Net Community Production and Carbon Exchange From Winter to Summer in the Atlantic Water Inflow to the Arctic Ocean

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The eastern Fram Strait and area north of Svalbard, are influenced by the inflow of warm Atlantic water, which is high in nutrients and CO₂, influencing the carbon flux into the Arctic Ocean. However, these estimates are mainly based on summer data and there is still doubt on the size of the net ocean Arctic CO₂ sink. We use data on carbonate chemistry and nutrients from three cruises in 2014 in the CarbonBridge project (January, May, and August) and one in Fram Strait (August). We describe the seasonal variability and the major drivers explaining the inorganic carbon change (C_{DIC}) in the upper 50 m, such as photosynthesis (C_{BIO}), and air-sea CO₂ exchange (C_{EXCH}). Remotely sensed data describes the evolution of the bloom and net community production. The focus area encompasses the meltwater-influenced domain (MWD) along the ice edge, the Atlantic water inflow (AWD), and the West Spitsbergen shelf (SD). The C_{BIO} total was 2.2 mol C m⁻² in the MWD derived from the nitrate consumption between January and May. Between January and August, the C_{BIO} was 3.0 mol C m⁻² in the AWD, thus C_{BIO} between May and August was 0.8 mol C m⁻². The ocean in our study area mainly acted as a CO₂ sink throughout the period. The mean CO₂ sink varied between 0.1 and 2.1 mol C m⁻² in the AWD in August. By the end of August, the AWD acted as a CO₂ source of 0.7 mol C m⁻², attributed to vertical mixing of CO₂-rich waters and contribution from respiratory CO₂ as net community production declined. The oceanic CO₂ uptake (C_{EXCH}) from the atmosphere had an impact on C_{DIC} between 5 and 36%, which is of similar magnitude as the impact of the calcium carbonate (CaCO₃, C_{CALC}) dissolution of 6–18%. C_{CALC} was attributed to be caused by a combination of the sea-ice ikaite dissolution and dissolution of advected CaCO₃ shells from the south. Indications of denitrification were observed, associated with sea-ice meltwater and bottom shelf processes. C_{BIO} played a major role (48–89%) for the impact on C_{DIC}.

Keywords: Atlantic water, sea ice melt water, Fram Strait and Svalbard shelf, ocean CO₂ sink, denitrification, primary production, ocean acidification

INTRODUCTION

The Arctic Ocean is changing, where warming, decreasing sea-ice extent, thinning of ice and increased freshwater addition have been reported recently (e.g., IPCC, 2007; Rabe et al., 2009; Morrison et al., 2012; Pachauri et al., 2014). The characteristics of the Arctic ice cover has changed from thick multi-year ice to thinner first- or second-year sea ice (e.g., Serreze et al., 2007;...
Rabe et al., 2009; Granskog et al., 2016; Rösel et al., 2018). As a result of all these changes, surface-water stratification, primary production, carbon export and ocean CO₂ uptake are expected to change (e.g., ACIA, 2005; AMAP, 2013, 2018; Slagstad et al., 2015; Fransson et al., 2017).

Part of the changes within the Arctic Ocean originates from trends already observed in the Pacific and Atlantic inflow waters (e.g., Jones et al., 2003; Shadwick et al., 2011b). One of the main deep gateways to the Arctic Ocean is the Fram Strait, where the inflow (eastern Fram Strait) into the Arctic consists of the relatively warm and salty Atlantic water (AW; e.g., Schauer et al., 2008). Recent findings show warming of the inflowing Atlantic water into the Arctic Ocean (e.g., Schauer et al., 2008; Beszczynska-Möller et al., 2012). The warmer inflowing water also affects the Arctic Ocean, and waters around Svalbard and in the Barents Sea, where less sea ice in summer has been reported (e.g., Árthun et al., 2012; Onarheim et al., 2014; Carmack et al., 2015), which will have implications for the biogeochemical processes and greenhouse-gas exchange (Damm et al., 2011; Fransson et al., 2017). The inflowing Atlantic water supplies nutrients, which is favorable for primary production, with consequences for the marine ecosystem (e.g., Fransson et al., 2001; Torres-Valdés et al., 2013; Haug et al., 2017). Moreover, the Atlantic water also transports inorganic carbon into the Arctic Ocean (e.g., Anderson et al., 1998; Fransson et al., 2001). Most of the atmospheric CO₂ uptake occurs as the Atlantic water is cooled, during its way north along the Norwegian coast, and consequently the AW contains high anthropogenic CO₂ content (e.g., Sabine et al., 2004; Olsen et al., 2006; Vázquez-Rodríguez et al., 2009). In addition, Chierici (1998) found that although most of the CO₂ uptake occurred before the AW enters the Arctic Ocean, it was due to processes within the Arctic, such as transport by brine shelf plumes that acted to sequester CO₂ into deep waters.

Ocean CO₂ uptake, as an effect of increased atmospheric CO₂ due to the increased emission of CO₂ from human activities (e.g., burning of fossil fuel, deforestation), is causing ocean acidification (OA) in the Arctic Ocean (e.g., AMAP, 2013, 2018). In addition, reported release of methane and CO₂ from the Siberian shelves may also contribute to OA in the Arctic (e.g., Semiletov et al., 2012; Anderson et al., 2017; Qi et al., 2017). With continued warming, freshening and changes to primary production, the rate of ocean acidification is expected to increase (AMAP, 2013, 2018). This will have consequences for the carbon export, ocean CO₂ uptake, anticipated consequences for the marine organisms and ecosystems around the Svalbard Archipelago. For example, several studies suggest that increased CO₂ in the Arctic Ocean would increase and stimulate spring bloom production (Holding et al., 2015; Sanz-Martín et al., 2018).

The seasonal variability in physical variables (salinity, temperature, and water masses), nutrients and chlorophyll a concentrations is well known and thoroughly described from the CarbonBridge cruises in Randelhoff et al. (2018). In this study, we focus on the seasonal variability of the carbonate chemistry parameters from surface to 800 m, such as dissolved inorganic carbon (DIC), total alkalinity (AT), pH, fugacity of CO₂ (fCO₂), and aragonite saturation (Ω₄₇). Especially, little is known about the seasonal variability of these parameters as well as the ocean acidification state and change in the Atlantic water inflow to the Arctic Ocean. In addition, we use the semi-conservative tracer N*, which indicates deviations from a conservative nitrogen to phosphate behavior during photosynthesis and gives an indication of effects due to denitrification and nitrogen fixation in our study area. Here, we explore the components explaining the DIC change, such as biological DIC (e.g., CO₂) uptake (by the nitrate loss from pre-bloom values in January), the net DIC exchange with the surrounding environment, such as the air-sea CO₂ flux, and the formation and dissolution of calcium carbonate in three domains: the ice-melt affected domain, the Atlantic core-water domain, and the shelf domain. Our estimates are compared with other studies in the same area and polar regions in general.

STUDY AREA

The study area (79°N-80°N, 4°E-10°E) that includes the eastern Fram Strait, west and north of Svalbard and main surface currents are shown in Figure 1. The eastern Fram Strait and western shelf off Spitsbergen are affected by the warm and saline Atlantic water (AW), transported in the West Spitsbergen Current (WSC; e.g., Cottier et al., 2005). This current brings heat, nutrients, and carbon into the Arctic Ocean and the Svalbard area (e.g., Randelhoff et al., 2018; Renner et al., 2018). In the western part of the WSC, sea ice forms in winter and seasonal heating creates a surface layer of meltwater. In 2014, the area north of Svalbard was covered by sea ice throughout the study (Figures 2A-D). In our main study region between 79 and 79.5°N, and 4–10°E, the sea-ice boundary (>10% of sea ice) was found at about 4°E (Figure 2A). This limited the ship’s ability to move further west in January and May 2014. In the area close to Svalbard, shelf water and shelf processes dominated, resulting in a mixture of AW, coastal water and locally formed water such as transformed Atlantic water (e.g., Cottier et al., 2005; Randelhoff et al., 2018).

Three different domains are defined based on the surface-water characteristics in January and on topography. The meltwater domain (MWD) includes the western and northern parts of the study area that are influenced by fresher surface water in spring and summer due to sea-ice melt from the sea ice formed the previous winter. The Atlantic core-water domain (AWD) is dominated by the AW and the shelf domain (SD) is located on the shelf near West Spitsbergen (Figure 1).

