentrations, resulted in an oversaturation of CO<sub>2</sub> in
- 4 the seawater with respect to the atmosphere, overwhelming any effect from DIC
- 5 drawdown via primary production and making this region a source for atmospheric CO<sub>2</sub>.
- 6 Turbulent mixing across and outside the sill, as well as through Sitakaday Narrows, likely
- 7 reduced stratification and enhanced air-sea flux, causing this region to be a source for
- 8 atmospheric CO<sub>2</sub>.
- 9 In the fall of 2011, winds increased slightly and all surface waters across the bay
- 10 experienced oversaturation with respect to the atmospheric CO<sub>2</sub>, with the lower bay
- acting as the strongest regional source (Fig. 7). The high  $pCO_2$  values observed during
- 12 fall, despite strong DIC drawdown during summer, may be the result of a variety of
- interactions. Reduced glacial runoff during fall increased <u>alkalinity</u> concentrations
- 14 (Reisdorph and Mathis, 2014) and surface water temperatures declined allowing them to
- 15 hold more CO<sub>2</sub> while mixing brought DIC-rich waters from depth to the surface.
- 16 Increased winds also likely led to enhanced turbulent mixing across the bay.
- During the winter of 2012 surface waters across all regions of the bay continued
- 18 to experience outgassing (Fig. 7), though to a lesser degree than during fall. The lower
- 19 bay experienced the largest degree of outgassing, likely due to its more turbulent mixing
- 20 than other regions. Despite winter having the lowest seawater temperatures, wind mixing
- 21 peaked and likely allowed for CO<sub>2</sub>-rich waters from depth and the air to enter the surface
- 22 waters, increasing  $pCO_2$  in all regions of the bay.

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1	Several regions of Glacier Bay transitioned to sinks for atmospheric CO <sub>2</sub> during	
2	the spring of 2012 as primary production increased and winds slowed. The lower bay was	
3	the exception, remaining oversaturated with respect to CO <sub>2</sub> and continuing to act as a	
4	minor source for atmospheric CO <sub>2</sub> . In the more northern regions, surface waters	
5	experienced a slight increase in surface temperatures, but due to the onset of spring	
6	productivity DIC was drawn down in the surface waters, decreasing the $pCO_2$ and	
7	allowing them to become sinks for atmospheric CO <sub>2</sub> . The east arm experienced the	
8	largest decrease in $pCO_2$ and became the largest sink region within the bay, while the	
9	west arm and central bay underwent similar flux transitions as primary production	
10	increased, drawing down DIC in the surface waters. Within the arms, the onset of glacial	
11	melt may have aided in setting up stratification, also helping to lead to larger sink statuses	
12	within these regions.	
13	During the summer of 2012, waters in the northern regions became increasingly  Jeremy Mathis 6/25/15 2:24 PM  Deleted: becoming	
14	saturated with respect to atmospheric $CO_2$ . While, $pCO_2$ in the east arm did increase from	
15	spring values, perhaps due to a small increase in surface water temperatures and	
16	reductions in alkalinity from glacial runoff, it was still undersaturated with respect to  Deleted: ed  Jeremy Mathis 6/25/15 2:24 PM  Deleted: ed	
17	atmospheric $pCO_2$ . Atmospheric $CO_2$ uptake within the central bay strengthened slightly  Stacey Reisdorph 6/14/15 8:58 AM  Deleted: TA	
18	from spring as $pCO_2$ in this region decreased, likely due to high levels of primary	
19	production in this region, as well as high nutrient replenishment from tidal mixing	
20	between the waters of lower bay and the stratified waters within the central bay (Hooge	

& Hooge, 2002). Conversely, the lower bay remained a minimal source for atmospheric

 $\mathrm{CO}_2$ , while the west arm transitioned into source during the summer. The lower bay

experiences the highest degree of turbulent or tidal mixing across the sill, within Cross

