

Key Points:

- We present the first multi-year, near-daily mixed layer total dissolved inorganic carbon time series in the Irminger Sea
- Annual net community production within the seasonally varying mixed layer is high ($9.8 \pm 1.6 \text{ mol m}^{-2} \text{ yr}^{-1}$) and has large interannual variability
- The greatest rates of inorganic carbon removal from the mixed layer via photosynthesis take place prior to mixed layer shoaling

Supporting Information:

Supporting Information may be found in the online version of this article.

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Net Community Production and Inorganic Carbon Cycling in the Central Irminger Sea

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Abstract The subpolar North Atlantic plays an outsized role in the atmosphere-to-ocean carbon sink. The central Irminger Sea is home to well-documented deep winter convection and high phytoplankton production, which drive strong seasonal and interannual variability in regional carbon cycling. We use observational data from moored carbonate chemistry system sensors and annual turn-around cruise samples at the Ocean Observatories Initiative's Irminger Sea Array to construct a near-continuous time series of mixed layer total dissolved inorganic carbon (DIC), $p\text{CO}_2$, and total alkalinity from summer 2015 to summer 2022. We use these carbonate chemistry system time series to deconvolve the physical and biological drivers of surface ocean carbon cycling in this region on seasonal, annual, and interannual time scales. We find high annual net community production within the seasonally varying mixed layer, averaging $9.8 \pm 1.6 \text{ mol m}^{-2} \text{ yr}^{-1}$ with high interannual variability (range of $6.0\text{--}13.9 \text{ mol m}^{-2} \text{ yr}^{-1}$). The highest daily net community production rates occur during the late winter and early spring, prior to the observed high chlorophyll concentrations associated with the spring phytoplankton bloom. As a result, the winter and early spring play a much larger role in biological carbon export from the mixed layer than traditionally thought.

Plain Language Summary The subpolar North Atlantic takes in more carbon from the atmosphere than other areas of the ocean relative to its size. This is partially caused by photosynthesis in the surface ocean, which turns inorganic carbon into organic carbon that is transported into the deep ocean in a process known as the biological carbon pump. Using measurements from sensors on moorings in the Irminger Sea, we construct a 7-year time series of the different parts of the inorganic carbon system. Using these, we separate out the forces that impact how much inorganic carbon has the potential to be exchanged with the atmosphere. We find that biological processes remove inorganic carbon from the surface ocean in the spring, summer, and early fall, while in the winter the surface ocean gets deeper and encompasses waters from below that have higher carbon content. The total amount of inorganic carbon removed from the surface ocean each year by biological process is extremely high in the Irminger Sea compared to other global ocean regions. This research highlights the importance of long-term, full year measurements to understand carbon cycle dynamics.

1. Introduction

The ocean acts as a key sink in the global carbon cycle, absorbing carbon from the atmosphere at its surface and then storing it in the deep ocean. Both biological and physical factors work in tandem to drive this uptake. The biological carbon pump transports organic carbon produced via photosynthesis into the deep ocean, causing a decrease in surface ocean $p\text{CO}_2$ that results in carbon dioxide moving from atmosphere to ocean (DeVries, 2022; Volk & Hoffert, 1985). Globally, the biological carbon pump moves approximately 10 PgC yr^{-1} into the ocean interior and is the main contributor to the 40-fold difference between the ocean and atmosphere carbon reservoirs (Caldera et al., 2018; Friedlingstein et al., 2022; Siegel et al., 2023). The North Atlantic accounts for a disproportionately large share of this global export relative to its size, with estimates ranging from 0.55 to 1.94 PgC yr^{-1} (average 1.27 PgC yr^{-1} ; Sanders et al., 2014). While extensive study has taken place in the North Atlantic on the myriad of pathways contributing to the region's biological carbon export, there is still high uncertainty in total flux values and the underlying mechanistic controls (Sanders et al., 2014).

Organic carbon meanders on its path from the surface to the deep ocean. It can be repeatedly aggregated, disaggregated, consumed, and respired by different classes of organisms, and is transported via sinking of particles, physical injection of high organic carbon waters, and vertical migration of heterotrophic grazers (Boyd, 2015; Boyd et al., 2019; Huang et al., 2022; Siegel et al., 2023; Stemmann & Boss, 2012). While known as the biological carbon pump, the physical processes involved in removing organic carbon from the surface cannot be siloed. In

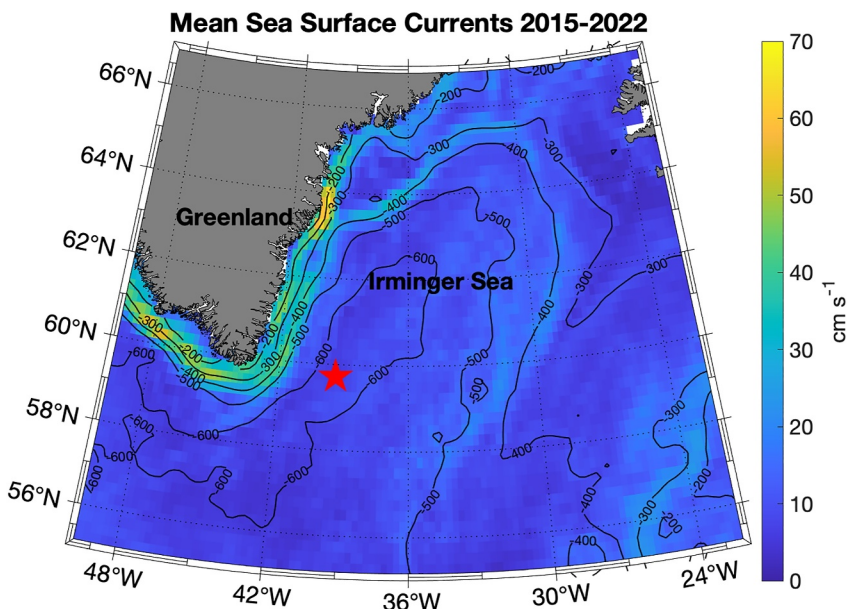


Figure 1. Map showing the Irminger Sea and location of Ocean Observatories Initiative Irminger Sea Array (red star). The colored shading is the mean sea surface currents and the black contours are the mean sea surface height in millimeters for 2015–2022, illustrating the counterclockwise rotation of the gyre, with the North Atlantic current bringing water northward to the east of the Array and faster moving currents bringing water southward to the west of the Array, along the eastern coast of Greenland. Data are from E.U. Copernicus Marine Service Information, <https://doi.org/10.48670/moi-00148>.

the North Atlantic, large spring phytoplankton blooms are thought to be the primary driver of high carbon export (Martin et al., 2011), however physical processes like deep convection have the potential to counterbalance this export by entraining previously exported carbon (Kortzinger et al., 2008; Palevsky & Nicholson, 2018; Quay et al., 2012, 2020). These dynamics are especially strong in the subpolar North Atlantic, which contains the deepest winter convection on Earth (de Jong & de Steur, 2016; de Jong et al., 2018; Holte et al., 2017) and high primary productivity (Boss & Behrenfeld, 2010; Henson et al., 2006, 2009).

To investigate the intertwined roles of biology and physics on carbon export, we use carbonate chemistry system measurements of total dissolved inorganic carbon (DIC) and total alkalinity (TA) which provide mechanistically agnostic tracers of the removal and addition of carbon from the surface ocean. Net community production (NCP) is a measure of the net biologically produced organic carbon in the upper ocean. It can be determined using inorganic carbon measurements and, when integrated throughout the year, represents the biological pump on an annual basis (ANCP, Emerson, 2014). Marine carbon cycling has been studied around the globe by making inorganic carbon system measurements at regular intervals throughout the year since the early 1980's (Bates et al., 2014). However, these historic discrete shipboard measurements have not provided adequate temporal resolution to constrain the full seasonal and interannual variability influencing ANCP using the carbonate chemistry system (Bates et al., 2014; Racapé et al., 2013). Technological advances since the turn of the century have increased observational capacity, providing high frequency time series measurements (more than once a day) for a number of chemical and physical parameters on both stationary mooring arrays and mobile floats and gliders. These have enabled the construction of DIC and TA mass balance budgets in the Northeast Atlantic, North Pacific, and Southern Ocean, and the subsequent disentanglement of biological and physical forcing on carbon cycling and quantification of annual net community production (Fassbender et al., 2016, 2017; Haskell et al., 2020; Huang et al., 2022; Knor et al., 2023; Kortzinger et al., 2008; Sauvé et al., 2023; Yang et al., 2021).