MATERIALS AND METHODS

Sampling

The data of carbonate chemistry and inorganic nutrients were collected during four cruises in 2014 (Figure 1 and Supplementary Table S1). Three cruises were conducted as
FIGURE 1 | Map of the study area with station locations of the four different cruises. Shown are the major surface currents (Arctic in blue and the Atlantic water in red), and ice edge at 13th January 2014 (marked blue line).

part of the CarbonBridge project (January, May, and August). A Fram Strait cruise was conducted in late August 2014, to obtain information on the Atlantic water inflow. The data analysis concentrates on two sections with the largest seasonal coverage on the eastern shelf of the Fram Strait (January, May, beginning of August and end of August) between ∼79–79.5°N and 4–10°E.

Water samples were collected from 8-L Niskin bottles mounted on a General Oceanics 12-bottle rosette equipped with a Conductivity-Temperature-Depth sensor system (CTD, Seabird SBE-911 plus). Water samples were collected at a total of 11 to 14 depths, from surface to 800 m depth (or at bottom) at each station, with the highest resolution in the upper 100 m. From these samples inorganic nutrients, nitrate (NO$_3^-$), phosphate (PO$_4^{3-}$), silicic acid [Si(OH)$_4$], and total DIC and total AT were determined.

Section figures and surface interpolation from weighted-gridding were performed in Ocean Data View software version 4.7 (Schlitzer, 2015).

**Chemical Analyses**
The DIC and AT were analyzed after the cruises at the Institute of Marine Research (IMR Tromsø, Norway) following the method described in Dickson et al. (2007). DIC was determined using gas extraction of acidified samples followed by coulometric titration and photometric detection using a Versatile Instrument for the Determination of Titration carbonate (VINDTA 3D, Marianda, Germany). The AT was
FIGURE 2 | Sea ice cover around Svalbard for selected dates during the four cruises: (A) 13th of January, (B) 19th of May, (C) 8th of August, and (D) 25th of August in 2014. Data is obtained from the Ice Service of the Norwegian Meteorological Institute (MET, http://polarview.met.no/) and use their ice chart color scheme for very open drift ice (1–4/10ths), open drift ice (4–7/10ths), close drift ice (7–10/10ths), and fast ice (10/10ths).
was calibrated using reference seawater from Ocean Scientific IV analyzer from O.I. Analytical, United States. The analyzer standard methods (Grasshoff et al., 2009) using a Flow Solution concentration throughout the study, but it is actually the sum of Si(OH)_4 the two most common forms of aragonite (Ar_5). We used AT, DIC, and nutrient concentrations as input parameters in a CO_2-chemical speciation model (CO2SYS program, Pierrot et al., 2006) to calculate other variables describing the carbonate chemistry, such as pH, fugacity of CO_2 (f_{CO_2}), saturation state of calcium carbonate (Ω) for the two most common forms of aragonite (Ω_{Ar}) and calcite (Ω_{Ca}). The calculations are based on the carbonate system dissociation constants (K^1 and K^2) estimated by Mehrbach et al. (1973), modified by Dickson and Millero (1987) and the HSO_4^- dissociation constant from Dickson (1990).

Calcium carbonate (CaCO_3) saturation state (Ω) is commonly used to indicate a change in the CO_2 chemistry and the ocean acidification state, and indicates the dissolution potential for solid CaCO_3, such as calcareous shells and skeleton of marine organisms. When Ω < 1, solid CaCO_3 is chemically unstable and prone to dissolution (i.e., the waters are undersaturated with respect to the CaCO_3 mineral). In the Arctic Ocean, increased freshwater supply from sea-ice melt and river runoff have shown to decrease Ω (and provide a positive feedback on OA; Chierici and Fransson, 2009; Fransson et al., 2013, 2015). However, Ω is a chemical parameter showing the dissolution potential; most organisms require higher saturation state to grow due to the high energy demand of calcification. For example, the aragonite forming pteropod Limacina helicina, showed decreased calcification at Ω_{Ar} value of <1.4 and that lower values had negative effects on the shell density and thickness (e.g., Comeau et al., 2009; Lischka and Riebesell, 2012; Bednaršek et al., 2014).

Calculation of Seasonal Drivers of the Carbonate System

The strength of the effects and direction of different drivers of the change of DIC are schematically summarized in Figure 3. Earlier studies have shown that biological processes, such as photosynthesis and respiration explain much of the observed seasonal changes of the carbonate system in the Arctic Ocean as well as on the air-sea CO_2 exchange (Fransson et al., 2001, 2017; Chierici et al., 2011; Tynan et al., 2016). During photosynthesis, DIC and f_{CO_2} decrease, and pH and CaCO_3 saturation (Ω) increase. AT increases slightly during photosynthesis as a result of nitrate and hydrogen ion consumption when proteins are formed (Eq. 1a) but is less affected by photosynthesis than DIC (Figure 3). However, the AT change is twice as much as DIC during the production of calcium carbonate (Eq. 1b). This means that if CaCO_3 production occurs simultaneously with photosynthesis, both AT and DIC change, but if no CaCO_3 production takes place, only DIC changes. Thus, AT can give indications on the presence of CaCO_3-forming organisms and the contribution of CaCO_3 formation in sea ice. Sea-ice melting and formation cause changes to the carbonate chemistry where CaCO_3 is formed inside the ice (Assur, 1958) producing high CO_2-rich brine, which is rejected to underlying water (e.g., Rysgaard et al., 2007, 2013; Fransson et al., 2013, 2017). The solid CaCO_3 can be trapped in the ice until ice melting when it dissolves in the water (e.g., Rysgaard et al., 2012, 2013). The CO_2-rich brine is considered an important process for transport and sequestering of CO_2 in the Arctic Ocean to depth (e.g., Chierici, 1998; Fransson et al., 2013; Rysgaard et al., 2013). In our study area, sea ice is formed in the western part of the Fram Strait and in the area north of Svalbard while the AW inflow keeps the West Spitsbergen shelf ice free. Precipitation of CaCO_3 from the brine produces CO_2 (aq) and reduces AT...
in the brine (Eq. 1b), where Ca\(^{2+}\) and HCO\(_3^-\) denote the concentration of the calcium ions (Ca\(^{2+}\)) and the bicarbonate ions (HCO\(_3^-\)) that are consumed as solid CaCO\(_3\)(s), CO\(_2\), and water (H\(_2\)O) are produced.

\[
2\text{CO}_2 + \text{NO}_3^- + \text{H}^+ + \text{H}_2\text{O} \rightarrow 
\text{NHCH}_2\text{CO} (\text{org}) + 3.5\text{O}_2
\]  
(1a)

\[
\text{Ca}^{2+} + 2\text{HCO}_3^- \rightarrow \text{CaCO}_3(s) + \text{H}_2\text{O} + \text{CO}_2(aq)
\]  
(1b)

Physical processes such as mixing of sub-surface water, usually high in CO\(_2\) (low pH), play a large role for transferring CO\(_2\) to the surface water, especially in fall due to increased wind-induced mixing and water column cooling. The DIC, fCO\(_2\), and pH values also change with air-sea CO\(_2\) exchange (C\(_{\text{EXCH}}\)). When the ocean fCO\(_2\) is lower than the atmospheric, CO\(_2\) is added to the water CO\(_2\) (ocean sink, referred as invasion in Figure 3), and if ocean fCO\(_2\) is higher, it loses CO\(_2\) to the atmosphere during ocean outgassing (ocean CO\(_2\) source, referred to as release in Figure 3). Since AT describes the ion-charge balance in the water, changes in the uncharged CO\(_2\) will not affect AT (Figure 3). Seasonal warming and cooling affect the CO\(_2\) solubility and explain part of the fCO\(_2\) and pH variability.

The full inorganic carbonate system is used together with nutrient data from the seasonal cruises, to estimate the different components causing a change in DIC (C\(_{\text{DIC}}\)) from pre-bloom situation in January, to May, August to the end of August, described in Eqs 2–5. The January values of a salinity > 35 in the upper 50 m were considered representative for concentrations before the onset of photosynthesis (Supplementary Table S2).

