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Key Points:
- New parameterizations for total alkalinity and dissolved inorganic carbon allow higher-resolution estimates of oceanic anthropogenic CO2
- Applying our parameterizations to autonomous biogeochemical Argo floats data could allow more detailed mapping of parameters influencing global oceanic CO2 cycles

Supporting Information:
- Supporting Information S1
- Figure S1

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Abstract Increasing marine uptake of anthropogenic CO2 (C_{am}) causes global ocean acidification. To obtain a high-resolution spatiotemporal distribution of oceanic carbon chemistry, we developed new parameterizations of the seawater total alkalinity, and dissolved inorganic carbon from the ocean’s surface to 2,000-m depth by using dissolved oxygen, water temperature (T), salinity (S), and pressure (P) data. Using the values of total alkalinity and dissolved inorganic carbon predicted by dissolved oxygen, T, S, and P data derived from autonomous biogeochemical Argo floats, we described the distribution of oceanic C_{am} in the 2000s in the subarctic North Pacific at high spatiotemporal resolution. The C_{am} was found about 300 m deeper than during the 1990s; its average inventory to 2,000 m was 24.8 ± 10.2 mol/m², about 20% higher than the 1990s average. Future application of parameterizations to global biogeochemical Argo floats data should allow the detailed global mapping of spatiotemporal distributions of CO2 parameters.

Plain Language Summary Ocean absorbs the increasing atmospheric CO2 by human activities from 1750s and encourages global ocean acidification. To obtain the human-activity-derived CO2 in the subarctic North Pacific in a high resolution, we applied our empirical ocean carbon chemistry equations using other hydrographic parameters to autonomous biogeochemical Argo floats data. The amount of human-activity-derived CO2 in this region was found about 300 m deeper than during the 1990s and about 20% higher than the 1990s average. Our method allows the development of a system for monitoring long-term trend changes in ocean carbon chemistry similar to other time series stations.

1. Introduction
The global carbon cycle is of increasing interest due to the greenhouse effect of carbon dioxide (CO2) on the Earth’s climate. Since the 1750s, the ocean has played an important role as a sink of anthropogenic CO2, absorbing almost one fourth of such emissions (Le Quere et al., 2014). Increasing marine CO2 levels have mostly negatively impacted marine organisms by causing ocean acidification and reducing the ocean’s buffering capacity (Doney et al., 2009).

To better understand changes in oceanic carbon chemistry, oceanographers have collected high-precision carbon chemistry data, including total alkalinity (TA), dissolved inorganic carbon (DIC), and pH within 0.1% precision, throughout the global ocean from the surface to the bottom waters at decadal-scale temporal resolutions since the last several decades of the twentieth century (Suzuki et al., 2013). However, such data were collected mainly during research cruises, limiting their value; ship-based data cannot provide information on annual, seasonal, or monthly variability, and their spatial resolution (distance between observation stations) tends to be only 2–5° of latitude and longitude (~100–500 km). Although time series data sets for long-term oceanic CO2 variations from stationary observation sites are available in some representative regions (e.g., ocean stations ALOHA, Papa, and KNOT in the North Pacific; BATS in the North Atlantic), time series data sets across the global ocean are spatially sparse (e.g., the Carbon Dioxide Information Analysis Center: http://cdiac.ornl.gov/oceans).

To overcome these limitations and complement observed carbon chemistry data, researchers have developed parameterizations and models of oceanic CO2 (Lee et al., 2006; Nakano & Watanabe, 2005; Williams et al., 2016). They use other hydrographical properties measured at higher spatial and temporal resolutions than carbon chemistry, including dissolved oxygen (DO), temperature (T), salinity (S), chlorophyll-a (Chl),
and nitrate (NO₃⁻), to clarify spatiotemporal changes in oceanic carbon chemistry (Alin et al., 2012; Juranek et al., 2011; Li et al., 2016; Nakano & Watanabe, 2005; Takeshita et al., 2013).