The Irminger Sea is an area of particular interest within the North Atlantic and is the location of the Irminger Sea Array (Figure 1, operated by the NSF's Ocean Observatories Initiative, OOI). The location was selected in order to provide sustained atmospheric, physical, and biogeochemical observations at a high latitude site, with a focus on the critical influences that affect the global ocean-atmosphere system (Ocean Observatories Initiative Science Prospectus, 2007; Smith et al., 2018). Fast moving currents bring water southward along the eastern coast of Greenland, and warm waters move northward along the eastern boundary as part of the North Atlantic current,

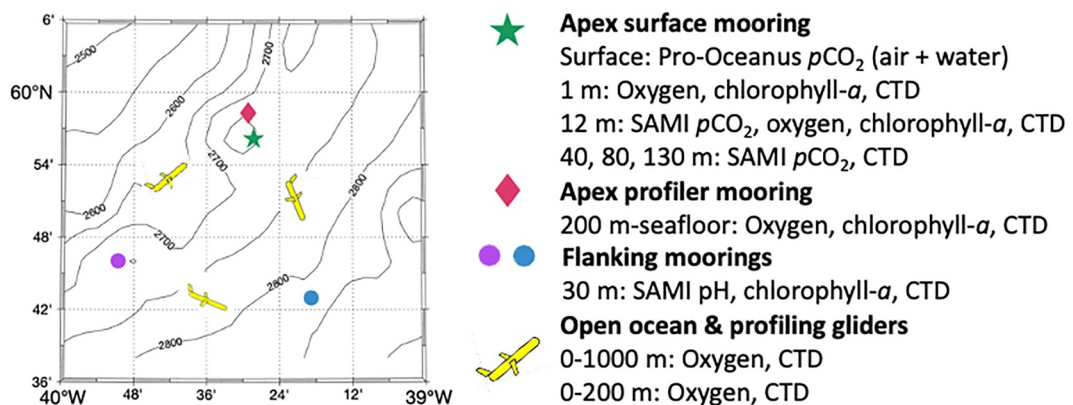


Figure 2. Schematic showing locations of the autonomous biogeochemical sensors and co-located CTDs on the OOI Irminger Array used in this analysis. Flanking Mooring A is west of Flanking Mooring B. Contour lines on the map show local bathymetry (m). The surface $p\text{CO}_2$ sensor is a Pro-Oceanus instrument; all other $p\text{CO}_2$ and pH sensors are Submersible Autonomous Moored Instruments (SAMI) made by Sunburst Sensors.

forming the central Irminger Gyre where the array is located (de Jong et al., 2018). A 19-year time series (2002–2020) using OOI and nearby mooring data finds extreme variability in the depth of winter convection (285–1,480 m, de Jong et al., 2024). Prior work in the region has used oxygen production as a tracer of photosynthesis and respiration (Palevsky & Nicholson, 2018) and the carbonate chemistry system seasonal cycle has been generally characterized (Bates et al., 2014; Racapé et al., 2013). Mixed layer net community production has been characterized throughout the subpolar North Atlantic during the highly productive period of the spring bloom, with rates often exceeding $60 \text{ mmol m}^{-2} \text{ d}^{-1}$ (e.g., Alkire et al., 2012; Baetge et al., 2020; Quay et al., 2012), however this strong seasonality prevents the individual snapshots from being used to calculate annual mean rates. As with many sites, there has not been sufficient data in the Irminger Sea to resolve full year-round net community production until now.

Understanding the controls on the biological pump in subregions like the Irminger Sea contributes to our overall understanding of the North Atlantic's current carbon export and the potential changes it may undergo as a result of climate change, including changes in primary productivity, surface and deep ocean respiration, and physical transport. Here, we use 7 years of high temporal frequency mooring data from the OOI Irminger Sea Array to provide the longest near-continuous daily carbonate chemistry data set to date in this region. We leverage these data to determine the biological and physical forcing on carbon cycling and quantify net community production over seasonal, annual, and interannual time scales in the subpolar North Atlantic.

2. Data

All data used in this analysis are publicly available; instructions for access are provided in Data Availability Statement at the end of this paper.

2.1. Ocean Observatories Initiative Irminger Sea Array

The Ocean Observatories Initiative (OOI) is a 25-year, NSF funded program that began collecting data in 2014, after nearly a decade of proposals and planning (Cowles et al., 2010; Isern & Clark, 2003; Smith et al., 2018). Our primary data source is the OOI Irminger Sea Array, which is comprised of 4 moorings, triangularly arranged and spaced approximately 20 km apart, and profiling and open ocean gliders (Figure 2). Repeat hydrography at the mooring locations is also available from annual turn-around cruises. Our analysis primarily uses the pH and partial pressure of carbon dioxide ($p\text{CO}_2$) sensors and their co-located CTDs in order to calculate DIC and TA, as described in Section 3. All carbonate chemistry sensors are located in the upper 130 m of the water column.

All carbonate chemistry system sensors, except at the surface, are Submersible Autonomous Moored Instruments (SAMI) made by Sunburst Sensors; both the pH and $p\text{CO}_2$ versions utilize spectrophotometry and a pH sensitive indicator dye (Álvarez et al., 2020; Lai et al., 2018). The surface $p\text{CO}_2$ sensor is a Pro-Oceanus CO_2 -Pro, which operates above and below the water, making air and seawater measurements using infrared detection (Jiang

et al., 2014). The pH and $p\text{CO}_2$ sensors below the surface make a measurement every 2 hr and the surface $p\text{CO}_2$ sensor makes a measurement every hour, though occasionally measurements are further apart due to battery limitations. Shallow sensors (pH sensors at 20 and 30 m; $p\text{CO}_2$ sensors at 0, 12, and 40 m) are in the mixed layer for the majority of the annual cycle, while deeper sensors (pH sensor at 100 m; $p\text{CO}_2$ sensors at 80 and 130 m) are only in the mixed layer in the winter. Two of the carbonate chemistry system sensors deployed by OOI (pH sensors at 20 and 100 m on the Apex Surface Mooring) have suffered from poor data return; given the limited raw data available prior to QA/QC, these sensors have been excluded from this analysis.

pH and $p\text{CO}_2$ data were quality controlled using gross range and spike tests based on Quality Assurance/Quality Control of Real Time Oceanographic Data (QARTOD) recommendations, followed by a moving median filter (Palevsky, Clayton, et al., 2023; U.S. Integrated Ocean Observing System, 2020; Table S1 in Supporting Information S1). We also remove data deemed suspect based on OOI operational annotations (for further detail, see the OOI Biogeochemical Sensor Data Best Practices and User Guide; Palevsky, Clayton, et al., 2023) and manually removed related stretches of data that indicate overall sensor failure. The fixed asset data within the mixed layer for pH, $p\text{CO}_2$, temperature, salinity, and pressure are at times sparse, with different depths and locations available across the time series (Figures S1 and S2 in Supporting Information S1). This sparsity is primarily due to infrastructure and instrument failures, which tend to occur during the late winter when seas are extremely rough and power becomes limited due to low sun angle. We remove all data from a given deployment if less than 5% of the data passes QA/QC.

Beyond the upper ocean carbonate chemistry system and CTD sensors, we use calibrated oxygen data from the wire-following profiler on the Apex profiler mooring (200–2,000 m, one cycle every 20 hr) and gliders (0–200 m and 0–1,000 m, varying temporal coverage). These are corrected using repeated stable deep oxygen measurements on the 3.1θ isotherm (\sim 1,800–2,000 m), based on the approach previous applied at the Irminger Sea Array by Palevsky and Nicholson (2018). These oxygen data are used for carbonate chemistry system predictions below the mixed layer using the CONTENT model (Bittig et al., 2018). We also use chlorophyll- α data from Sea-Bird/WETLabs fluorometers deployed on the OOI moorings at 1, 12, and 30 m during periods they are within the mixed layer. These data are presented for qualitative rather than quantitative interpretation, as end-user quality control and calibration of the fluorometer data are beyond the scope of this work.

2.2. Cruise Data

During the turn-around cruise each summer, discrete DIC, pH, and total alkalinity (TA) samples are collected at depths corresponding to the deployment depths of moored pH and $p\text{CO}_2$ sensors (Palevsky, Clayton, et al., 2023). In addition to the discrete samples routinely collected and analyzed by the OOI program, we collected and analyzed additional DIC and TA samples from the 2018 and 2019 turn-around cruises (Palevsky, Fogaren, et al., 2023). Quality control of the discrete DIC, pH, and TA samples provided by the OOI program identified inconsistencies among these three carbonate chemistry system variables that indicate data quality issues with a subset of the TA samples (Figure S3 in Supporting Information S1). We therefore rely primarily on directly measured discrete DIC data rather than on other measured or calculated carbonate chemistry system variables in calibrating and validating the sensor-based time series. Our analysis also uses calibrated salinity, temperature, and oxygen depth profiles from sensors on the CTD rosette (Fogaren & Palevsky, 2023; McRaven, 2022a, 2022b, 2022c, 2022d).

3. Methods

3.1. Carbonate Chemistry System Time Series

We synthesize the fixed mooring assets to construct near-continuous mixed layer DIC and TA, and surface $p\text{CO}_2$ time series over the 7-year period from summer 2015 through summer 2022. Neither DIC nor TA can currently be measured with commercially available autonomous sensors. Therefore, we leverage the fact that with any two measured carbonate chemistry system parameters (pH , $p\text{CO}_2$, DIC, and TA) and temperature, salinity, and pressure, the other two unknowns can be calculated.