In order to eliminate the change in concentrations due to salinity changes (i.e., dilution), all data were salinity-normalized to 35.1 (January salinity in the Atlantic water), after 35.1/S × C, where S refers to the observed salinity and C refers to the concentration of either DIC, AT, or NO\(_3\). Since the area is considered to be nitrogen limited (Smith et al., 1987; Kattner and Becker, 1991; Randelhoff et al., 2018), we used the change in salinity-normalized nitrate concentrations from January to May/August (ΔNO\(_3\), µmol kg\(^{-1}\)) converted to carbon using the carbon-to-nitrate (C:N) stoichiometric ratio based on 106:16 (Redfield et al., 1963), to estimate the biological component (C\(_{\text{BIO}}\), mol C m\(^{-2}\)) of the total DIC change (C\(_{\text{DIC}}\), mol C m\(^{-2}\)). Similar C:N ratios were found in the particulate organic matter during nitrogen consumption in May and August based on data collected in the same study (Paulsen et al., 2018). The difference between C\(_{\text{DIC}}\), C\(_{\text{BIO}}\), and C\(_{\text{CALC}}\) gives an estimate of the ocean’s role as an atmospheric CO\(_2\) sink or source, C\(_{\text{EXCH}}\) (mol C m\(^{-2}\)) according to Eq. 5. C\(_{\text{BIO}}\), C\(_{\text{EXCH}}\), and C\(_{\text{CALC}}\) were integrated in the top 50 meters. This assumption was valid because that was the maximum depth of nitrate drawdown observed in Randelhoff et al. (2018). Following from the discussion above, the change in total alkalinity (ΔAT, mol C m\(^{-2}\)) corrected for the effect of photosynthesis by subtracting the NO\(_3^-\) change, was used to estimate the change in the calcification component of the CO\(_2\) (mol C m\(^{-2}\)), referred to as C\(_{\text{CALC}}\), mol C m\(^{-2}\). The mean concentrations for each parameter in January at S > 35 were used to represent the pre-bloom state (Supplementary Table S2).

\[
\text{C}_{\text{DIC}} = \text{C}_{\text{BIO}} + \text{C}_{\text{EXCH}} + \text{C}_{\text{CALC}}
\]  
(2)

\[
\text{C}_{\text{BIO}} = \Delta\text{NO}_3 \times \text{C} : \text{N}
\]  
(3)

\[
\text{C}_{\text{CALC}} = 0.5(\Delta\text{AT} + \Delta\text{NO}_3)
\]  
(4)

\[
\text{C}_{\text{EXCH}} = \text{C}_{\text{DIC}} - \text{C}_{\text{BIO}} - \text{C}_{\text{CALC}}
\]  
(5)

**Calculation of N**

The semi-conservative tracer N\(^*\) (µmol L\(^{-1}\)) allows us to easily identify high and low anomalies relative to the global mean concentration of fixed nitrogen lost relative to the phosphate concentration (PO\(_4^{3-}\), µmol L\(^{-1}\), Eq. 6). Deviations from conservative behavior are meaningful in identifying regions of denitrification and nitrogen fixation but only give indications for either nitrogen loss or replenishment, which are not always caused by nitrification. This means that negative (positive) values of N\(^*\) cannot be directly associated with denitrification (nitrogen fixation). The distribution and seasonal change in N\(^*\) in our study area was calculated using the relationship described in Eq. 6 (Deutsch et al., 2001). This relationship includes processes leading to deviations in the entire water column and not only in the euphotic zone as described in Gruber and Sarmiento (1997).

\[
\text{N}^* = \text{NO}_3^- - 16 \times \text{PO}_4 + 2.9
\]  
(6)

In the equation, the constant 16 refers to the stoichiometric relationship between NO\(_3^-\) and PO\(_4^{3-}\) based on linear regression of world ocean nutrient data, and 2.9 is the constant derived by setting the global mean values to zero (Gruber and Sarmiento, 1997). The change in N\(^*\) reflects the net effect of denitrification and N\(_2\) fixation, thus negative N\(^*\) values suggests a deficit of nitrogen relative to the global mean, whereas positive values denotes larger than the world mean thus suggesting a nitrogen excess possibly linked to nitrogen fixation. In this study N\(^*\) is used to study the seasonal change of the deficit or source that may be explained by denitrification or nitrogen fixation, but also by advection of waters of different N\(^*\).

**Remotely Sensed Data**

To obtain information on the timing and development of the phytoplankton bloom in our study area we used data on chlorophyll (Chl\(_{\text{alt}}\)) and particulate inorganic carbon (PIC, Figure 12) from the Moderate Resolution Imaging Spectroradiometer (MODIS) onboard Aqua spacecraft downloaded from NASA Goddard Space Flight Center, Ocean Ecology Laboratory, Ocean Biology Processing Group. Level 3, 8-day binned, 9 × 9 km resolution arrays were further sampled into grid cells limited by 1° longitude and 0.5° latitude (Børsheim et al., 2014).
Estimates of net primary production from the Vertically Generalized Production Model (VGPM, Behrenfeld and Falkowski, 1997) were downloaded from www.science.oregonstate.edu. Annual primary production was estimated by integrating the production time series from each grid cell throughout the productive season (Borsheim et al., 2014).

The information on sea-ice coverage for the Svalbard area during part of our field periods displayed in Figure 2 was obtained from ice charts of the Ice Service of the Norwegian Meteorological Institute (MET) where the ice chart color scheme shows for very open drift ice (1–4/10ths), open drift ice (4–7/10ths), close drift ice (7–10/10ths), and fast ice (10/10ths).

RESULTS
Seasonal Variability in the Atlantic Water Inflow and Shelf Water Hydrography
Salinity and temperature in January, May, and August showed large variability between months (Figures 4–6). In January, there were clear longitudinal differences in the water-mass characteristics in the upper 100 m, which defined our study domains. At 4–5°E, there was a relatively thin surface layer in the upper 20 m, with lower salinity (<34.5) and temperature (<0°C) than in the surface water further to the east (Figures 4A,B). Between 5 and 8°E, the upper 100 m was more saline (>35) and warmer (4–6°C) than the other domains, while on the shelf (8–10°E) the salinity and temperature were intermediate, 34.8–35 and 2–4°C, respectively (Figures 4A,B). In January and May, a relatively warm (2–3°C) water off shelf, with high salinity (>35), reached from >600 m depth to the surface. The warm (>4°C) core in the upper 150 m in January was not observed in May (Figures 4B, 5B). The low-salinity (<34.5) and cold water (<2°C) in January in the upper 50 m observed in the western part of the study area (3–5°E), was located further to the east, closer to the slope (5–8°E) in May (Figures 4–6). In August, this low-salinity upper-50 m layer had spread eastward to the slope and shelf, where increased temperature (>6°C) was observed, compared to in January and May (Figures 6A,B). The relatively low salinity and relatively high temperature contributed to a water stratification in the upper 50 m.

Carbonate Chemistry and Ocean Acidification State
The AT, DIC, and fCO$_2$ values varied between the months. The AT values were lowest (about 2270 µmol kg$^{-1}$) in the upper 50 m at 4–5°E (MWD) in January (Figure 4C) and at 5–8°E (AWD) in May (Figure 5C), similar to the pattern observed in salinity (Figures 4A, 5A, 6A). The highest AT values of approximately 2325 µmol kg$^{-1}$ were observed in the locations of AWD and SD in the upper 50 m. In August, this low-alkalinity water was spread in the upper 50 m over the entire study area 4–10°E (Figure 6C), coinciding with low salinity (Figure 6A). The linear relationship between AT and salinity in the upper 50 m (AT = 57.07 × S + 315, R$^2$ = 0.95, N = 99), indicated a freshwater end-member of 315 µmol kg$^{-1}$. The DIC values in January and May were also low in the upper 50 m, coinciding with observations of low salinity and AT in the upper 50 m (Figures 4A,D, 5A,D). In August, the DIC trends were different than those observed in January and May, with the lowest DIC (<2100 µmol kg$^{-1}$), coinciding with both the lowest salinity and the highest temperature, in the upper 50 m (Figures 6A,B,D). At a few depths on the slope at 50–400 m depth, DIC was elevated compared to the same depth off the slope and on the shelf during all three cruises: in January (Figure 4D), May (Figure 5D) and August (Figure 6D). The fCO$_2$ values were undersaturated relative to the atmospheric fCO$_2$ levels of about 406 µatm (Fransson et al., 2017), with the lowest fCO$_2$ values in the upper 50 m in May (<200 µatm) and August (about 250 µatm), following a similar pattern to DIC (Figures 5D,F, 6D,F). In the water column on the shelf at about 9°E, the DIC and fCO$_2$ values were higher than the surrounding water. This was most evident in the fCO$_2$ that reached near atmospheric concentration of 406 µatm. The pH, Ω$_{Ar}$, and Ω$_{Ca}$ values also showed temporal variability, highest in May (Figures 5E,G,H) and August (Figures 6E,G,H) in the upper 50 m, coinciding with low DIC and fCO$_2$ values, in the stratified upper 50 m due to changes in salinity and temperature (Figures 5C,D, 6C,D). In May, the Ω$_{Ar}$ and Ω$_{Ca}$ values reached values up to a maximum of 2.5 and 4, respectively (Figures 5G,H), coinciding with low-salinity water (Figure 5A). The minimum Ω$_{Ar}$ and Ω$_{Ca}$ values of approximately 1.3 and 2.05, respectively, were observed at depths below 600 m in January (Figures 4G,H).