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Sounds, and through Sitakaday Narrows, inhibiting stratification and primary production 1 2 and causing it act as a source for atmospheric CO<sub>2</sub> year-round. The difference in the 3 sink/source status of the east and west arms of the bay was likely the result of differences Stacey Reisdorph 6/14/15 8:58 AM 4 in glacial influences, with the west arm more influenced by low-alkalinity glacial runoff Deleted: TA 5 as it has the majority of the tidewater glaciers along its length. These glaciers caused a Stacey Reisdorph 6/14/15 8:58 AM 6 higher degree of <u>alkalinity</u> and DIC dilution than was observed within the west arm. Deleted: TA 7 8 7.0 Conclusions 9 Glacier Bay experiences a high degree of spatial and temporal throughout the 10 year. Environmental influences vary seasonally along a gradient from the glacially-11 influenced northern regions within the arms to the marine-influenced lower bay. This 12 imparts spatial differences in stratification and macronutrient availability that effect 13 biological processes and thus, rates of NCP, Despite Glacier Bay's limited exchange with Deleted: within each of the four pre-defined regions of the Glacier Bay 14 the marine waters of the Gulf of Alaska, it has been observed to support elevated primary 15 production through most of the year (Hooge & Hooge, 2002). However, rapid 16 deglaciation within Glacier Bay has imparted a high volume of fresh glacial runoff, a 17 portion of which has been from tidewater glaciers that melt directly into the bay, 18 affecting stratification, macronutrient concentrations and influencing air-sea CO<sub>2</sub> Stacey Reisdorph 6/29/15 8:35 AM

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exchange and net community production. For this study, we calculated rates of NCP and

air-sea CO2 exchange in each of the four regions of Glacier Bay in order to assess current

production levels in the bay and how these processes may impact the carbon dynamics.

southeastern Alaska fjords, despite playing an important role in the global carbon cycle.

To date, there are no NCP or air-sea flux estimates for Glacier Bay or similar

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1	Rates of NCP were positive across the bay between the summer and fall of 2011,	
2	as well as between the spring and summer of 2012 during peak times of primary	
3	production. NCP was highest during the transition between summer and fall of 2011,	
4	with regional NCP rates ranging from ~54 to ~80 mmol C m <sup>-2</sup> d <sup>-1</sup> . Rates during the	
5	summer of 2012 were lower, between $\sim\!6$ and $\sim\!20$ mmol C m <sup>-2</sup> d <sup>-1</sup> .	
6	Between the fall of 2011 and winter of 2012, as well as between the winter and	
7	spring of 2012, air-sea gas exchange overwhelmed any production signal across the bay,	
8	especially during the fall (Fig. 7; Table 1). The one exception was lower bay between	
9	winter and spring where NCP rates were positive, likely due to earlier replenishment of	
10	nutrients from marine waters outside the bay.	
11	The impact of rapid deglaciation in Glacier Bay can be observed in the seasonal	
12	impacts on the carbon cycling and NCP in this estuarine system. This study enhances the	
13	limited biogeochemical literature regarding Glacier Bay and includes one of the more	
14	robust datasets from Glacier Bay. We found the highest level of NCP to occur between	
15	the summer and fall seasons in 2011, with the greatest production within the glacially-	
16	<u>influenced arms of the bay.</u> The influence of <u>the surrounding glaciers</u> has the potential to	Stacey Reisdorph 6/23/15 9:59 AM
17	significantly impact the efficiency and makeup of the marine food web within Glacier	<b>Deleted:</b> , especially tidewater glaciers,
18	Bay in unknown ways with unknown consequences. <u>However, additional study</u> of the <u>se</u>	Stacey Reisdorph 6/23/15 9:59 AM
19	influences and their effects on the rate of NCP is needed to fully understand the impacts	Deleted: Better understanding  Stacey Reisdorph 6/24/15 9:35 AM
20	of future deglaciation.	Deleted: of  Jeremy Mathis 6/25/15 2:26 PM
		<b>Deleted:</b> can help identify possible these outcomes
21		