In order to create as close to a complete mixed layer time series as possible for use in the mass balance budget, it was necessary to combine measurements and estimates of carbonate chemistry system parameters from a number of different sensors. For the DIC time series in particular, significant effort was undertaken to calibrate the

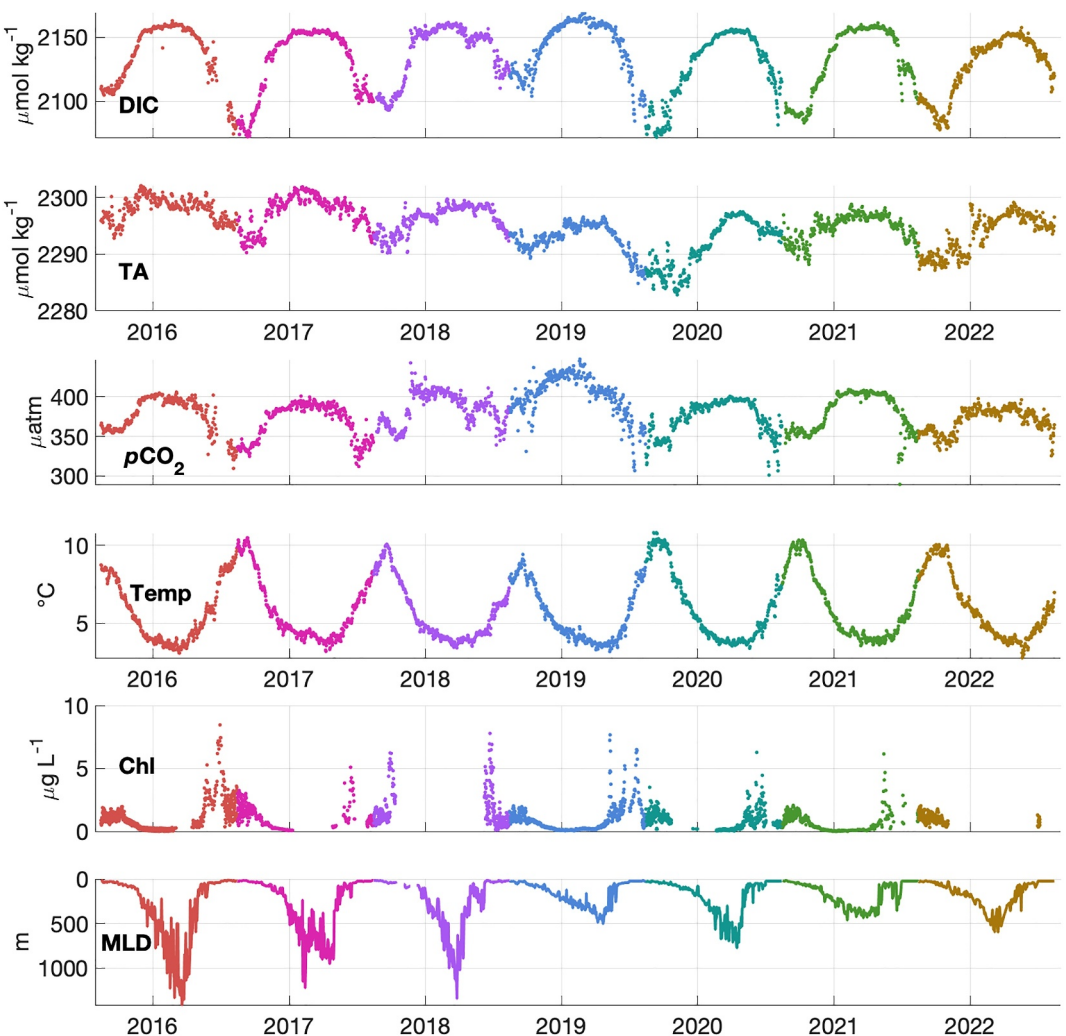


Figure 3. Daily mixed layer DIC, TA, $p\text{CO}_2$, temperature, chlorophyll- a , and mixed layer depth (MLD) time series. Colors correspond with annual cycles beginning on August 15th.

multiple partial datastreams prior to synthesis; a detailed description of the DIC moored time series calibration can be found in Text S3 of the Supporting Information S1. After calibration, the individual DIC time series from the separate sensor data streams are combined to create a daily mixed layer average. For the mixed layer time series, we use only sensors deployed at nominally 40 m and shallower. The mixed layer depth fluctuates significantly throughout the year, which results in the carbonate chemistry system sensors below the nominal 12 m sensor alternating being in the mixed layer and below it, depending on their deployment depths and the current depth of mixing (Text S4 in Supporting Information S1). Previous work has shown that the horizontal spatial variability is low enough to analyze sensors across the array as one data stream (de Jong et al., 2018; Palevsky & Nicholson, 2018). In the case of brief mooring data overlap (less than a week in summers 2018, 2019, and 2021), data from both moorings are presented in daily resolution figures (Figure 3). For the weekly mean budget calculations, data from both deployments are used to calculate the mean for that week.

3.1.1. Total Alkalinity (TA)

Due to the close co-varying relationship between total alkalinity (TA) and salinity, regressions between the two have long been used to estimate total alkalinity for carbonate chemistry system calculations (Millero et al., 1998), with other variables having been incorporated in more recent work. We use the Locally Interpolated Alkalinity Regression (LIRv2) model to predict TA from mixed layer sensor data, using temperature, salinity, and location as predictors (Carter et al., 2018). Uncertainty for estimated TA is generated from the LIRv2 model (Carter

et al., 2018), which incorporates the uncertainty in CTD measurements of temperature (0.005°C) and salinity (0.01). While other global and regional models exist to predict TA, we chose LIRv2 because it required fewer input variables than other models, such as CANYON-B which needs calibrated dissolved oxygen (Bittig et al., 2018). We found that the difference in the outputs is within the given uncertainty of the models. Comparison of the time series TA from the LIRv2 model agrees within uncertainty with nearly all available discrete TA calculated from DIC and pH samples (94%, Figure S8 in Supporting Information S1).

3.1.2. Total Dissolved Inorganic Carbon (DIC)

For each $p\text{CO}_2$ and pH sensor, DIC was calculated from daily averages of the sensor measurements and estimated total alkalinity (described above) using CO2SYS, with temperature, pressure, and salinity from co-located CTDs (Lewis & Wallace, 1998; Orr et al., 2018; Sharp et al., 2020; van Heuven et al., 2011). Silicate and phosphate are also calculation inputs as they contribute to the acid-base system, however DIC outputs are very insensitive to these inputs (<1 $\mu\text{mol kg}^{-1}$ difference calculated using the minimum and maximum values ever recorded in the region). As a result, we use mean nutrient values from regional GLODAPv2_2021 data (Lauvset et al., 2021; Olsen et al., 2016). We used the dissociation constants of Lueker et al. (2000), the KSO_4 constant of Dickson (1990), and the total boron constant of Uppström (1974). Days with fewer than 2 $p\text{CO}_2$ or pH measurements are excluded for each sensor. There are several short gaps in the time series where no carbonate chemistry system data is available. In the later part of the time series (summer 2018–summer 2022) we fill these gaps using DIC predicted using CONTENT from the calibrated 1 and 12 m moored oxygen sensors, with the longest gap being about 2 months in summer 2019 and all other gaps being less than 2 weeks (Figure S9 in Supporting Information S1). CONTENT is a Bayesian neural-network that predicts the carbonate chemistry system using dissolved oxygen, temperature, salinity, location, and time (Bittig et al., 2018). CONTENT is similar to its sister model (CANYON-B), however it is designed to assess the internal agreement of carbonate chemistry system parameters and prescribes an uncertainty value based on the goodness of agreement (Bittig et al., 2018). Along with the inputs of the data themselves, CONTENT also allows the user to input the measurement uncertainty, which is then factored into the uncertainty of the CONTENT predictions.

3.1.3. Calibrating the DIC Time Series

Our use of a multitude of carbonate chemistry system sensors requires confirmation that these sensors are accurately measuring our parameters of interest. Intercomparison among multiple array sensors when simultaneously measuring within the mixed layer show offsets greater than expected based on the manufacturer-provided uncertainties for factory-calibrated instruments (0.005 for pH and 2 μatm for seawater $p\text{CO}_2$; Palevsky, Clayton, et al., 2023), indicating that further calibration is needed. However, once calibrated, the DIC time series calculated from individual sensors in the mixed layer match well with validation data from deeper moored assets, as well as discrete cruise samples (Figure S4c in Supporting Information S1). We present the calibration process and the rationale for each calibration step below, with further details provided in Text S3 of the Supporting Information S1.

The SAMI $p\text{CO}_2$ sensors show significant offsets from one another and from the Pro-Oceanus $p\text{CO}_2$ sensor during some deployments, but follow the same seasonal, weekly, and daily patterns when in the mixed layer (Figure S4a in Supporting Information S1). The SAMI pH sensors also show offsets, though much less substantial. The offsets between the SAMI sensors are not consistent annually, and there is no offset seen in the temperature data of the co-located CTDs, which indicates a known sensor calibration issue that require correction rather than a true signal (DeGrandpre et al., 2004; Kortzinger et al., 2008). The shallower $p\text{CO}_2$ and pH sensors (12–40 m) used in the mixed layer time series are generally within a steep thermocline during turn-around cruises in summer, precluding aligning the moored sensor data with the discrete samples collected from co-located CTD casts as a calibration approach itself.

Unlike the SAMI sensors, when the Pro-Oceanus surface $p\text{CO}_2$ sensor measures for the entire deployment, the newly deployed sensor matches the previously deployed sensor while both are operational, which instills confidence in the accuracy of the surface Pro-Oceanus (summer 2020 and 2021, Figure S1 in Supporting Information S1). The Pro-Oceanus, however, fails after only weeks to months in five out of the seven deployments used in this analysis. In order to create a complete mixed layer time series, we leverage the strengths of both types of carbonate chemistry system sensors on the array, namely the longevity of the SAMI sensors and the accuracy of

the Pro-Oceanus. We correct for offsets in the DIC time series calculated from the shallow (12–40 m) SAMI pH and $p\text{CO}_2$ sensors by aligning them with the available DIC time series calculated from the Pro-Oceanus, after which we average all DIC data in the mixed layer (Text S3, Figure S4 in Supporting Information S1). The uncertainty in the individual components of the mixed layer DIC time series is estimated using CO2SYS's error propagation, with measured temperature, salinity, pressure, pH or $p\text{CO}_2$, and estimated TA uncertainties as inputs (Orr et al., 2018).