Nutrients, N$^*$, and Chlorophyll
In the AWD, the water column had high concentrations of NO$_3^-$, PO$_4^{3-}$, and Si(OH)$_4$ in January throughout the water column with values of 11, 0.8, and 5 µM, respectively (Figures 4I–K). Interestingly, the PO$_4^{3-}$ concentrations were higher near the ice edge (MWD) in the upper 200 m relative to the concentrations in AWD, whereas NO$_3^-$ and SiOH$_4$ concentrations were lower in this region, relative to those in the AWD. In the upper 50 m, nutrient concentrations changed from relatively high (values) in January to depleted or near-depleted values in May (Figures 4, 6I–K). The Si(OH)$_4$ concentration had the largest decrease from January to May in the AWD, whereas PO$_4^{3-}$ and NO$_3^-$ concentrations showed largest decrease in the low-salinity water. This was where high pH and high Ω$_{Ar}$ and Ω$_{Ca}$ values were observed (Figures 5E,G,H). By August, Si(OH)$_4$ concentrations in the upper 50 m, were low (<2 µM) from off shore to the shelf (Figure 6K). The ratio between ΔNO$_3$ and ΔPO$_4$ over the period of maximum observed nutrient decrease (i.e., January to May), based on data in upper 50 m was 15.2 ± 0.5 µM, similar to the N:P ratio of 16 reported by Redfield et al. (1963).

In January, N$^*$ values were generally positive, in excess of >3 µmol kg$^{-1}$ throughout the water column in the Atlantic-water influenced domain (Figure 4L). Similar values were also found on the shelf. Negative values of less than <−1 µmol kg$^{-1}$ were observed in the upper 100 m and below 650 m in the MWD (Figure 5L). By May, negative N$^*$ remained in the MWD (Figure 5L). The most striking difference from January to May was the change of N$^*$ in the western part of the
FIGURE 4 Variability of physical and chemical parameters in the eastern Fram Strait (latitude: 79–79.5°N, longitude: 4–10°E) in January 2014 from top: (A) salinity, (B) temperature (°C), (C) total alkalinity (AT, µmol kg⁻¹), (D) total dissolved inorganic carbon (DIC, µmol kg⁻¹), (E) pH, (F) fugacity of carbon dioxide (fCO₂, µatm), (G) aragonite saturation (ΩAr), (H) calcite saturation (ΩCa), (I) nitrate (NO₃, µmol L⁻¹), (J) phosphate (PO₄, µmol L⁻¹), (K) silicic acid (Si(OH)₄, µmol L⁻¹), and (L) N* (µmol kg⁻¹, negative values denote denitrification, and positive values show nitrification). The sampling locations and depths are denoted as black dots. The section plot and interpolation were performed by the Ocean Data View program (Schlitzer, 2015).
FIGURE 5 | Variability of physical and chemical parameters in the eastern Fram Strait (latitude: 79–79.5°N, longitude: 4–10°E) in May 2014 from top; (A) salinity, (B) temperature (°C), (C) total alkalinity (AT, µmol kg⁻¹), (D) total dissolved inorganic carbon (DIC, µmol kg⁻¹), (E) pH, (F) fugacity of carbon dioxide (fCO₂, µatm), (G) aragonite saturation (ΩAr), (H) calcite saturation (ΩCa), (I) nitrate (NO₃, µmol L⁻¹), (J) phosphate (PO₄, µmol L⁻¹), (K) silicic acid (SiOH₄, µmol L⁻¹), and (L) N⁺ (µmol kg⁻¹, negative values denote denitrification, and positive values show nitrification). The sampling locations and depths are denoted as black dots. The section plot and interpolation were performed by the Ocean Data View program (Schlitzer, 2015).
FIGURE 6 | Variability of physical and chemical parameters in the eastern Fram Strait (latitude: 79–79.5°N, longitude: 4–10°E) in August 2014 from top: (A) salinity, (B) temperature (°C), (C) total alkalinity (AT, µmol kg⁻¹), (D) total dissolved inorganic carbon (DIC, µmol kg⁻¹), (E) pH, (F) fugacity of carbon dioxide (fCO₂, µatm), (G) aragonite saturation (ΩAr), (H) calcite saturation (ΩCa), (I) nitrate (NO₃, µmol L⁻¹), (J) phosphate (PO₄, µmol L⁻¹), (K) silicic acid (SiOH₄, µmol L⁻¹), and (L) N* (µmol kg⁻¹, negative values denote denitrification, and positive values show nitrification). The sampling locations and depths are denoted as black dots. The section plot and interpolation were performed by the Ocean Data View program (Schlitzer, 2019).
AWD, which shifted from high positive $\mathrm{N^+}$ values in January, to negative values of less than $-2$ $\mu$mol kg$^{-1}$ at 7°E (Figure 5L). The change from positive to negative values were also observed on the slope (8°E), and on parts of the shelf at 9°E (Figure 5L). In August, the negative $\mathrm{N^+}$ values remained only in the intermediate water on the shelf, while the rest of the study area showed positive $\mathrm{N^+}$ values with a maximum of 2.8 $\mu$mol kg$^{-1}$ in the AWD (Figure 6L).

Remotely sensed chlorophyll data ($\mathrm{Chl}_{\text{sat}}$) gives an overview of the succession of the phytoplankton bloom and clearly shows the onset of a bloom near the ice edge (Figures 7B,C), white areas show ice cover. In the first week of May (Figure 7A), the bloom increased around the ice edge (Figure 7C), continuing until late June (Figure 7D), when the $\mathrm{Chl}_{\text{sat}}$ decreased. At the end of July, $\mathrm{Chl}_{\text{sat}}$ increased again in our study area and persisted through August (Figures 7F,G) until the beginning of September after which the bloom ceased completely and reached undetectable values (Figure 7H). Largest values were observed in between May and mid-June (Figures 7B,C).

### Seasonal Drivers of DIC Change

Negative values of $\mathrm{C_{BIO}}$ signify loss of DIC in the surface water (i.e., $\mathrm{CO_2}$) through net photosynthesis, whereas positive $\mathrm{C_{BIO}}$ values signify DIC release through net respiration. The $\mathrm{C_{BIO}}$ between January and May was between 1.9 and 2.2 mol C m$^{-2}$ (23–26 g C m$^{-2}$) in the MWD at about 4–6°E (Figure 8A). In the AWD, further east (> 6 and 8°E) and along 79°N, the $\mathrm{C_{BIO}}$ averaged between $-0.7$ mol C m$^{-2}$ and near 0 mol C m$^{-2}$ at the shelf break (Figure 8A). On the West Spitsbergen shelf, $\mathrm{C_{BIO}}$ was $-0.8$ mol C m$^{-2}$. The $\mathrm{C_{BIO}}$ between January and August was $-3.0$ mol C m$^{-2}$ ($-36$ g C m$^{-2}$) in the AWD (Figure 8B), implying a 0.8 mol C m$^{-2}$ net community production in summer (between May and August).

Negative $\mathrm{C_{EXCH}}$ indicates an ocean loss, for example through $\mathrm{CO_2}$ outgassing to the atmosphere (loss, negative values), while positive values denote an oceanic uptake of atmospheric $\mathrm{CO_2}$ (Figure 9, Eq. 4). Between January and May, $\mathrm{C_{EXCH}}$ ranged from being a net oceanic $\mathrm{CO_2}$ sink of about 0.5 mol C m$^{-2}$ on the SD, northern MWD and the northeast Svalbard, to become a net atmospheric $\mathrm{CO_2}$ source (loss) between 0.5 and 0.2 mol C m$^{-2}$ in the southern MWD (Figure 9A). By August, the southern part of the study area (around 79°N), especially evident in the AWD, $\mathrm{C_{EXCH}}$ changed from an insignificant ocean $\mathrm{CO_2}$ sink to a larger $\mathrm{CO_2}$ sink of about 2.3 mol C m$^{-2}$ (Figure 9B). In the northern part, $\mathrm{C_{EXCH}}$ values were changed from a net sink to a net $\mathrm{CO_2}$ source reaching a maximum release of 0.5 mol C m$^{-2}$. This was particularly pronounced in the area north of Svalbard (Figure 9B).