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Sharman, Natalie Monacci, Kristen Shake, Seth Danielson and the entire NPS staff in 2 Gustavus and Juneau, AK for their help in sample collection, logistics and editing. We 3 4 also want to thank the staff and visitors of Glacier Bay National Park and Preserve, as well as the community of Gustavus for their support and interest in this project. 5 6 7 8 9 References 10 Anderson, L.A., Sarmiento, J.L., 1994. Redfield ratios of remineralization determined by 11 nutrient data analysis. Global Biogeochem. Cycles 8, 65-80. 12 13 Aracena, C., Lange, C.B., Luis Iriarte, J., Rebolledo, L., Pantoja, S., 2011. Latitudinal 14 patterns of export production recorded in surface sediments of the Chilean 15 Patagonian fjords (41–55°S) as a response to water column productivity. Cont. Shelf 16 Res. 31, 340–355. doi:10.1016/j.csr.2010.08.008 17 18 Bates, N.R., Best, M.H.P., Hansell, D. A., 2005. Spatio-temporal distribution of dissolved 19 inorganic carbon and net community production in the Chukchi and Beaufort Seas. 20 Deep Sea Res. Part II Top. Stud. Oceanogr. 52, 3303-3323. 21 doi:10.1016/j.dsr2.2005.10.005 22 23 Cross, J.N., Mathis, J.T., Bates, N.R., 2012. Hydrographic controls on net community 24 production and total organic carbon distributions in the eastern Bering Sea. Deep 25 Sea Res. Part II Top. Stud. Oceanogr. 65-70, 98-109. 26 doi:10.1016/j.dsr2.2012.02.003 27 Dickson, A.G., 1990. Standard potential of the reaction:  $AgCl_{(s)} + \frac{1}{2}H_{2(g)} = Ag_{(s)} + HCl_{(aq)}$ 28 and the standard acidity constant of the ion HSO<sub>4</sub> in synthetic seawater from 29 273.15 to 318.15. The Journal of Chemical Thermodynamics, 22, 113–127. 30 31 doi:10.1016/0021-9614(90)90074-Z 32 33 Dickson, A.G., Millero, F.J., 1987. A comparison of the equilibrium constants for the 34 dissociation of carbonic acid in seawater media. Deep Sea Research, 34: 1733-35 1743. doi:10.1016/0198-0149(87)90021-5

G7224 to the University of Alaska Fairbanks. We would also like to thank Lewis

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## 1 Figures and Tables

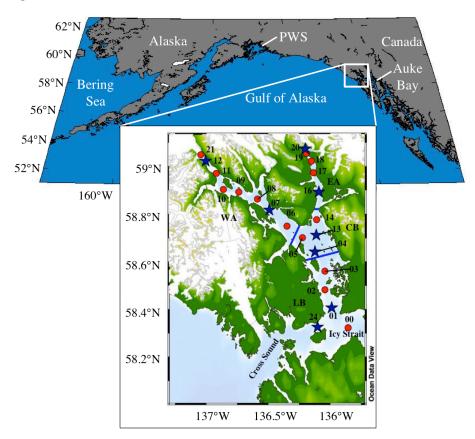
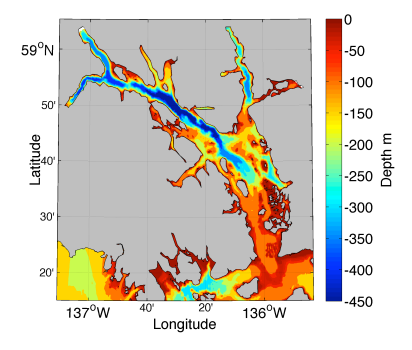
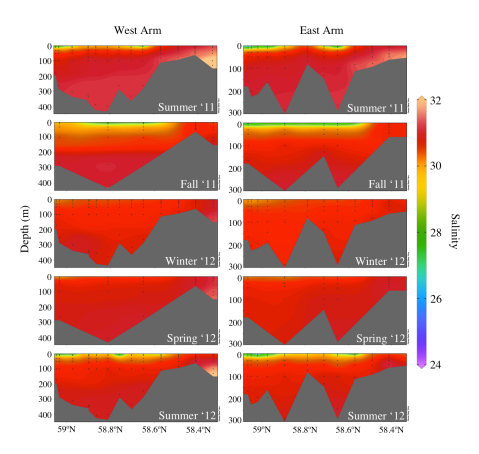


Fig. 1: Glacier Bay location and oceanographic sampling station map - Blue lines denote regional boundaries. Red dots show all oceanographic station locations with station number. Blue stars represent 'core' station location. lower bay, central bay, east, west arm.