While we do not use the 80 and 130 m $p\text{CO}_2$ sensor data as part of our mixed layer time series, they provide a useful check on our calibration process. The final mixed layer DIC time series shows good agreement with these deeper sensor DIC data, calibrated using predicted DIC from turn-around cruise CTD dissolved oxygen measurements, once they reenter the mixed layer (Figure S4c in Supporting Information S1, Bittig et al., 2018; Palevsky, Fogaren, et al., 2023). This agreement with the independently calibrated deep sensors, as well as with the available discrete cruise data points, further instills confidence in the choice to align the 12–40 m DIC time series with the DIC time series from the Pro-Oceanus (Figure S4c in Supporting Information S1). A detailed explanation of the calibration processes can be found in Text S3 of the Supporting Information S1.

3.1.4. $p\text{CO}_2$

We fill in the gaps in the surface $p\text{CO}_2$ record directly measured by the Pro-Oceanus using $p\text{CO}_2$ calculated from the mixed layer DIC time series, mixed layer estimated total alkalinity, surface buoy salinity, and the fifth generation European Centre for Medium-Range Weather Forecasts atmospheric reanalysis (ERA5) sea surface temperature (Figure S10 in Supporting Information S1, Hersbach et al., 2020). When there are gaps in the surface buoy salinity data due to surface mooring platform failures, we assume a homogeneous mixed layer and use 30 m salinity (Figure S13 in Supporting Information S1). When both calculated and measured $p\text{CO}_2$ are available, the mean difference between the two is 2 μatm , which is within the measurement error of the Pro-Oceanus direct measurements (Palevsky, Clayton, et al., 2023).

3.2. Mixed Layer Budget

Seasonal changes in the observed mixed layer DIC concentration are driven by physical transport (entrainment), gas exchange, evaporation and precipitation (EP), and biological processes, namely net community production and calcium carbonate production and dissolution (Equation 1, Fassbender et al., 2016, 2017; Haskell et al., 2020; Kortzinger et al., 2008; Palevsky & Quay, 2017; Yang et al., 2021). Changes in total alkalinity are driven by the same processes as DIC, with the exception of gas exchange (Equation 2). The time rate of change terms for each tracer (left hand side of Equations 1 and 2, respectively) are determined from the DIC and TA time series data. Observational data allows us to calculate the non-biological drivers ($d\text{DIC}/dt$ Gas Exchange, $d\text{DIC}/dt$ EP, and $d\text{DIC}/dt$ Entrainment; $d\text{TA}/dt$ EP, and $d\text{TA}/dt$ Entrainment) and subtract from the weekly overall change ($d\text{DIC}/dt$; $d\text{TA}/dt$), leaving the biological drivers ($d\text{DIC}/dt$ Biology; $d\text{TA}/dt$ Biology), as the remainder. The sections below describe how each of the right hand side terms in these equations are calculated. All right-hand side terms in both budgets are calculated at weekly resolution. We smooth the observed carbonate chemistry system time series used in the budget with a 3 week running mean to remove high-frequency variability.

$$\frac{d\text{DIC}}{dt} = \frac{d\text{DIC}}{dt} \text{ |Gas Exchange} + \frac{d\text{DIC}}{dt} \text{ |EP} + \frac{d\text{DIC}}{dt} \text{ |Entrainment} + \frac{d\text{DIC}}{dt} \text{ |Biology} \quad (1)$$

$$\frac{d\text{TA}}{dt} = \frac{d\text{TA}}{dt} \text{ |EP} + \frac{d(\text{TA})}{dt} \text{ |Entrainment} + \frac{d\text{TA}}{dt} \text{ |Biology} \quad (2)$$

3.2.1. Gas Exchange

The rate of DIC change due to gas exchange is influenced by the $p\text{CO}_2$ in the atmosphere and surface waters, wind speed, temperature, and salinity. For calculating this rate, these driving forces are encapsulated by the piston velocity (k), $p\text{CO}_2$ solubility (K_H), atmospheric and surface water $p\text{CO}_2$ ($p\text{CO}_2_{\text{Air}}$, $p\text{CO}_2_{\text{Seawater}}$), and density (ρ), with positive values indicating a flux from the atmosphere into the ocean (Equation 3).

$$\frac{d\text{DIC}}{dt} \Big|_{\text{Gas Exchange}} = k * K_H * (p\text{CO}_2_{\text{Air}} - p\text{CO}_2_{\text{Seawater}}) * \rho \quad (3)$$

The piston velocity (k) is calculated using the Ho et al. (2006) wind speed-dependent gas transfer parameterization, implemented in the MATLAB `gas_toolbox` (Manning & Nicholson, 2022) and the Schmidt constant (Wanninkhof, 1992). Of the OOI moored assets, the surface buoy faced the most challenges with continuous data collection, limiting the ability to use directly measured meteorological data. Therefore, ERA5 products from the nearest grid cell (60°N, 39.5°W) are used for hourly sea surface temperature and 10 m wind speeds (calculated from v - and u -winds at 10 m, Hersbach et al., 2020). The ERA5 winds match well with the OOI 10 m measured wind speeds when available (mean difference -1.5 m s^{-1} , Figure S11 in Supporting Information S1). The slight overestimate of wind speed by the ERA5 product may mean that, though gas exchange is a small term in our DIC budget, our calculated contribution of gas exchange may be an overestimate. The solubility of CO₂ is calculated using the temperature and salinity dependent constant (K_H) from Weiss (1974). The mixed layer seawater $p\text{CO}_2$ data is calculated as described in Section 3.1.4. While the Pro-Oceanus $p\text{CO}_2$ sensor measures air $p\text{CO}_2$ on the OOI Irminger Sea Array surface buoy, the sensor fails frequently and there are no calibration or validation samples taken that can confirm the accuracy of this sensor. For this analysis, therefore, the air $p\text{CO}_2$ data is from NOAA's Global CarbonTracker gridded daily average product at 59°N, 40.5°W using CO₂ mole fraction, humidity, and barometric pressure (CT_2022 and CT-NRT.v2023-4, Jacobson et al., 2023; uncertainty estimated as 5.8°utm based on comparison with Pro-Oceanus air $p\text{CO}_2$ and with independent flask measurements, see Figure S12 in Supporting Information S1). Multiplying by the density (ρ) leaves us with the amount of carbon exchanged in units of $\text{mmol m}^{-2} \text{ d}^{-1}$.

3.2.2. Physical Transport

The fixed nature of moored sensors necessitates a Eulerian approach in order to account for the influence of horizontal and vertical transport on mixed layer DIC. The primary physical influence is entrainment, whereby the mixed layer increases in depth and adds water from below into the mixed layer. When the mixed layer shoals (detrainment), the mixed layer DIC concentration is not impacted. Given that this study site is in the center of the Irminger Gyre where there are low horizontal velocities (Figure 1), we exclude horizontal transport from our budget. We also exclude estimating the budget contributions of diffusive vertical mixing across the base of the mixed layer or of upwelling due to Ekman-driven mass divergence. Although these have been shown to be significant in mixed layer mass balance budgets in other regions (Fassbender et al., 2016, 2017; Knor et al., 2023; Palevsky et al., 2016; Yang et al., 2017), measured values for these terms have been shown to be low in the Irminger Sea and exceptionally deep winter convection at this site means that entrainment dominates over these second-order processes (Fratantoni, 2001; Painter et al., 2014; Våge et al., 2008).

In order to calculate the rate of physical entrainment, we use weekly mixed layer depths (MLDs). There has been extensive discussion within the literature regarding how to define the MLD with respect to timescales of mixing and biological processes (Carranza et al., 2018; Carvalho et al., 2017; Lacour et al., 2019). A challenge in defining a MLD appropriate for our analysis is differentiating between the actively mixing layer and remnant layers of prior deep mixing events that retain uniform physical properties but are no longer biogeochemically uniform because they are actively exchanging with the surface ocean (for a more extensive discussion, see Carranza et al., 2018). We use a combination of methods including temperature thresholds and bio-optical methods to determine the MLD depending on the time of year and availability of both OOI and external data (Table S2 in Supporting Information S1), a full explanation of which can be found in Text S9 of the Supporting Information S1.

We determine the effects of entrainment on mixed layer DIC, TA, and salinity using a one-dimensional model that accounts for the influence of mixing between the prior mixed layer and newly entrained waters, calculated at weekly time steps. This model uses the mixed layer depth time series to calculate physical entrainment rates. To determine the properties of the water being entrained from below the base of the mixed layer, we use dissolved oxygen, temperature, and salinity data from gliders and the wire-following profiler, from which DIC and TA are estimated using the CONTENT model (described in Section 3.1.2; Bittig et al., 2018). A detailed explanation of the entrainment model can be found in Text S10 of the Supporting Information S1. The entrainment model results track closely with observed concentrations in the fall, showing that the increase in mixed layer DIC during this

period of time is primarily the result of the entrainment of high DIC waters (Figure S16 in Supporting Information S1).