Starting in 2000, researchers deployed an array of autonomous floats across the global ocean to monitor large and long-term oceanic changes as part of the Argo mission. These floats measure T and S from the sea surface to a depth of 2,000 m every 5–10 days (Carval et al., 2017). Over 3,000 Argo floats are now active and delivering real-time quality controlled data with high accuracy and horizontal/vertical resolutions, making it possible to analyze basin-scale oceanic changes more accurately than ever before. Furthermore, recently developed biogeochemical Argo floats (BGC-Argo) that can sense DO, NO₃⁻, Chl, and more have been increasingly deployed (Johnson et al., 2017; Thierry & Bittig, 2016). Although the BGC-Argo project intends to develop a global array, only 8% of Argo floats currently have DO sensors (DO-Argo), and quality control is not yet complete due to complex procedures (Johnson et al., 2017; Takeshita et al., 2013; Thierry & Bittig, 2016). Thus, there is great potential for improved understanding of oceanic carbon chemistry processes when the current challenges are overcome although the use of DO-Argo data has so far been limited to some pilot studies of eddies, mixed layer depth and ocean circulation, and biological properties for net production (Bushinsky et al., 2017; Inoue & Kouketsu, 2015; Riser & Johnson, 2008; Takeshita et al., 2013).

To obtain an oceanic carbon chemistry data set with the same spatiotemporal scale as the high-resolution Argo physical and biological data sets, parameterization techniques have been applied to the Argo data sets (Juranek et al., 2011; Li et al., 2016; Williams et al., 2015). Simple, high-precision parameterizations of oceanic TA, DIC, and pH have previously been applied to DO, T, and S observational data collected by Argo floats in the subarctic North Pacific and used to produce quasi-real-time high-resolution maps of the vertical distributions of carbon chemistry to 400-m depth (Li et al., 2016). Moreover, the North Pacific Intermediate Water, a cold low-salinity mass produced in the western subarctic North Pacific, affects the climate and ocean circulation of the entire North Pacific (e.g., Ueno & Yasuda, 2003; Yasuda, 1997). Recent studies have shown strengthening ocean stratification in the subarctic North Pacific (Watanabe et al., 2001), which is expected to decrease both biological productivity and the ocean’s capacity to buffer anthropogenic CO₂ (Doney et al., 2009; Intergovernmental Panel on Climate Change, 2013).

Therefore, we chose this region to test new parameterizations of oceanic carbon chemistry and their application to hydrographical data with high spatiotemporal resolutions obtained from Argo profiling floats. We used these reconstructed TA and DIC values along with oceanic carbon chemistry thermodynamics to calculate the quasi-real-time distribution of anthropogenic CO₂ (Cₐ) at a high resolution in the subarctic North Pacific and examined the regional changes in Cₐ over the last several decades.

2. Materials and Methods

2.1. Improved Parameterizations of TA and DIC in the Subarctic North Pacific

A previous study (Li et al., 2016) provided parameterizations for TA, DIC, and pH from 40- to 400-m depth in the subarctic North Pacific. By adding a pressure (P) parameter, we could provide new parameterizations for TA, DIC, and pH to 2000-m depth in the same region covering most of the subarctic North Pacific (40–56°N, 145°E–130°W, Figure S1). Because DO, T, S, and P have F values of more than 2.4 (Wilks, 2011, Tables 1 and S2), we can use the values of DO (μmol/kg), T (°C), S (dimensionless), and P (dbar) to reconstruct the predicted TA and DIC (TApred and DICpred; see equation S1 for pH):

\[ TApred = 486.3 - 0.1031 \text{DO} - 4.992 \ T + 54.03 \ S + 0.04155 \ P \]  \quad (1)

\[ DICpred = 963.5 - 0.6485 \text{DO} - 17.39 \ T - 42.70 \ S + 0.009313 \ P \]  \quad (2)

Equations (1) and (2) were obtained using the observed high-precision carbon chemistry and hydrographic data sets for DO, T, S, and P collected by the Climate Variability and predictability (available from the Carbon Dioxide Information Analysis Center: http://cdiac.ornl.gov/oceans) and Pacific Ocean Interior Carbon international programs from 2000 to 2010 (Suzuki et al., 2013). The data quality after 2000 were controlled by using Certified Referenced Materials distributed by A.G. Dickson of the Scripps Institution of Oceanography.
2.2. Application of Parameterizations and Argo-Based DO, \(T\), and \(S\) Data to the Estimation of the Cant and Disequilibrium DIC (Cdiseq)