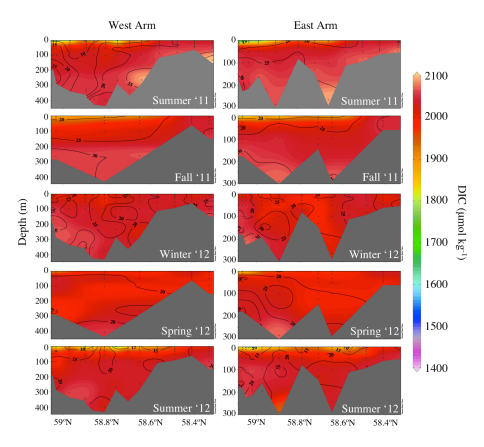


3 Figure 2: Bathymetry of Glacier Bay – Bathymetric map of Glacier Bay



2 Figure 3: Seasonal distribution of salinity. Spatial and seasonal distribution of salinity in

the water column.



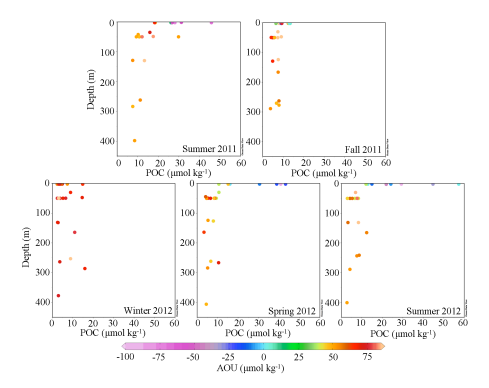
2 Figure 4: Spatial distribution of DIC and nitrate. Spatial and seasonal distribution of DIC

3 in the water column. Contours represent nitrate concentrations

Seasonal transition	Region	Regional Area	NCP rate	NCP mass
		$(m^2)$	$(\text{mmol C m}^{-2} d^{-1})$	(kg C d <sup>-1</sup> )
Summer and Fall	Lower Bay	5.44x10 <sup>8</sup>	$68.9 \pm 3.5$	$4.5 \times 10^5 \pm 2.3 \times 10^4$
	Central Bay	$3.40 \times 10^8$	$53.6 \pm 2.7$	$2.2 \times 10^5 \pm 1.1 \times 10^4$
	West Arm	$1.80 \times 10^{8}$	$81.3 \pm 4.1$	$1.8 \times 10^5 \pm 8.8 \times 10^3$
	East Arm	$9.00 \times 10^7$	$70.3 \pm 3.5$	$7.6x10^4 \pm 3.8x10^3$
Fall and Winter	Lower Bay	$5.44 \times 10^{8}$	$-14.2 \pm 0.7$	$-9.3x10^4 \pm 4.6x10^3$
	Central Bay	$3.40 \times 10^{8}$	$-11.5 \pm 0.6$	$-4.7x10^4 \pm 2.3x10^3$
	West Arm	$1.80 \times 10^{8}$	$-1.3 \pm 0.1$	$-2.7 \times 10^3 \pm 135.7$
	East Arm	$9.00 \times 10^7$	$-0.5 \pm 0.0$	-515.7 ± 25.8
Winter and Spring	Lower Bay	$5.44 \times 10^{8}$	$17.6 \pm 0.9$	$1.1 \times 10^5 \pm 5.7 \times 10^3$
	Central Bay	$3.40 \times 10^{8}$	$-17.5 \pm 0.9$	$-7.1 \times 10^4 \pm 3.6 \times 10^3$
	West Arm	$1.80 \times 10^{8}$	$-26.6 \pm 1.3$	$-5.7 \times 10^4 \pm 2.9 \times 10^3$
	East Arm	$9.00 \times 10^7$	$-36.4 \pm 1.8$	$-3.9 \times 10^4 \pm 2.0 \times 10^3$
Spring and Summer	Lower Bay	$5.44 \times 10^{8}$	$19.4 \pm 1.0$	$1.3 \times 10^5 \pm 6.3 \times 10^3$
	Central Bay	$3.40 \times 10^8$	$17.2 \pm 0.9$	$7.0 \times 10^4 \pm 3.5 \times 10^3$
	West Arm	$1.80 \times 10^8$	$6.0 \pm 0.3$	$1.3 \times 10^4 \pm 652.1$
	East Arm	$9.00 \times 10^7$	$15.7 \pm 0.8$	$1.7 \times 10^4 \pm 846.9$