The influence of calcification or dissolution of $\mathrm{CaCO_3}$ on the DIC change ($\mathrm{C_{CALC}}$) was investigated following Eq. 4, where positive values signify a gain in DIC through $\mathrm{CaCO_3}$ dissolution and negative values a loss through $\mathrm{CaCO_3}$ formation/precipitation. Equation 4 is based on the salinity-normalized AT, corrected for photosynthesis using the nitrate change. Since AT is not affected by air-sea $\mathrm{CO_2}$ exchange, it is only $\mathrm{CaCO_3}$ dissolution that explains the increase in AT and ultimately the increase in $\mathrm{C_{CALC}}$. Generally, the area showed a DIC gain (positive $\mathrm{C_{CALC}}$) for the whole region, except for the small loss of 0.1 mol C m$^{-2}$ found in the SD between January and May (Figure 10A). At this time, the largest DIC gain of up to 0.5 mol C m$^{-2}$ was found in the MWD and the lowest DIC gain of less than 0.10 mol C m$^{-2}$ in the area north of Svalbard (Figure 10A). In the AWD, the largest $\mathrm{C_{CALC}}$ values were 0.3 mol C m$^{-2}$. Between January and August, the DIC gain derived from $\mathrm{C_{CALC}}$ had generally increased throughout the study area except in the MWD and north of Svalbard compared to the change between January and May. The largest DIC gain derived from $\mathrm{C_{CALC}}$ was observed in the AWD to a maximum of 0.7 C m$^{-2}$ between January and August, hence an increase between May and August of about 0.4 mol C m$^{-2}$ (Figure 10B).

Figure 11 shows a composite of the seasonal change of $\mathrm{C_{BIO}}$, $\mathrm{C_{EXCH}}$, and $\mathrm{C_{CALC}}$ focusing on variability of the area between 79–79.5°N and 4–10°E between January and May (Figure 11A), May and August (difference in the DIC between January and August, Figure 11B), and between beginning August and the end of August (Figure 11C). Figure 11 clearly shows a large biological DIC uptake, $\mathrm{C_{BIO}}$, in the MWD between January and May (Figure 11A). $\mathrm{C_{BIO}}$ DIC uptake increased (more negative $\mathrm{C_{BIO}}$) in the SD and AWD from May to August to a maximum $\mathrm{C_{BIO}}$ DIC change of $-3.0$ mol C m$^{-2}$ (Figure 11B). By the end of August, $\mathrm{C_{BIO}}$ showed a net DIC gain in the AWD and MWD of up to 2.3 mol C m$^{-2}$, sustaining biological DIC uptake (negative $\mathrm{C_{BIO}}$) in part of the MWD and SD of about 0.2 mol C m$^{-2}$ (Figure 11C). Between January and May, $\mathrm{C_{EXCH}}$ showed that the ocean generally acted as a small net oceanic sink of atmospheric $\mathrm{CO_2}$ of about 0.4 mol C m$^{-2}$ (Figure 11A). By beginning of August, the $\mathrm{CO_2}$ sink increased in the AWD of up to 2.3 mol C m$^{-2}$ (Figure 11B). At the end of August, the $\mathrm{CO_2}$ sink decreased greatly and changed $\mathrm{C_{EXCH}}$ by $-3.8$ mol C m$^{-2}$ from a sink to become a $\mathrm{CO_2}$ source, releasing $\mathrm{CO_2}$ to the atmosphere in the AWD (Figure 11C). In January to May and August, throughout the study area, the $\mathrm{C_{CALC}}$ resulted in a DIC gain of maximum of 0.5 mol C m$^{-2}$ in the MWD in January to May (Figures 11A,B). In the SD, $\mathrm{C_{CALC}}$ showed less importance than the other domains (Figures 11A–C). By the end of August, the $\mathrm{C_{CALC}}$ changed from a DIC gain to a DIC loss (Figure 11C) of a maximum of about $-0.5$ mol C m$^{-2}$.

The large $\mathrm{C_{CALC}}$ values in May and August were found in the area influenced by sea-ice formation and melt, such as in the MWD in May, and in the AWD in August. In our study area and time of year, the remotely sensed data on particulate inorganic carbon ($\mathrm{PIC}_{\text{sat}}$) showed a clear seasonal trend (based on data in the area 78.5–79.5°N and 4–10°E) from values less than 0.1 $\mu$mol kg$^{-1}$ in May to maximum PIC values of up to 0.6 $\mu$mol kg$^{-1}$ in August (Figure 12). The seasonal trend agrees with our $\mathrm{C_{CALC}}$ estimates for the AWD, but the values are too low to explain the $\mathrm{C_{CALC}}$ in the AWD of up to 0.5 mol C m$^{-2}$ between May and August (Figure 11B). Part of the difference between the $\mathrm{PIC}_{\text{sat}}$ and $\mathrm{C_{CALC}}$ are based on the methodological difference. The $\mathrm{C_{CALC}}$ values are integrated to 50 m and $\mathrm{PIC}_{\text{sat}}$ values are based on the surface ocean.

The succession of $\mathrm{C_{BIO}}$ from May to end of August was also observed from the remotely sensed chlorophyll data ($\mathrm{Chl}_{\text{sat}}$). The
FIGURE 7 | Remotely sensed chlorophyll a (Chlₐ, 8-day average) in May (A,B), June (C,D), July (E), August (F,G), and September (H), in 2014. Red values show the highest concentration and blue the lowest values on a logarithmic scale in mg m⁻³.
maximum 8-day mean values of chl\textsubscript{sat} in the area 79.75–78.75°N shown by longitude reached >0.5 mg m\textsuperscript{-3} (in the MWD) by 23rd April and reached the highest values of 6 mg m\textsuperscript{-3} in the beginning of July at 6°E (Figure 13A). After the peak, chl\textsubscript{sat} rapidly decreased to reach values below 1 mg m\textsuperscript{-3} (Figure 13A). The succession of the bloom was also investigated in the three domains, where chl\textsubscript{sat} > 0.5 mg m\textsuperscript{-3} was observed one week later in the AWD and SD domains than in MWD, on the 1st May (Figures 13B,C). In the AWD, the chl\textsubscript{sat} varied between 3 and 1.5 mg m\textsuperscript{-3} throughout the season and showed less seasonal variability than the other domains (Figure 13B). In the shelf domain at 9°E, generally higher chl\textsubscript{sat} of about 3 mg m\textsuperscript{-3} was observed than on the shallow shelf at 10°E, where chl\textsubscript{sat} values steadily increased and reached the highest values of 1.7 mg m\textsuperscript{-3} by the 23rd August. One week later, chl\textsubscript{sat} values rapidly declined to undetectable values (Figure 13C). The chl\textsubscript{sat} in the MWD
clearly showed a spring bloom between end of April to end of May, followed by a secondary bloom from mid-June to end of July (Figure 3A).

**DISCUSSION**

**Succession of the Bloom and Variability in Primary Productivity Estimates**

Our study agrees with several studies that show evidence of extensive spring blooms near the ice edge in the Arctic Ocean and its shelf seas. These blooms are mainly caused by relatively high-light conditions, meltwater-induced stratification and high nutrient availability, controlled by the nitrate concentration (i.e., Sakshaug, 2004; Wassmann and Reigstad, 2011; Assmy et al., 2017). Randelhoff et al. (2018) found an overall increase in ammonium values from May to August, explained by the remineralization after the spring bloom when the plankton community shifted toward a nutrient-recycling state during summer. Later in summer these authors suggested that thermal convection added nutrients and likely CO₂-rich sub-surface water to the surface ocean. Increased remineralization (where C\textsubscript{BIO} results in increased DIC) and mixing at the end of the summer would explain the rapidly changing conditions from a net biological DIC consumption and ocean CO₂ sink in August to a net biological DIC source in late August (Figure 11C).
FIGURE 10 | The interpolated and integrated $C_{\text{CALC}}$ in the top 50 meters (mol C m$^{-2}$) in 2014 calculated from the difference between January and: (A) May and, (B) August 2014. Positive values denote a gain in DIC through CaCO$_3$ dissolution and negative values a loss through formation of CaCO$_3$.

Increased mixing of CO$_2$-rich sub-surface water would also explain the change in $C_{\text{EXCH}}$ from an ocean CO$_2$ sink area in August to atmospheric CO$_2$ release by the end of August. Our study supports these findings: the spring bloom started near the ice edge in the west and moved eastward as the season progressed with most of the biological DIC consumption estimated in the AW domain and on the shelf (SD; Figure 11).