3.2.3. Evaporation and Precipitation

To account for dilution and concentration effects on DIC and TA due to evaporation and precipitation, we find the difference between the observed change in salinity over time and the entrainment-modeled change in salinity over time (Fassbender et al., 2016; Haskell et al., 2020).

We attribute the difference between the observations and the modeled change to evaporation and precipitation, as any change in salinity not due to entrainment is the result of these surface processes, again assuming minimal horizontal advection. We multiply the difference in the two by the ratio of DIC (and TA) to absolute salinity on the first day of each deployment, working under the assumption that the ratio at which salinity changes due to evaporation and precipitation relative to DIC (and TA) is constant throughout each deployment (Equation 4). The same process is followed for TA.

$$\frac{d(\text{DIC, TA})}{dt} \Big|_{\text{EP}} = \left(\frac{d\text{Sal}}{dt} - \frac{d\text{Sal}}{dt} \Big|_{\text{Entrainment}} \right) * \frac{(\text{DIC, TA})}{\text{Sal}} \Big|_{t=1} \quad (4)$$

3.2.4. Biological Processes

The biological drivers are calculated as the residual terms of Equations 1 and 2. These drivers are net community production (NCP) via photosynthesis and respiration and calcium carbonate (CaCO_3) formation and dissolution (Equation 5).

$$\frac{d(\text{DIC, TA})}{dt} \Big|_{\text{Biology}} = \frac{d(\text{DIC, TA})}{dt} \Big|_{\text{NCP}} + \frac{d(\text{DIC, TA})}{dt} \Big|_{\text{CaCO}_3} \quad (5)$$

In order to separate the soft tissue processes (photosynthesis and respiration, which influence NCP) from calcium carbonate production and dissolution, we leverage the known production ratio of CO_2 and H^+ (117 to -17) from respiration using 1 mol of phosphate (HPO_4^{2-}) (Anderson & Sarmiento, 1994; Fassbender et al., 2016). This formation and precipitation of CaCO_3 influences TA and DIC in a 2:1 ratio. This allows us to calculate the influence of NCP on the mixed layer DIC budget (Equation 6). We also calculate weekly and annual NCP within the seasonally varying mixed layer by integrating to the weekly mixed layer depth.

$$\frac{d\text{DIC}}{dt} \Big|_{\text{NCP}} = \frac{\left(\frac{d\text{TA}}{dt} \Big|_{\text{Bio}} - 2 * \frac{d\text{DIC}}{dt} \Big|_{\text{Bio}} \right)}{-2 + \frac{-17}{117}} \quad (6)$$

3.3. Uncertainty

Here we present the calculated uncertainties of the calibrated DIC, TA, and $p\text{CO}_2$ time series (described in Sections 3.1.1–3.1.4), as well as uncertainties of all terms in the DIC budget (Table 1). There is uncertainty in each of the terms of the mass balance budget (Equations 1 and 2, Table 1). The 95% confidence bounds of entrainment are calculated by taking the minimum and maximum values from the entrainment model runs with a combination of systematically over- and under-estimated mixed layer depths and the lower and upper uncertainties of the DIC concentration below the mixed layer. The evaporation-precipitation uncertainty is estimated using a Monte Carlo simulation iterated 5,000 times with the salinity and DIC entrainment uncertainty and the uncertainty of the DIC concentration on the first week of each deployment. The gas exchange uncertainty is also calculated using a Monte Carlo simulation to account for the uncertainties in temperature, salinity, $p\text{CO}_2$ of air and seawater, and a 20% assigned uncertainty in the gas transfer coefficient to account for uncertainty in the wind speed data as well as in the parameterized relationship between wind speed and gas transfer (Ho et al., 2006; Wanninkhof, 2014; Yang et al., 2021). Calculating the biological processes as the remainder from the mass balance budget means that the biological term contains the accumulation of the uncertainties from all the other budget terms. We use a wrap-around Monte Carlo containing all the previously discussed uncertainties (Table 1) in which we subtract the

Table 1
Uncertainties of Analyzed Parameters

Time series	Uncertainty	Primary sources of error
Calibrated DIC ($\Delta\text{DIC}_{\text{Observed}}$)	11.2–11.6 mmol m ⁻³	Measurement of pH and $p\text{CO}_2$, calibration of DIC time series (see Section 3.1.3)
TA ($\Delta\text{TA}_{\text{Observed}}$)	9.7–10.2 mmol m ⁻³	Estimate from LIRv2 model, measurement of salinity (see Section 3.1.1)
$p\text{CO}_2$	2 μatm	Measurement of surface $p\text{CO}_2$, calculation from DIC and TA (see Section 3.1.4)
$\Delta\text{DIC}_{\text{Entrainment}}$	8.7–10.3 mmol m ⁻³	DIC concentration below the mixed layer, mixed layer depth
$\Delta\text{DIC}_{\text{EP}}$	1.1–2.0 mmol m ⁻³	Entrainment salinity model from mixed layer depth and glider/WFP salinity measurements, ratio of DIC/TA to salinity on first day of deployment
$\Delta\text{DIC}_{\text{GE}}$	0.02–0.05 mmol m ⁻³	Air and sea $p\text{CO}_2$, wind speed, gas transfer coefficient
$\Delta\text{DIC}_{\text{NCP}}$	14.7–16.0 mmol m ⁻³	Combined uncertainty of all other budget terms, stoichiometry of photosynthesis-respiration and calcium carbonate formation-dissolution
$\Delta\text{DIC}_{\text{CaCO}_3}$	28.5–30.5 mmol m ⁻³	Combined uncertainty of all other budget terms, stoichiometry of photosynthesis-respiration and calcium carbonate formation-dissolution

Note. Values shown are the range of yearly means over all 7 years in the time-series.

physical transport, EP, and gas exchange terms from the overall change in DIC in order to determine the uncertainty in the biological processes.

4. Results and Discussion

4.1. Mixed Layer Time Series

Here we present mixed layer carbonate chemistry system variables (DIC, TA, and $p\text{CO}_2$) as well as their potential drivers (mixed layer depth, temperature, and chlorophyll-*a*) in the central Irminger Sea (Figure 3). The carbonate chemistry system variables follow a similar annual cycle of highs and lows, with the minimums occurring in late summer at the end of the productive season, then climbing throughout the fall to a maximum in winter, after which they are drawn down again in the spring (Figure 3).

In the Irminger Sea, sea surface temperatures vary significantly, from around 3°C in the winter up to 12°C in the summer. The $p\text{CO}_2$ is significantly affected by temperature (Weiss, 1974); if there were no biological processes reducing the $p\text{CO}_2$, we would expect to see the highest $p\text{CO}_2$ in the summer and the lowest $p\text{CO}_2$ in the winter, as solubility declines as temperature increases (Takahashi et al., 2002). Instead, as has been well documented in prior literature, the greatest $p\text{CO}_2$ are recorded in the winter and lowest are found in the summer due to biological drawdown, indicating that biophysical effects, rather than temperature, are the primary drivers of $p\text{CO}_2$ at our site (Bates et al., 2014; Landschützer et al., 2018; Takahashi et al., 2002).

The influence of vertical mixing and primary productivity can be clearly seen on the other, temperature-insensitive, carbonate system parameters (DIC and TA). In the spring and summer, the mixed layer is strongly stratified and high chlorophyll concentrations are recorded. At the same time, mixed layer DIC concentrations reach their minimum in the annual cycle, ranging from $2,074 \pm 8$ to $2,113 \pm 8 \mu\text{mol kg}^{-1}$ during our period of observation (Figure 4). It is important to note that while chlorophyll concentration does indicate primary productivity, the amount of inorganic carbon being utilized is not directly reflected, as widely varying chlorophyll to carbon ratios occur at different global locations, depths in the water column, and times of year (Sathyendranath et al., 2009). Due to infrastructure failures, the sensors are not always successful in capturing the entire spring bloom; however, in situ chlorophyll measured by fluorometry and satellite chlorophyll data confirm annual spring and fall blooms at the Irminger site (Painter et al., 2014).

In the fall, convection begins and the mixed layer starts deepening. The Irminger Sea's uniquely deep mixed layers are not observed until later in the season, with mixed layer depths rarely exceeding 100 m before mid November. Most of the interannual variability in mixed layer depth occurs in the winter, both in the timing of deepening and the maximum depth reached. Some years, such as 2015–2016, experience early and rapid deepening, but the deepest sustained winter mixing is usually reached in mid to late March, ranging from ~400 to ~1,300 m. The patterns observed here match previous analysis of MLDs in this region (de Jong et al., 2024). Shoaling in the spring begins gradually and then intensifies, going from near maximum depths to less than 100 m over the span of