<sup>2</sup> Table 1: Regional rates and masses of NCP – NCP by region in Glacier Bay based the

<sup>3</sup> change in salinity-normalized DIC concentrations between seasons.



3 Fig. 5: Seasonal POC vs. depth vs. AOU - Seasonal scatter plots of POC concentrations

- 4 vs. depth for each season between the summer of 2011 through the summer of 2012.
- 5 Color bar represents AOU in μmol kg<sup>-1</sup>.



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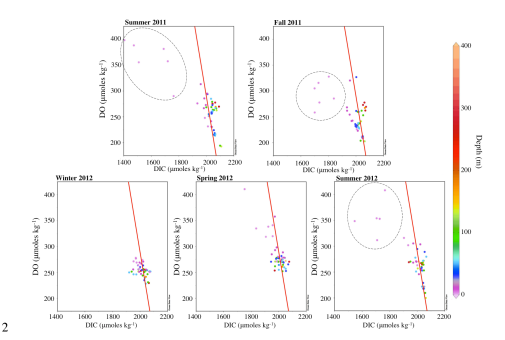


Fig. 6: Seasonal DIC vs. oxygen vs. depth - Scatter plots of DIC concentrations vs.

oxygen concentrations for each season between the summer of 2011 and the summer of

2012. Color bar represents depth in m. The red line depicts the C:O Redfield ratio of 106:

6 -170. Dotted circles highlight samples that deviate from Redfield.

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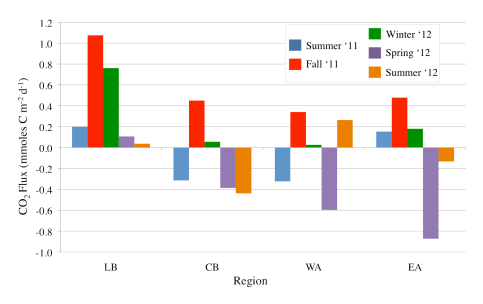
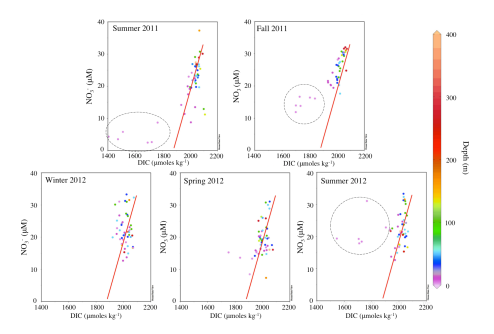


Fig. 7: Air-sea CO<sub>2</sub> flux – Seasonal air-sea CO<sub>2</sub> fluxes by region in mmol C m<sup>-2</sup> d<sup>-1</sup>. Blue represents the summer of 2011, red = fall of 2011, green = winter of 2012, purple =

4 spring of 2012, yellow = summer of 2012.

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2 Fig. 8: Seasonal DIC vs. NO<sub>3</sub> vs. depth - Scatter plots of DIC concentrations vs. NO<sub>3</sub>

- 3 concentrations for each season between the summer of 2011 and the summer of 2012.
- 4 Color bar represents depth in m. The red line depicts the C:N Redfield ratio of 106:16.
- 5 Dotted circles highlight samples that deviate from Redfield.