Based on the difference between January and May (2.2 mol C m$^{-2}$ and 26 g C m$^{-2}$) and between January and August (3.0 mol C m$^{-2}$ and 36 g C m$^{-2}$) estimates (Figure 8 and Table 1), about 75% of the $C_{\text{BIO}}$ occurred in spring, whereas the remaining 25% (0.8 mol C m$^{-2}$ and $\sim$9 g C m$^{-2}$) occurred in summer (between May and August). This spring value is similar to the annual export production estimates of 28–32 g C m$^{-2}$ in the Barents Sea presented by Fransson et al. (2001) using a similar approach as in our study. Our $C_{\text{BIO}}$ estimates are based on the loss of NO$_3^-$, sometimes referred to as new production, and do not consider production based on recycled nitrogen, other
FIGURE 11 | The integrated change of $C_{\text{BIO}}$ (green columns), $C_{\text{EXCH}}$ (orange columns), and $C_{\text{CALC}}$ (black columns) in top 50 meters ($y$-axis, mol C m$^{-2}$) between:
(A) January and May, (B) May to August, and (C) August to end of August along the 79°N section in eastern Fram Strait ($x$-axis Longitude°E). The dashed vertical lines divide the section in the three domains; meltwater domain (MWD), the Atlantic water domain (AWD), and the shelf domain (SD). Positive numbers denote a gain of carbon, and negative denote a loss of carbon.

than nitrate (Muggli and Smith, 1993). In the Labrador Sea, primary production showed a similar bloom succession as in our area, with high nitrate-based spring bloom in May and recycled production in August (Tremblay et al., 2006). They estimated the ratio between the relative contribution of NO$_3^-$ to total nitrogen uptake ratio, the $f$-ratio, to be 0.8 in May and 0.2 in August (Tremblay et al., 2006). The $f$-ratios from their study was used to convert our $C_{\text{BIO}}$ estimates to total production based on all nitrogen sources ($C_{\text{BIO} \text{TOT}}$). This resulted in a $C_{\text{BIO} \text{TOT}}$ of up to 2.6 mol C m$^{-2}$ for the spring bloom (between January and May), 1.4 mol C m$^{-2}$ in summer (between May and August), and a total annual $C_{\text{BIO} \text{TOT}}$ of 4.0 mol C m$^{-2}$. 
Based on chl\textsubscript{sat} data, the onset of the bloom occurred close to the 1st of May, resulting in a daily C\textsubscript{BIO} estimate of 0.10 and 0.14 mol C m\textsuperscript{-2} d\textsuperscript{-1}, or 1.2 to 1.7 g C m\textsuperscript{-2} d\textsuperscript{-1} using data collected between 16th and 23th of May (equivalent to 16 to 23 "bloom days," Figure 13). The daily total production estimate (C\textsubscript{BIOTOT}) for May range between 0.11 and 0.16 mol C m\textsuperscript{-2} d\textsuperscript{-1} (1.4–2.0 g C m\textsuperscript{-2} d\textsuperscript{-1}). Sanz-Martín et al. (2018) estimated total production (Gross Primary Production, GPP) west and north of Svalbard in May 2014. They used oxygen (O\textsubscript{2}) production in incubations with water collected from the spring-bloom conditions. Their study estimated a daily GPP of about 6.2 µmol O\textsubscript{2} l\textsuperscript{-1} d\textsuperscript{-1} (converted to 4.8 µmol C l\textsuperscript{-1} d\textsuperscript{-1} by C:O ratio of 106:138) from northwest Svalbard shelf stations (P1 and P5, Supplementary Table S1). Converting their value to the same bloom period as ours, integrated to 50 m results in a daily estimate of 0.24 mol C m\textsuperscript{-2} (2.9 g C m\textsuperscript{-2} d\textsuperscript{-1}) which is larger than our daily C\textsubscript{BIOTOT} estimates from 0.11 to 0.16 mol C m\textsuperscript{-2} d\textsuperscript{-1} b (2.6 mol C m\textsuperscript{-2} for 16–23 days). This is likely due to the integration of their value to 50 m, which probably overestimates the Sanz-Martín et al. (2018) GPP value. Wassmann et al. (2010) presented model estimates of GPP in the area west of Svalbard between 74 and 80°N, where they found a maximum annual GPP of 120 g C m\textsuperscript{-2} (10 mol C m\textsuperscript{-2}) for the full period between January and August was likely caused by methodological considerations. A difference in the integration depth can be expected, NCP\textsubscript{sat} was based on a surface chlorophyll a and PAR values, and variable euphotic zone depth, likely shallower than the 50-meter C\textsubscript{BIO} estimate. For example, Randelhoff et al. (2018) measured a mixed layer depth of 10–15 m at the ice edge.

Dissolved inorganic carbon uptake by Arctic productivity is similar to Antarctic estimates. In the Atlantic sector of the Southern Ocean in the Weddell Sea, Hoppema et al. (2007) estimated the NCP using integrated nutrient-deficiency methods such as in our study, and estimated NCP (same as our C\textsubscript{BIO}) to be 1.5 mol C m\textsuperscript{-2} for a 4-month period between November and March 2005 (Hoppema et al., 2002). Extrapolated values to

\begin{figure}
\centering
\includegraphics[width=\textwidth]{pic_sat.png}
\caption{The seasonal evolution of the remotely sensed particulate inorganic carbon (PIC\textsubscript{sat}, µmol kg\textsuperscript{-1}) from mid-April to end of August 2014, in the area between longitude 4 and 10°E, and latitude between 78.5 and 79.5°N.}
\end{figure}
FIGURE 13 | The seasonal variability of the maximum value of remotely sensed chlorophyll a (mg m$^{-3}$) in the area between 79.75 and 78.75°N from 4 to 10°E divided in the: (A) meltwater influenced domain (MWD), (B) the Atlantic water domain (AWD), and (C) the shelf domain (SD) from 4th of February to 12th of October in 2014. Values are Level 3, 8-day binned, 9 × 9 km resolution arrays from the Moderate Resolution Imaging Spectroradiometer (MODIS) onboard Aqua spacecraft downloaded from NASA Goddard Space Flight Center, Ocean Ecology Laboratory, Ocean Biology Processing Group.
TABLE 1 | Summary and direction of the estimated drivers of the mean and standard deviation for the seasonal DIC change (C$_{\text{DIC}}$, mol C m$^{-2}$) integrated in the top 50 meters in the three study domains: meltwater influenced domain, MWD, Atlantic water domain, AWD, and the shelf domain, SD, in May, August, and end of August 2014.

<table>
<thead>
<tr>
<th>Month</th>
<th>Domain</th>
<th>$C_{\text{BIO}}$ mol C m$^{-2}$</th>
<th>$C_{\text{EXCH}}$ mol C m$^{-2}$</th>
<th>$C_{\text{CALC}}$ mol C m$^{-2}$</th>
<th>$C_{\text{BIO}}$ % of abs.tot</th>
<th>$C_{\text{EXCH}}$ % of abs.tot</th>
<th>$C_{\text{CALC}}$ % of abs.tot</th>
</tr>
</thead>
<tbody>
<tr>
<td>May</td>
<td>MWD</td>
<td>$-2.2 \pm 0.3$</td>
<td>$0.1 \pm 0.4$</td>
<td>$0.3 \pm 0.5$</td>
<td>83</td>
<td>5</td>
<td>12</td>
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<tr>
<td></td>
<td>AWD</td>
<td>$-0.3 \pm 0.2$</td>
<td>$0.1 \pm 0.2$</td>
<td>$0.3 \pm 0.1$</td>
<td>48</td>
<td>8</td>
<td>44</td>
</tr>
<tr>
<td></td>
<td>SD</td>
<td>$-0.8 \pm 0.0$</td>
<td>$0.2 \pm 0.4$</td>
<td>$0.2 \pm 0.4$</td>
<td>70</td>
<td>15</td>
<td>15</td>
</tr>
<tr>
<td>Average</td>
<td></td>
<td>$-1.1 \pm 1.0$</td>
<td>$0.1 \pm 0.1$</td>
<td>$0.3 \pm 0.2$</td>
<td>67</td>
<td>9</td>
<td>24</td>
</tr>
<tr>
<td>August</td>
<td>MWD</td>
<td>$-2.3 \pm 0.9$</td>
<td>$0.4 \pm 0.1$</td>
<td>$0.2 \pm 0.2$</td>
<td>80</td>
<td>12</td>
<td>8</td>
</tr>
<tr>
<td></td>
<td>AWD</td>
<td>$-3.0 \pm 0.2$</td>
<td>$2.1 \pm 0.1$</td>
<td>$0.7 \pm 0.5$</td>
<td>53</td>
<td>36</td>
<td>11</td>
</tr>
<tr>
<td></td>
<td>SD</td>
<td>$-2.1 \pm 0.8$</td>
<td>$0.4 \pm 0.5$</td>
<td>$0.2 \pm 0.1$</td>
<td>78</td>
<td>13</td>
<td>9</td>
</tr>
<tr>
<td>Average</td>
<td></td>
<td>$-2.5 \pm 0.6$</td>
<td>$1.0 \pm 0.2$</td>
<td>$0.3 \pm 0.3$</td>
<td>70</td>
<td>20</td>
<td>9</td>
</tr>
<tr>
<td>Aug end</td>
<td>MWD</td>
<td>$-1.7 \pm 1.7$</td>
<td>$0.1 \pm 1.1$</td>
<td>$0.1 \pm 0.1$</td>
<td>89</td>
<td>5</td>
<td>6</td>
</tr>
<tr>
<td></td>
<td>AWD</td>
<td>$-2.2 \pm 0.1$</td>
<td>$-0.7 \pm 0.8$</td>
<td>$0.2 \pm 0.2$</td>
<td>70</td>
<td>24</td>
<td>6</td>
</tr>
<tr>
<td></td>
<td>SD</td>
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</tr>
</tbody>
</table>