According to Sarmiento and Gruber (2006), the sum of the anthropogenic DIC (\(C_{\text{ant}}\)) and disequilibrium DIC (\(C_{\text{diseq}}\)) can be calculated as \(C_{\text{res}}\) as follows:

\[
C_{\text{res}} = C_{\text{ant}} + C_{\text{diseq}} = C_{\text{DIC}} - C_{\text{pi, sat}} - C_{\text{soft}} - C_{\text{carb}}
\]  

where \(C_{\text{pi, sat}}\) (preindustrial saturated DIC) and \(C_{\text{res}}\) are preformed components, and \(C_{\text{soft}}\) (derived from the remineralization of organic matter) and \(C_{\text{carb}}\) (derived from the dissolution of CaCO₃, see below) are regenerated components. We used the predicted DIC (\(\text{DIC}_{\text{pred}}\)) calculated by equation (2) instead of the observed DIC. For the parameterization of \(\text{DIC}_{\text{pred}}\), we used observed DIC data from the 2000s, which were influenced by a direct-anthropogenic component derived from the increasing level of anthropogenic CO₂ and an indirect-anthropogenic component derived from changes in \(\text{DO}\), \(T\), \(S\), and \(P\); thus, \(\text{DIC}_{\text{pred}}\) consists of \(C_{\text{ant}}\) from the 2000s and indirect-anthropogenic DIC at an arbitrary time. Here we assumed that the \(C_{\text{ant}}\) component of \(\text{DIC}_{\text{pred}}\) was constant at the \(C_{\text{ant}}\) value in 2005, determined by calculating the average \(C_{\text{ant}}\) value during 2000–2010.

\(C_{\text{pi, sat}}\) can be calculated from the equilibrium carbon chemistry relationships as follows:

\[
C_{\text{pi, sat}} = f(T\text{A}_0, \text{pCO}_2 = 280 \text{ ppm}, T, S, P)
\]  

where \(T\text{A}_0\) is preformed TA from the sea surface obtained from the linear relationship between \(T\text{A}_0 = 399 + 54.629\times S\), Millero et al., 1998). We calculated the contribution of organic matter remineralization to DIC as

\[
C_{\text{soft}} = 117/170 \text{ AOU}
\]  

where 117/170 is the stoichiometric ratio of carbon to oxygen (Anderson & Sarmiento, 1994) and AOU is apparent oxygen utilization, calculated as the difference between saturated DO and observed DO. Here we assumed that the stoichiometric ratio was constant vertically and used the N: C: O₂ ratio of 16: 117: −170 (Anderson & Sarmiento, 1994). We calculated the contribution from the dissolution of CaCO₃ as

\[
C_{\text{carb}} = 0.5(\text{TA}_{\text{pred}}-\text{TA}_0-16/170 \text{ AOU})
\]  

where 16/170 is the stoichiometric ratio of nitrogen to oxygen (Anderson & Sarmiento, 1994). The term 16/170AOU corrects the calculation for the fact that TA also includes nitrate formed by the remineralization of organic matter. The coefficient 0.5 indicates that a change of one mole in TA corresponds to a change of one-half mole in DIC.

We did not use the traditional method to calculate the disequilibrium DIC in seawater (Gruber et al., 1996), because we did not have any seawater age information. Therefore, assuming that the CO₂ disequilibrium value would be constant from the surface mixed layer to the ocean interior, we estimated the CO₂ disequilibrium value at the mixed layer depth (MLD, \(C_{\text{diseq, MLD}}\)) by subtracting the preformed and regenerated

| Parameterizations for TA and DIC in the North Pacific subarctic region |
|-----------------------------|-----------------------------|
| TA (\(n^a = 3956\))         | DIC (\(n = 4004\))          |
| \(R^2^b\)                   | RMSE^c  \(F^d\)  \(\text{VIF}^e\) | \(R^2\)  RMSE \(F\)  \(\text{VIF}\) |
| DO\(^f\)                   | 0.98  8.18  1394  4.3  0.99  5.60  118721  4.2 |
| \(T^g\)                    | 2762  2.0  5414  6.9  7228  6.9 |
| \(S^h\)                    | 7935  4.0  |