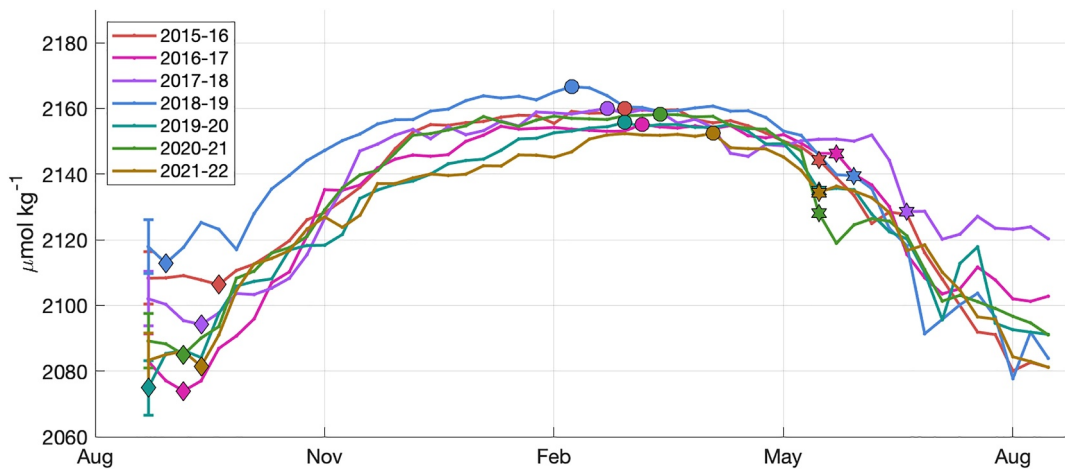


Figure 4. The annual seasonal cycle of DIC concentration in the mixed layer, beginning on August 15th. The diamonds are the lowest DIC concentration at the end of the productive season and the circles are the annual DIC maximum. The stars indicate the date at which the mixed layer first shoals above 100 m in the spring. The error bars at the beginning of the time series show uncertainty derived from the Monte Carlo simulation and apply across the time series. Gaps less than 2 weeks have been filled via linear interpolation.

a few weeks from early April to mid-May (Sterl & de Jong, 2022). During this time, we observe repeated shoaling and deepening, a process previously observed in the North Atlantic (Lacour et al., 2019). After the final shoaling, increased chlorophyll concentrations are observed again. It should be noted that the 7-year OOI time series captures a period of time in which there is particularly strong winter convection (3 winters deeper than 1,000 m), even for the Irminger region. Before the OOI time series, winter mixed layer depths for only 1 year between 2002 and 2013 (2011–2012) were close to 1,000 m (de Jong et al., 2024).

The mixed layer DIC signal generally increases and decreases in concert with the seasonal changes in mixed layer depth. The increase in mixed layer DIC coincides with mixed layer deepening (Figure 3). It is rapid in mid-September through mid-November and then continues to rise with a less significant slope between mid-February to late March, when it reaches the maximum annual concentration. The DIC concentration begins to decrease moderately in the late winter and then decreases more rapidly in late April–mid-May, coinciding with the shoaling of the mixed layer and high chlorophyll concentrations.

The amount of fall-winter increase and spring-summer decrease of mixed layer DIC varies from year to year. While there is a broad interannual range in the amplitude of the DIC seasonal cycle (mean 66 ± 13 , range 48 – $80 \mu\text{mol kg}^{-1}$), the range of the winter maximum weekly DIC concentration is only $14 \mu\text{mol kg}^{-1}$ (Figure 4). This is almost within the uncertainty of the mixed layer DIC time series ($\sim 11 \mu\text{mol kg}^{-1}$), making the annual maximum DIC effectively indistinguishable from year to year. The summer minimum is far more variable, reflecting more variability in the impact of biological DIC drawdown than winter convection on the mixed layer DIC annual cycle. The DIC seasonal cycle amplitude is similar to previous measurements in the Irminger Sea ($\sim 60 \mu\text{mol kg}^{-1}$, Bates et al., 2014) and other high latitude regions including the North Pacific ($\sim 56 \pm 7 \mu\text{mol kg}^{-1}$ at Ocean Station Papa, $73 \pm 2 \mu\text{mol kg}^{-1}$ in the Kuroshio Extension, Fassbender et al., 2016, 2017) and Western Antarctic Peninsula ($\sim 55 \mu\text{mol kg}^{-1}$, Yang et al., 2021), however the interannual variability observed in the Irminger Sea is higher.

4.2. Mixed Layer Carbon Mass Balance

To investigate the specific drivers of the carbonate chemistry system annual cycle and its interannual variability, we use a mass balance approach. Comparison among each of the terms influencing the DIC seasonal cycle (Equation 1) shows that entrainment of deep waters and biological processes have the greatest impact on the DIC seasonal cycle, with gas exchange and evaporation-precipitation playing relatively minor roles (Figure 5).

While gas exchange has very little impact on the overall mixed layer DIC concentration, the average total addition of DIC to the mixed layer due to CO_2 influx from the atmosphere is $2.1 \text{ mol} \pm 1.1 \text{ mol m}^{-2} \text{ yr}^{-1}$ over our 7 year

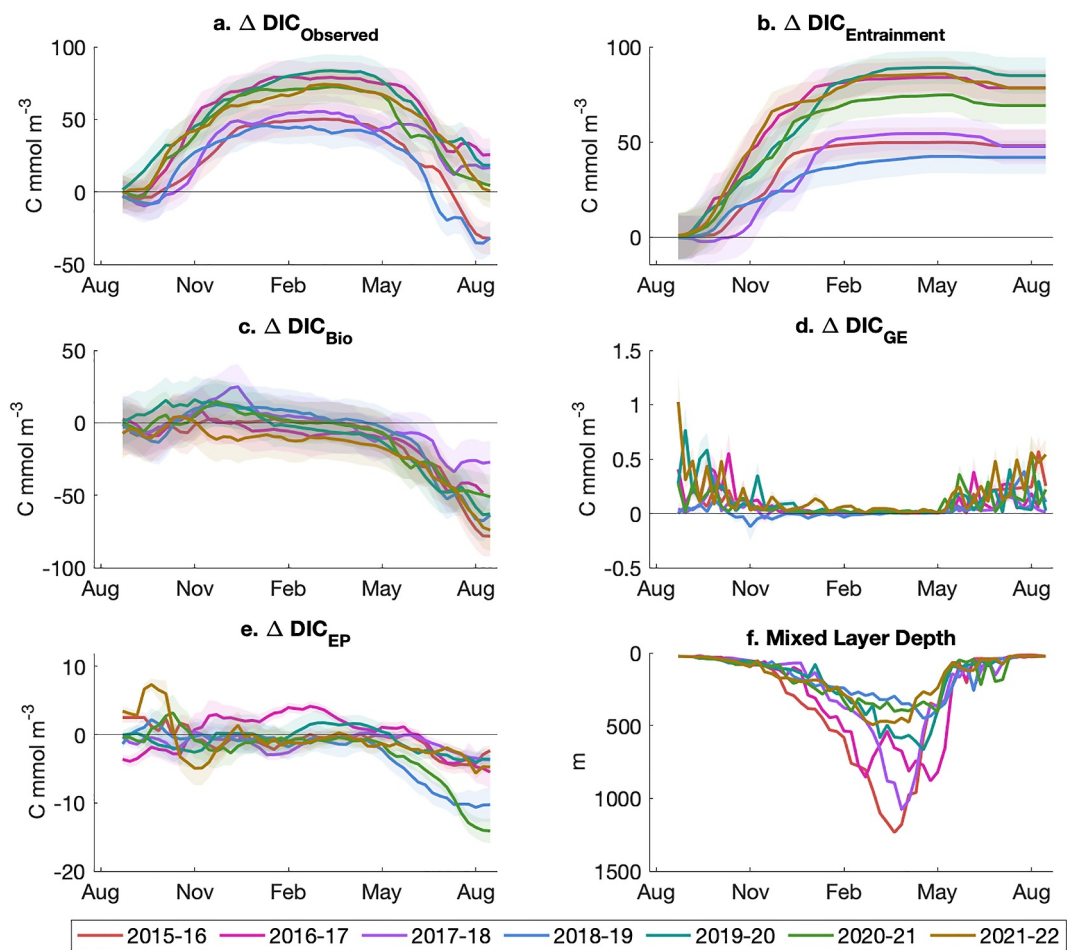


Figure 5. Influences of each term in the mixed layer DIC mass balance budget (Equation 1) on (a) the observed change in DIC concentration since August 15 over each of the 7 years in the time series. Changes in mixed layer DIC are driven by: (b) the change in DIC concentration due to entrainment, (c) the change in DIC concentration due to biological processes, (d) the change in DIC concentration due to gas exchange, and (e) the change in DIC concentration due to evaporation and precipitation, each presented as the cumulative change since August 15 in each year of the time series. Each of these terms is calculated within the seasonally varying mixed layer depth at weekly resolution (f). Note that the y-axes are scaled differently on each plot.

period. A recent $p\text{CO}_2$ climatological product reports a mean of $3.0 \pm 0.7 \text{ mol m}^{-2} \text{ yr}^{-1}$ during our study period (range $2.1\text{--}3.8 \text{ mol m}^{-2} \text{ yr}^{-1}$, Landschützer et al., 2020). This matches our observations within the uncertainty, and the somewhat higher values can likely be attributed to our differing choice of gas transfer parameterization (Atamanchuk et al., 2020). From our observational data, the years with the lowest annual carbon uptake via gas exchange are also the years when the maximum mixed layer depth exceeds 1,000 m (2015–2016 and 2017–2018), potentially indicating that the continued entrainment of high DIC waters suppresses winter uptake driven by cooling waters. Gas exchange occurs at the surface and therefore has the highest potential to impact DIC concentration when the mixed layer is shallowest (has the least volume). The most significant stratification occurs during the summer when the difference between air and seawater $p\text{CO}_2$ is greatest. These conditions would support strong influx, however the winds tend to be slow this time of year and do not drive vigorous gas exchange. In the winter when extremely high winds occur, the volume of the mixed layer has expanded by almost two orders of magnitude, so even the fastest gas exchange has very little impact on overall mixed layer DIC concentration, making gas exchange ($d\text{DIC}/dt_{\text{GE}}$) the least impactful driver of changes in mixed layer DIC (Figure 5d).