Negative values denote a loss of DIC and positive values denotes a gain in DIC caused by the following processes: nitrate based biological carbon uptake ($C_{\text{BIO}}$, carbon lost from the water), air-sea CO$_2$ exchange, $C_{\text{EXCH}}$, where positive values denotes an ocean CO$_2$ sink and negative an ocean CO$_2$ source, and the net effect of calcium carbonate dissolution (gain, positive) or formation (loss, negative), $C_{\text{CALC}}$. All values in mol C m$^{-2}$ total over the specified period. The role of each driver relative to the total absolute change is indicated in as percentage (%) in the three right columns. nd indicates no data and na not applicable.

FIGURE 14 | Remotely sensed net primary production (NCP$_{\text{sat}}$, mol C m$^{-2}$) for 2014 from the Vertically Generalized Production Model (VGPM, Behrenfeld and Falkowski, 1997) downloaded from www.science.oregonstate.edu. NCP$_{\text{sat}}$ was estimated by integrating the production time series from each grid cell throughout the productive season (Børsheim et al., 2014).
productivity is constrained to the spring bloom. Areas west of Svalbard present twice as much annual production than in the Arctic, north of Svalbard. Finally, the depth of integration and the length of the sampling period can bias estimates in regions with shallow, and variable, mixed layer depth. Our analysis, encompassing from winter to autumn, highlights the importance of the seasonal signal in this high-latitude region.

The Oceanic Sink of Atmospheric CO₂ in Fram Strait Area and Its Variability

Several studies estimating the air-sea CO₂ fluxes and the annual net CO₂ sink show that the Arctic Ocean, the Fram Strait and waters around Svalbard act as atmospheric CO₂ sinks (e.g., Tynan et al., 2016; Yasunaka et al., 2016, 2018; Fransson et al., 2017). The 18-year annual average of observed seawater fCO₂ and a self-organizing mapping technique showed that the Arctic Ocean acted as an ocean CO₂ sink between 8 and 12 mmol m⁻² d⁻¹ and a daily mean in a full annual cycle of 5 mmol m⁻² d⁻¹ (Yasunaka et al., 2018). In our estimates, the study area showed a small oceanic CO₂ sink until May at an average of 0.12 ± 0.06 mol C m⁻² for the period January to May. By August, the ocean CO₂ uptake increased to a large net ocean CO₂ sink of an average of 1.1 ± 1.0 mol C m⁻² and a maximum CO₂ sink of 2.1 mol C m⁻² (25 g C m⁻²) in the AWD for the January to August period (Table 1). This suggests a mean daily ocean CO₂ uptake of 10 mmol m⁻² d⁻¹ and a maximum of 21 mmol m⁻² d⁻¹ in August (100 days). By the end of August, the AWD shifted to a significant CO₂ source of about 0.7 mol C m⁻² (Table 1). Accounting for the study period between January and August, this results in an ocean net annual CO₂ sink between 4 and 8 mmol m⁻² d⁻¹, which are similar to the results by Yasunaka et al. (2018). The winter-to-summer ocean CO₂ uptake estimates of 3.7 mol C m⁻² (44 g C m⁻²) by Fransson et al. (2001) in the Barents Sea in the upper 50 m, are nearly twice as large as our maximum estimate of 2.1 mol C m⁻² (25 g C m⁻²). Their relatively large CO₂ uptake were mainly driven by biological DIC uptake in the biologically productive waters of the Barents Sea. Model estimates for the Barents Sea showed an annual oceanic CO₂ uptake between 10 and 40 g C m⁻², related to warm and cold years (Slagstad and Wassmann, 1996), which were similar to our estimates of the annual net oceanic CO₂ uptake between 10 and 25 g C m⁻² for the whole period from January to May and August. In the Greenland Sea, the ocean acted as a CO₂ sink throughout the year of about 53 g C m⁻² (Anderson et al., 2000). On the other hand, our estimate is much larger than the Atlantic water influenced Kara-Laptev Sea of 1 g C m⁻² (Fransson et al., 2001). In the southern Beaufort Sea, surface waters were undersaturated with respect to atmospheric CO₂ throughout the year and constituted a net sink of 14 g C m⁻² (1.2 mol C m⁻² yr⁻¹), with ice coverage and ice formation limiting the CO₂ uptake during winter (Shadwick et al., 2011a). They explained that the CO₂ uptake was largely driven by under-ice and open—water biological activity, with high subsequent export of organic matter to the deeper water column. These results emphasize the large regional variability of the annual net oceanic sink of atmospheric CO₂ and the importance to consider the local processes driving the exchange.

In the high-latitude Southern Ocean, Fransson et al. (2004) used a similar method as in our study, i.e., winter-to-summer deficits of nitrate in several regions, such as the polar front (APF), winter ice edge (WIE), and the seasonal ice edge (SIE). In the SIE and the APF, a net ocean release of CO₂ to the atmosphere of 0.1–0.5 mol m⁻², respectively, was calculated over a time scale of several months (from austral winter to January). In the WIE, the ocean acted as a net atmospheric CO₂ sink of about 0.1 mol m⁻², which is similar to our values for the MWD for the period January to May (Table 1).

Dissolution and Sources of CaCO₃

Increased C_CALC suggests dissolution of CaCO₃, which may be caused by breakdown of CaCO₃ shells and skeleton from calcifying organisms or dissolution of ikaite minerals from sea ice melt water (Rysgaard et al., 2012, 2013; Fransson et al., 2013). Our estimate of C_CALC showed that the corresponding DIC change due to calcification or dissolution, was less significant than the C_BIG but at times similar to the values of C_EXCH (Figure 11 and Table 1). We found that the large C_CALC values in May were either found in the area influenced by sea ice formation and melt such as in MWD, or in the AWD domain in August. The shells (coccoliths) affects the reflectance of the surface water and results in a turquoise milky surface which is clearly observed using remotely sensed sensors (Tyrrell et al., 1999). In our study area and time of year, the remotely sensed data on particulate inorganic carbon (PIC_sat) showed a clear seasonal trend from values less than 0.1 µmol kg⁻¹ in May to maximum PIC values of up to 0.6 µmol kg⁻¹ in August (Figure 12). The seasonal PIC_sat trend agrees with our C_CALC estimates for the AWD but as previously stated, the values are too low to explain the DIC gain through CaCO₃ dissolution estimated from C_CALC of maximum of 0.3 and 0.7 mol C m⁻² in May and August, respectively (Table 1). Calcifying phytoplankton blooms are known to occur after the spring bloom and may sustain growth in relatively nutrient depleted waters during summer. This is observed on the Arctic inflow shelves such as the Bering Sea and Barents Sea and very common in the North Atlantic (e.g., Robertson et al., 1994), but not common in the Arctic Ocean (i.e., Tyrrell and Merico, 2004 and references therein). The low PIC_sat during our study agreed with observations of very low cell numbers of calcifying phytoplankton observed during the CarbonBridge study (Egge et al., 2018). Perhaps these shells were not locally formed and dissolved but transported from the south with the Atlantic water inflow or from the West Spitsbergen fjords (e.g., Lalande et al., 2016). This was the explanation for the E. huxleyi blooms, the most common and opportunistic calcifying phytoplankton, that was observed in the upper 50 m of the marginal ice zone in the Barents Sea in August 2003 (Hegseth and Sundsfjord, 2008). These blooms were attributed to intrusions of Atlantic water bringing cells of oceanic phytoplankton species via the subsurface circumpolar boundary current west of Svalbard and then eastward along the Eurasian Shelf break.