Note. DIC = dissolved inorganic carbon; TA = total alkalinity. 
\(^a\)Number of data points. \(^b\)Coefficient of determination. \(^c\)Root-mean-square error. \(^d\)Value from \(F\) test. \(^e\)Variance inflation factor. VIF > 10 is a standard threshold for assessing collinearity (Kutner et al., 2004).
components at the MLD [saturated DIC at MLD (Csat, MLD) and Csoft at MLD (Csoft, MLD, and Ccarb, MLD, respectively)] from DICpred at the MLD (DICpred MLD). Thus, Cdiseq, MLD in 2005 was calculated as

\[ C_{\text{sat,MLD}} = f(\text{TA}_0, \text{pCO}_2 = 375 \text{ ppm}, \text{T}, \text{S}, \text{P}) \]  

(7)

\[ C_{\text{diseq, MLD}} = \text{DIC}_{\text{pred, MLD}} - C_{\text{sat,MLD}} - C_{\text{soft,MLD}} - C_{\text{carb,MLD}} \]  

(8)

Because the DICpred value in 2005 included Cant, we used 375 ppm as the pCO2 value in 2005 in equation (7). We used the mean values of Cdiseq, MLD of each Argo float as Cdiseq for all periods of that float. Finally, we estimated Cant as

\[ C_{\text{ant}} = C_{\text{res}} - C_{\text{diseq}} \]  

(9)

### 3. Results and Discussion

By the 1990s, Cant had penetrated as deep as ~1,500 m in the subarctic North Pacific (Sabine et al., 2004). This suggests that parameterizations of ocean carbon chemistry should be considered to 2,000-m depth (the
measurement limit for BCG-Argo floats) in this region to examine how the distribution of Cant has changed. In addition, $P$ is a main factor affecting the dissociation constants of carbonic acid and the dissolution of calcium carbonate in the ocean, so that we developed parameterizations of TA and DIC to 2,000-m depth that considered $P$ (equations (1) and (2)).

For equation (1), the root-mean-square error (RMSE) was 8.2 μmol/kg and the coefficient of determination $R^2$ was 0.98; for equation (2), RMSE was 5.6 μmol/kg and $R^2$ was 0.99. These relationships were consistent.
Comparisons of Cdiseq Between Sabine et al. (2002) and This Study

Table 2

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<th>ID</th>
<th>C_diseq (μmol kg⁻¹)</th>
<th>C_diseq (μmol kg⁻¹)</th>
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aDifference between C_diss calculated in this study and in Sabine et al. (2002). bWater types defined in Sabine et al. (2002): 1c, 33.2 < S < 33.75 and θ < 4; 1e, S < 33.65 and θ < 4.2; 2a, S < 33.94 and θ > 2.6; 3c, 33.2 < S < 34.5 and DO < 50 μmol/kg (θ = potential temperature in degree Celsius).

with their parameterizations to 400-m depth, used to reconstruct oceanic carbon chemistry at weekly to decadal scales (Li et al., 2016, sections S1 and S2 and Table 1). Combining these parameterizations with C_{ant} while only considering the RMSEs of T_{A pred} and DIC_{pred}, we calculated C_{ant} the disequilibrium value of DIC (C_{diseq}), and their sum (C_{res}) with a precision of ±6.9 μmol/kg using DO, T, S, and P data over the subarctic North Pacific.

DO-Argo floats used in this study are mostly deployed in the western and eastern parts of the subarctic North Pacific. Although Argo DO data can be validated by comparison with discrete DO observation data obtained from bottle samples collected near the float locations, in this region such data are insufficient and too shallow to validate all DO-Argo data (Bushinsky et al., 2016). Therefore, we chose and tested two DO-Argo floats (WMO ID: 2900540 [west DO-Argo] and 5904106 [east DO-Argo], details in section S3) in the western and eastern gyres of the subarctic North Pacific. We then corrected the DO-Argo data using a linear fit to bottle DO data collected near each DO-Argo float (section S3 and Figure S5) and applied the new TA and DIC parameterizations to the DO, T, S and P of BGC-Argo data sets collected from the two gyres (Figure 1). The results showed that the vertical distributions of T_{A pred} and DIC_{pred} varied over a wider range and changed more smoothly in the east than in the west, mostly consistent with previous findings (Li et al., 2016).