The impact of evaporation and precipitation on DIC concentration ($d\text{DIC}/dt_{\text{EP}}$) is also low throughout the year, and, as with gas exchange, has the greatest impact when the water column is stratified due to its influence occurring only at the air-sea interface. The impact of evaporation and precipitation varies from year to year but, on

average, mild dilution occurs in the summer months, reducing DIC concentration by $6.4 \pm 4.3 \text{ mmol m}^{-3} \text{ yr}^{-1}$ (Figure 5e). The dominant driver of DIC increase in the mixed layer is entrainment ($d\text{DIC}/dt_{\text{Entrainment}}$), which increases mixed layer DIC by $64 \pm 17 \text{ mmol m}^{-3} \text{ yr}^{-1}$. The DIC concentration rises concurrently with an increase in mixed layer depth as high DIC waters from below mix with the DIC-depleted surface waters. The concentration rises rapidly even though the mixed layer does not deepen extremely quickly because of the outsized role of the addition of water into the shallow end-of-summer mixed layer and the very low summer DIC concentration. A mixed layer of 20 m with DIC of $2,080 \text{ mmol m}^{-3}$ that entrains 80 m of $2,170 \text{ mmol m}^{-3}$ DIC waters then has a DIC concentration of $2,152 \text{ mmol m}^{-3}$, an increase of 72 mmol m^{-3} and roughly the amplitude of the seasonal cycle. Alternatively, a 100 m mixed layer with DIC of $2,152 \text{ mmol m}^{-3}$ that entrains 200 m of $2,170 \text{ mmol m}^{-3}$ DIC waters results in a DIC concentration of $2,164 \text{ mmol m}^{-3}$, only increasing the overall concentration by 8 mmol m^{-3} . The total amount of carbon added via entrainment varies based on each year's maximum mixed layer depth, but brings the surface concentration close to the same value each winter ($\sim 2,220 \text{ mmol m}^{-3}$). While we have primarily considered the extremely deep winter convection for its role in returning carbon to the surface ocean in the winter, the subsequent springtime shoaling of the mixed layer detains water that contains both inorganic and organic carbon in a process known as the seasonal mixed layer pump (Dall'Olmo et al., 2016). The intraseasonal and seasonal mixed layer pump has been shown to significantly contribute to carbon export in the subpolar North Atlantic, a process which is likely occurring at our study site (Lacour et al., 2019).

Biological processes ($d\text{DIC}/dt_{\text{Biology}}$) lead to a net drawdown of DIC on an annual basis (Figure 5c), with photosynthesis and calcium carbonate formation outpacing respiration and calcium carbonate dissolution. In the fall, the influence of biology is near zero, with slight heterotrophy in some years and autotrophy in others; however, the uncertainty crosses zero most years, obscuring any definitive fall trend. Each winter shows the same pattern: from February to May there is a slow but persistent decrease in DIC concentration due to biological processes. Once the mixed layer shoals, there is much more rapid drawdown, which comprises the bulk of the change in concentration due to biology over the annual cycle. While we cannot quantitatively relate the measured chlorophyll to DIC drawdown, the chlorophyll data clearly supports the rapid reduction due to primary productivity in the late spring (Figure 3). The calcium carbonate formation and dissolution is a very small portion of the overall change in DIC due to biological processes, accounting for $5.1 \pm 1.1 \text{ mmol m}^{-3}$ drawdown in DIC annually (Figure S17 in Supporting Information S1). This value falls within the average annual uncertainty of the $d\text{DIC}/dt_{\text{CaCO}_3}$ budget term (6.1 mmol m^{-3}) and thus we do not further interpret the role of calcium carbonate dissolution and formation.

4.3. Annual Net Community Production

The annual net community production (ANCP) within the mixed layer is the upper bound on the amount of carbon removed from the surface ocean by the biological pump each year ($\text{NCP} = -\Delta\text{DIC}_{\text{NCP}}$). The choice of depth of integration for calculating ANCP varies depending on the method of export research. This can lead to discrepancies when intercomparing ANCP rates, particularly in high latitude regions with deep winter convection, where a significant fraction of the carbon removed from the mixed layer during spring and summer is subsequently respired within the seasonal thermocline and re-entrained into the mixed layer during winter (Palevsky & Doney, 2018). Here, we leverage our mixed layer DIC budget to determine the seasonal cycle and interannual variability of NCP and ANCP integrated to the seasonally varying mixed layer depth. While not all of this NCP will lead to long-term carbon sequestration at depth, this value is of particular interest as the surface ocean $p\text{CO}_2$, and in turn the influx of carbon from the atmosphere to the upper ocean, is strongly influenced by NCP-driven changes in mixed layer DIC concentration.

We find net autotrophy in the Irminger Sea for most of the year, with the highest NCP rates occurring in the early spring prior to mixed layer shoaling (Figure 6). NCP rates in this paragraph are the mean and standard deviation across all 7 years. In the late summer into the late fall, we observe NCP from near neutral to mildly heterotrophic ($-5 \pm 17 \text{ mmol m}^{-2} \text{ d}^{-1}$ from August 15 to November 15). In January, there is a transition to autotrophy with an NCP of $24 \pm 43 \text{ mmol m}^{-2} \text{ d}^{-1}$ until the end of February. By April, shoaling has begun and the mixed layer is strongly autotrophic, with an NCP of $77 \pm 26 \text{ mmol m}^{-2} \text{ d}^{-1}$ from the beginning of April to mid-May. From then until August 15th, the mixed layer is relatively shallow and continues to be autotrophic at a much lower NCP of $28 \pm 24 \text{ mmol m}^{-2} \text{ d}^{-1}$.

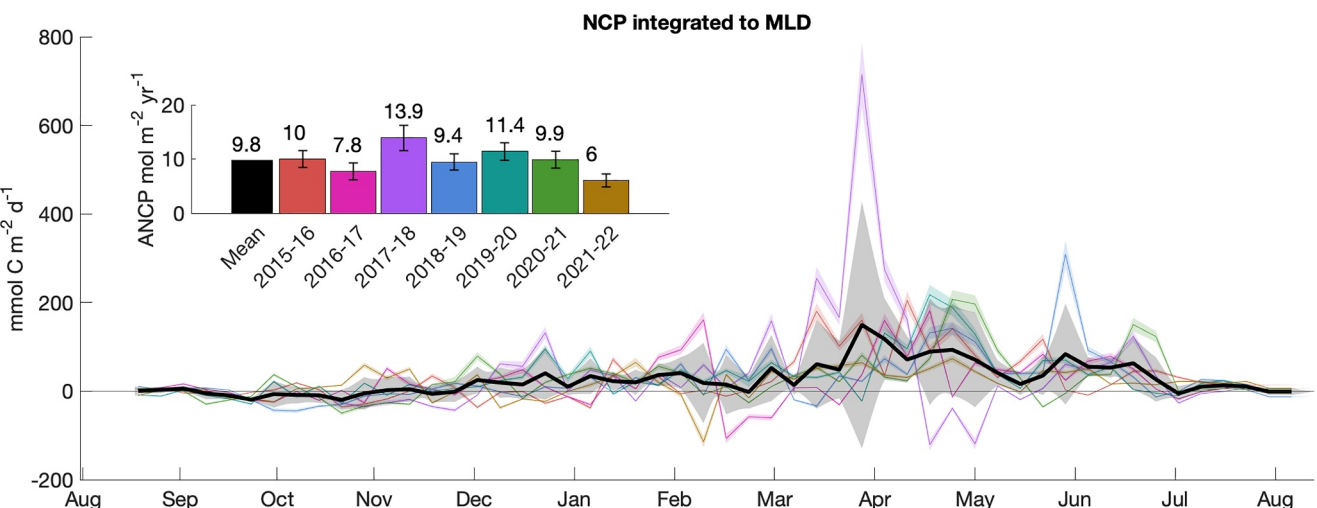


Figure 6. Net community production (NCP) rates integrated to the seasonally varying mixed layer depth. The colored lines show each individual year, the thick black line shows the mean across all years, and the gray shading is one standard deviation of the interannual mean. The bar chart shows total annual net community production (ANCP). ANCP error bars are the uncertainty as calculated through Monte Carlo analysis (details in Section 3.3).

We find that the central Irminger Sea has high ANCP as well as high interannual variability, with an annual mean of $9.8 \pm 1.6 \text{ mol C m}^{-2} \text{ yr}^{-1}$ (\pm signifies propagated error, not standard deviation, Figure 6). The highest ANCP recorded during our study period was $13.9 \pm 2.4 \text{ mol C m}^{-2} \text{ yr}^{-1}$ during 2017–2018. This year had a later-than-average onset of mixed layer deepening, resulting in a shallow fall mixed layer that caused the high fall chlorophyll concentration without a high depth-integrated rate of NCP. This is the only year with a decrease in DIC due to entrainment in the fall, likely indicating photosynthesis below the mixed layer during the large fall bloom. There is a late fall respiration signal, potentially fueled by sinking organic carbon from the fall bloom. 2017–2018 had very deep winter convection, as well as a decrease in DIC due to NCP from early March to mid-April that outpaced the time series mean (12 mmol m^{-3} reduction in DIC over this time period compared to the time series mean of 5 mmol m^{-3}), which together resulted in high integrated NCP over this time period and for the yearly total. The lowest ANCP recorded occurred during 2021–2022 ($6.0 \pm 1.3 \text{ mol m}^{-2} \text{ yr}^{-1}$), which also had the earliest spring shoaling of the observed period (Figure 3). DIC increased in the late fall due to both entrainment and respiration and there were relatively low NCP rates in the winter and early spring. These findings indicate that the depth and timing of winter mixing substantially impacts the overall annual net community production; however, the rate of DIC drawdown by biology prior to mixed layer shoaling is also an important control on ANCP.