Another possible source for DIC gain through CaCO₃ dissolution is the dissolution of ikaite particles (which is a form
of CaCO$_3$ formed in sea ice) that has been recently released from melting sea ice and dissolved in the upper water column. Sea-ice cover will obstruct remotely sensed observations of ikaite and is most likely not included in the PIC$_{\text{sat}}$ values (Figure 12). Consequently, sea-ice derived ikaite could explain the relatively high DIC gain from C$_{\text{CALT}}$ of about 0.3 mol C m$^{-2}$ estimated between January and May. This implies that most of the C$_{\text{CALT}}$ increase of about 0.4 mol C m$^{-2}$ (from 0.3 to 0.7 mol C m$^{-2}$; Table 1) in the AWD between May and August is attributed to dissolution of advected CaCO$_3$ shells (Table 1). By the end of August, the C$_{\text{CALT}}$ values are the lowest throughout the study area (Table 1), agreeing well with the drastic decrease in PIC$_{\text{sat}}$ values (Figure 12).

**The Role of Nitrogen Fixation and Denitrification Based on N$^*$**

N$^*$ values from our study varied with season and ranged between $-3$ and $+2.5 \mu$mol kg$^{-1}$, which is the same range as reported from other ocean basins (Gruber and Sarmiento, 1997). The high values ($>1 \mu$mol kg$^{-1}$) in the AWD is generally found in well-oxygenated waters, such as in the North Atlantic > 35$^\circ$N (Gruber and Sarmiento, 1997). Consequently, the high N$^*$ in the eastern Fram Strait may be enriched by the inflowing Atlantic waters and does not necessarily imply a local source due to on-site N$_2$ fixation. To estimate the different nitrogen sources and sinks requires information on the isotopic nitrogen ratios (Granger et al., 2011). Sipler et al. (2017) estimated the depth-integrated N$_2$ fixation in the ice-free season (June to September) west of Svalbard and the Barents Sea to about 1.5 g N m$^{-2}$ in the upper 50 m. In the Nansen Basin, north of Svalbard, integrated N$_2$-fixation was significantly lower and between 0 and 0.5 g N m$^{-2}$ (Sipler et al., 2017).

The negative N$^*$ values observed on the shelf slope and in shelf bottom water, especially evident in August, may indicate a nitrogen loss due to benthic denitrification (Figure 6L). This was observed to be the case in the Pacific Arctic inflow region, in the eastern Bering Sea shelf where benthic denitrification caused a nitrogen loss of between 2 and 13 $\mu$mol L$^{-1}$ in April 2007 and 2008 (Granger et al., 2011). This implies that modifications such as increased nitrogen loss on the Bering Sea shelf may decrease the nitrogen concentrations in the Pacific water inflow waters and the surface water column in the Arctic Ocean (e.g., Jones et al., 2003). The Arctic outflow water exits through the Fram Strait, mainly in the East Greenland Current along the Greenland shelf (e.g., de Steur et al., 2014). It is unlikely that the negative N$^*$ values observed in the water column in January and May in the MWD was caused by Pacific water outflow. Denitrification has also been found in melting Arctic sea ice before the under-ice spring bloom oxygenates the surface water (Rysgaard et al., 2008). This could explain the nitrogen loss in the upper water column in January as well as the increasing N$^*$ values in the surface water as the bloom progressed and oxygenated the water from May to August (Figures 4L, 5L, 6L). However, sea-ice processes cannot explain the negative N$^*$ values before May in the deeper parts of the water column (>200 m). The decreased N$^*$ from January to May below 200 m depth for whole area, may be caused by contribution of another water mass. About half of the AW transported by the WSC, recirculated between 76 and 81$^\circ$N, exits the Fram Strait from the north, and has modified the chemical and physical properties of AW (MAW; Rudels et al., 2000; Marnela et al., 2013). According to Sipler et al. (2017), N$_2$ fixation was much lower in the Arctic Ocean than in the AW inflow area. Consequently, one explanation for the decreased N$^*$ between January and May could be that the AW looses nitrogen as it resides in the low N$_2$ fixation area further north and during its return to the Fram Strait contains less nitrogen as well as less heat. By August, the nitrogen increased by 2 $\mu$mol L$^{-1}$, showing a nitrogen gain, which implies a larger contribution of original AW. The strength and magnitude of the recirculation of MAW in the Fram Strait show large interannual variability (Rudels et al., 2000; de Steur et al., 2014). Moreover, observations and models show that eddy activity results in substantial seasonal and spatial variability in the recirculation and facilitates subduction of AW (Hattermann et al., 2016).

**CONCLUSION AND FUTURE OUTLOOK**

Phytoplankton DIC uptake (C$_{\text{BIO}}$) played by far the most important role for the observed DIC change throughout the study area and explained up to 89% of the total DIC change. The C$_{\text{EXCH}}$ played a minor to moderate role and was most significant in August in the Atlantic water domain, explaining about 36% of the relative importance of the DIC drivers. In May, dissolution of sea-ice derived CaCO$_3$ (ikaite) played a moderate but important role to explain the net effect of the DIC gain in all domains. By August, the biological DIC uptake (C$_{\text{BIO}}$) had increased in all domains, and at this time we observed the largest C$_{\text{EXCH}}$ gain, and continuing gain from CaCO$_3$ dissolution, most likely from an advected source. Of the total DIC gain between January and August (sum of C$_{\text{EXCH}}$ and C$_{\text{CALT}}$ ~2.8 mol C m$^{-2}$, Table 1), 25% was explained by CaCO$_3$ dissolution and the remaining 75% of C$_{\text{EXCH}}$ was due to ocean CO$_2$ uptake from the atmosphere as a result of the high biological CO$_2$ demand during photosynthesis between May and August.

In a future scenario, decreased sea ice and more open water exposed to atmosphere will facilitate direct ocean CO$_2$ uptake through increased air-sea CO$_2$ flux as long as the surface water is undersaturated in CO$_2$ relative to the atmospheric CO$_2$ level. In contrast, less sea-ice associated CaCO$_3$ dissolution (ikaite) will decrease the addition of total alkalinity, thus the buffering capacity against acidic input (e.g., CO$_2$), decreasing the CO$_2$ uptake potential in spring. However, with the influence of advected CaCO$_3$ shells later in summer, the buffering potential could be fully or partly restored but at a later stage. Moreover, less influence of melting sea ice may potentially decrease nitrate removal caused by denitrification. A larger inflow of well-oxygenated Atlantic water will also result in lower potential for denitrification to take place. Several studies show that increased CO$_2$ concentrations enhance primary production in spring in this area (Holding et al., 2015; Sanz-Martín et al., 2018), which would allow larger DIC uptake through biological CO$_2$ consumption. In that case, progressing ocean acidification in the
surface water would be mitigated fully or partly by biological CO₂ consumption. In a scenario of increased advection of Atlantic water, this buffer may become more important for the CO₂ uptake capacity and on-going ocean acidification. However, the net effect of the studied processes on the DIC change and ocean CO₂ uptake in the Arctic inflow region will ultimately depend on a combination of several processes such as changes in primary production, stratification, nutrient availability, the net carbon export out of the mixed layer, as well as changes in the advection of warm Atlantic water. Warming of the surface ocean will decrease the ocean CO₂ uptake solubility due to decreased CO₂ dissolution in warmer compared to colder water. With less meltwater in spring, the sea-ice ikaite contribution to the surface water would decrease, hence having consequences for the ocean to act as a net CO₂ sink in future. Furthermore, the increase in wind-induced vertical mixing due to increased open water in winter could contribute to increased DIC in the surface water, perhaps resulting in a CO₂ source.

DATA AVAILABILITY

The datasets generated for this study are available on request to the corresponding author.

AUTHOR CONTRIBUTIONS

MC, AF, and MV involved in the sampling and study design. MC performed the data analyses, and created the tables and figures. MC completed the writing of the manuscript with contributions from MV and AF. MV contributed with the calculations. KB contributed with the calculations based on the remotely sensed data. MC and AF collected and analyzed the carbonate chemistry.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmars.2019.00528/full#supplementary-material

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