We used the DIC_{pred} and T_{A pred} to calculate C_{ant}, C_diseq, and C_{res} using Argo DO, T, S, and P data over the subarctic North Pacific (Figure 2). Here C_{res} and C_{ant} represent the anthropogenic injection of CO_2 into the ocean until 2005 as an average from 2000 to 2010.

As we had no data on seawater age, we could not use the traditional method of estimating C_{diseq} due to disequilibrium in the air-sea exchange (Gruber et al., 1996). Instead, we assumed that the C_{diseq} would be constant vertically from the surface mixed layer and subtracted the preformed and regenerated components from DIC_{pred} at the mixed layer depth (MLD, C_{diseq, MLD}) as the C_{diseq} value. We used the mean C_{diseq, MLD} value of each Argo float as C_{diseq} for all periods. The C_{diseq, MLD} for west and east DO-Argo was −9.2 μmol/kg and −5.1 μmol/kg, respectively (Figure 2), corresponding to the order of magnitude for C_{diseq} (−13 ± 5 μmol/kg) calculated by the traditional trace age method in a previous study of the North Atlantic (Gruber et al., 1996). We also compared our C_{diseq} to another C_{diseq} determined by the mixing of different water types (Sabine et al., 2002); the difference was less than 6 μmol/kg (Table 2).

Most C_{diseq, MLD} values were negative because they represented an air-sea CO_2 exchange period of about 1/2 year (Figure 2). Some positive C_{diseq, MLD} values appeared in the summer (April–June, Figures 2a and 2d) for both Argo floats. This result indicates that a shallower MLD may facilitate the air-sea exchange of CO_2, making it temporarily easier to reach equilibrium. As we found that C_{diseq} was not always constant in a given region and changes both seasonally and locally, our new method for estimating C_{diseq} without seawater age information is useful for C_{ant} calculation and can be broadly applied to Argo or other higher-resolution ocean hydrographic data in the future.

In the subarctic North Pacific, C_{ant} reached depths deeper than 1,700 m, about 300 m deeper than C_{ant} in the 1990s (Figure 2). C_{ant} penetrated more deeply in the west than in the east and was relatively high; this result was consistent with the isopycnal distribution of C_{ant} calculated by a previous global-scale study (Sabine et al., 2004). In addition, west DO-Argo drifted at relatively low latitudes and therefore had a lower buffering factor than east DO-Argo, which drifted at higher latitudes. The average C_{ant} inventories of west and east DO-Argo were 32.3 and 17.3 mol/m², respectively. The average inventory of C_{ant} in the subarctic North Pacific to 2,000-m depth was therefore 24.8 ± 10.2 mol/m², about 20% higher than the average during the 1990s (Sabine et al., 2004). The rate of ocean uptake of CO_2 was 0.5 mol·m⁻²·year⁻¹ in this region, a relatively low value for the North Pacific (Sabine et al., 2008). This low value may reflect a strengthening of ocean stratification in the subarctic North Pacific (Watanabe et al., 2001).
4. Conclusions

In this study, we showed the Lagrangian variations of carbon species and \( C_{\text{sat}} \) when using Argo data. Furthermore, we demonstrated that Argo floats in a specific region can be connected to produce time series data. Combining Argo data with our parameterizations allows the development of a system for monitoring long-term trend changes in ocean carbon chemistry similar to other time series stations (e.g., ALOHA and KNOT). Argo floats equipped with pH sensors in the subarctic North Pacific will inevitably be deployed in the near future. Since the pH sensors give us the high-resolution spatiotemporal distribution of ocean pH, we will easily use the sensor pH and our parameterized pH to separate the direct and indirect anthropogenic effect on ocean acidification (Watanabe et al., 2018).

By using BGC-Argo or other automatic sensor data sets for DO, \( T, S, \) and \( P \), our parameterizations can produce detailed spatiotemporal distributions of ocean carbon chemistry and anthropogenic \( CO_2 \) above 2,000-m depth in quasi-real time over the subarctic North Pacific, which traditional ship-based methods cannot achieve. In the future, application of our parameterization method to global BGC-Argo data should make it possible to obtain a high-resolution map and detailed time series distribution of global oceanic \( CO_2 \) cycle parameters.

References