This study adds to a growing body of research using DIC and TA as mass balance tracers to quantify ANCP within the seasonally varying mixed layer depth. The majority of sites have yielded lower ANCP than we find in the Irminger Sea ($2 \pm 1 \text{ mol C m}^{-2} \text{ yr}^{-1}$ in the subpolar Northeast Pacific, $1.2 \pm 2.8 \text{ mol C m}^{-2} \text{ yr}^{-1}$ in the North Pacific subtropical gyre, and $2.8 \pm 2.4 \text{ mol C m}^{-2} \text{ yr}^{-1}$ on the West Antarctic Peninsula shelf; Fassbender et al., 2016; Knor et al., 2023; Yang et al., 2021). However, in the Kuroshio extension in the western North Pacific, ANCP was $7 \pm 3 \text{ mol C m}^{-2} \text{ yr}^{-1}$ (Fassbender et al., 2017). The Kuroshio site has the deepest winter mixed layer depths of other sites where these methods have been used (up to 300 m), and similarly to the Irminger Sea, there are extremely high NCP rates seen in early spring as a result. A mixed layer carbon budget in the eastern subpolar North Atlantic (Porcupine Abyssal Plain site, 49°N , 16.5°W) found NCP within the seasonally varying mixed layer of $6.4 \pm 1.1 \text{ mol m}^{-2} \text{ yr}^{-1}$, with over two thirds of the production occurring prior to the spring shoaling of the mixed layer, a similar seasonal partitioning and annual magnitude as at our site (Kortzinger et al., 2008). Kortzinger et al. also found that 40% of the carbon exported from the seasonally varying mixed layer was subsequently offset by entrainment of respiration carbon during deep winter mixing. Reduced ANCP when integrating to the winter mixed layer depth as compared to ANCP within the seasonally varying mixed layer has been corroborated by oxygen budgets using data from across the subpolar North Atlantic (Quay et al., 2012, 2020). Given the uniquely deep winter mixing in the Irminger Sea, it will be important for future work to contextualize our ANCP results by determining what fraction of this NCP contributes to long-term carbon

sequestration, as well as the mechanistic relationship between NCP-driven DIC-drawdown in the mixed layer and air-sea CO_2 flux.

Our chemical tracer approach cannot provide insight into the complex ecosystem level dynamics that likely contribute to the observed high interannual variability in ANCP, however there is a clear connection between strong convection and high ANCP. Prior work investigating interannual variability of phytoplankton productivity in the Irminger Sea based on satellite chlorophyll data (which therefore does not capture early spring NCP discussed here) found that the onset of elevated chlorophyll concentrations varies by as much as 30 days (Henson et al., 2006). Their analysis found that timing of spring mixed layer shoaling influenced the timing and magnitude of the spring bloom; stormier winters with a high number of days with gale-force winds in turn delayed spring stratification, corresponding to later onset of elevated chlorophyll and lower peak chlorophyll concentrations (Henson et al., 2006). However, our work shows significant DIC drawdown by NCP within the deep late winter-early spring mixed layer, prior to mixed layer shoaling and increase in surface chlorophyll concentrations. During our period of observation, between 59% and 90% (mean across all years of 73%) of annual NCP occurs prior to the mixed layer shoaling above 100 m for the first time in the spring. Our finding of elevated NCP during deep winter mixing is consistent with the disturbance-recovery spring bloom hypothesis, in which the decoupling of zooplankton predator and prey during deep mixing enables enhanced primary productivity due to relief of grazing pressure (Behrenfeld & Boss, 2014). Historically, increased light availability with a shoaling mixed layer has been thought to be the catalyst of increased phytoplankton growth (Sverdrup, 1953), however more recent work has demonstrated that though high chlorophyll concentrations may be driven by this critical depth hypothesis, a combination of more complex ecosystem interactions caused by winter mixing are at play (Behrenfeld & Boss, 2014 and references within; Mignot et al., 2018). Our results suggest that deeper mixed layers in the later winter and early spring may actually drive higher ANCP, potentially the result of inhibition of grazing due to mixing.

5. Conclusions

This work constructs the first multi-year, high-frequency time series of the inorganic carbon system in the subpolar North Atlantic using daily measurements of the carbonate chemistry system. Strong biological drawdown is the primary removal mechanism of inorganic carbon from the mixed layer. Increases in mixed layer DIC are primarily controlled by entrainment of high DIC waters as the mixed layer deepens due to winter convection. Similar maximum DIC concentrations are found each winter despite interannual variability in winter mixing, implying that variations in the depth of winter convection do not drive wintertime mixed layer DIC concentrations. While previous analysis at this site has emphasized the role of deep winter mixing reintroducing respired carbon to the mixed layer that has been previously exported (Palevsky & Nicholson, 2018), we find that strong NCP in the late winter and early spring begins reducing DIC concentration prior to shoaling in all years, such that years with extremely deep convection actually drive higher ANCP from the seasonally varying mixed layer than those with shallower convection. The highest rates of NCP occur prior to the appearance of high chlorophyll concentrations, highlighting the utility of in situ sensing of carbonate parameters rather than relying on chlorophyll measurements as a proxy for biological productivity in the mixed layer. It is probable that the detrainment of water containing freshly produced organic carbon in the spring contributes to a high magnitude of carbon export, a subject which warrants further exploration. Future work is also needed to determine what fraction of the ANCP contributes to long-term carbon sequestration and to elucidate the connection between biologically driven DIC-drawdown and air-sea CO_2 flux.

The average annual net community production within the seasonally varying mixed layer is $9.8 \pm 1.6 \text{ mol C m}^{-2} \text{ yr}^{-1}$ and ranges from 6.0 to $13.9 \text{ mol C m}^{-2} \text{ yr}^{-1}$ over the 7-year study period. Sparsity of data often leads to averaging across multiple years in ocean biogeochemistry and specifically when determining ANCP, however doing so can blur the important differences from year to year in both the drivers and magnitude. Averaging across multiple years does not sufficiently capture the complex carbonate chemistry system dynamics in the central Irminger Sea, driven by high interannual variability of winter convection and primary productivity. Collecting observational data is both costly and challenging, however if only 1 year of data is collected or multiple years are averaged together, ANCP in areas with high interannual variability like the Irminger Sea will be misrepresented. This is highly relevant in the context of efforts to detect the emerging impacts of ongoing anthropogenic climate change on biogeochemical cycles relative to baseline natural variability (Henson et al., 2016). The intended 25-year time series of the OOI Irminger Sea Array will provide more information about

the interconnected physical and biogeochemical controls on interannual variability in ANCP, as well as potential long-term trends.

Data Availability Statement

Data Availability: Ocean Observatories Initiative mooring and glider data used in this analysis are all from the THREDDS Gold Copy catalog (<https://thredds.dataexplorer.oceanobservatories.org/thredds/catalog.html>).

Reference designators for each depth and deployment can be found in the OOI Data Explorer (<https://oceanoceans.org/knowledgebase/how-to-decipher-a-reference-designator/>, <https://dataexplorer.oceanobservatories.org/>). OOI DIC and TA bottle samples were accessed through the OOI Alfresco portal (<https://alfresco.oceanobservatories.org/alfresco/faces/jsp/browse/browse.jsp>).

DIC and TA bottle samples run at Boston College can be accessed at <https://www.bco-dmo.org/dataset/904722> (Palevsky, Fogaren, et al., 2023). ERA5 Reanalysis hourly sea surface temperature and 10 m wind speed data, used for mixed layer calculations and air-sea gas exchange, are from <https://doi.org/10.24381/cds.adbb2d47> (Hersbach et al., 2023). Sea surface height and current data used in Figure 1 are from E.U. Copernicus Marine Service Information, <https://doi.org/10.48670/moi-00148> (Global Ocean Gridded L4 Sea Surface Heights And Derived Variables Reprocessed, 1993-Ongoing, n.d.). NOAA's CarbonTracker data, used in air-sea gas calculations, are from <https://doi.org/10.15138/ffxv-2z26> (Jacobson et al., 2023). Nutrient mean values at our site are from GLODAPv2_2021, accessible at <https://www.ncei.noaa.gov/data/oceans/ncei/ocads/data/0237935/> (Lauvset et al., 2021).

Software Availability: All analyses were conducted in MATLAB from MathWorks. Basic physical oceanographic conversions were done using the Gibbs SeaWater (GSW) Oceanographic Toolbox of TEOS-10, available at <http://www.TEOS-10.org>. Calculations within the carbonate chemistry system used CO2SYS, accessible at <https://doi.org/10.5281/zenodo.3952803> (Sharp et al., 2020). Predicted DIC and TA were determined using the CONTENT model, accessible at <https://github.com/HCBScienceProducts/CANYON-B> (Bittig et al., 2018). Gas transfer parameterizations were calculated using the Gas_Toolbox, accessible at <https://doi.org/10.5281/zenodo.6126685> (Manning & Nicholson, 2022).

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