Variability of Sea-Air Carbon Dioxide Flux in Autumn Across the Weddell Gyre and Offshore Dronning Maud Land in the Southern Ocean

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Sea surface fugacity of carbon dioxide ($fCO_{2}ssw$) was measured across the Weddell gyre and the eastern sector in the Atlantic Southern Ocean in autumn. During the occupation between February and April 2019, the region of the study transect was a potential ocean CO$_2$ sink. A net CO$_2$ flux (FCO$_2$) of $-6.2$ ($\pm 8$; sink) mmol m$^{-2}$ d$^{-1}$ was estimated for the entire study region, with the largest average CO$_2$ sink of $-10.0$ ($\pm 8$) mmol m$^{-2}$ d$^{-1}$ in the partly ice-covered Astrid Ridge (AR) region near the coast at 68°S and $-6.1$ ($\pm 8$) mmol m$^{-2}$d$^{-1}$ was observed in the Maud Rise (MR) region. A CO$_2$ sink was also observed south of 66°S in the Weddell Sea (WS). To assess the main drivers describing the variability of $fCO_{2}ssw$, a correlation model using $fCO_{2}$ and oxygen saturation was considered. Spatial distributions of the $fCO_{2}$ saturation/O$_2$ saturation correlations, described relative to the surface water properties of the controlling variables (chlorophyll a, apparent oxygen utilization (AOU), sea surface temperature, and sea surface salinity) further constrained the interplay of the processes driving the $fCO_{2}ssw$ distributions. Photosynthetic CO$_2$ drawdown significantly offsets the influence of the upwelling of CO$_2$-rich waters in the central Weddell gyre and enhanced the CO$_2$ sink in the region. FCO$_2$ of $-6.9$ mmol m$^{-2}$ d$^{-1}$ estimated for the Weddell gyre in this study was different from FCO$_2$ of $-2.5$ mmol m$^{-2}$ d$^{-1}$ in autumn estimated in a previous study. Due to low CO$_2$ data coverage during autumn, limited sea-air CO$_2$ flux estimates from direct sea-surface CO$_2$ observations particularly for the Weddell gyre region are available with which to compare the values estimated in this study. This highlights the importance of increasing seasonal CO$_2$ observations especially during autumn/winter to improving the seasonal coverage of flux estimates in the seasonal sea ice-covered regions of the Southern Ocean.

Keywords: CO$_2$ and biogeochemical drivers, chlorophyll, Antarctic coast, Maud Rise, Astrid Ridge, sea ice, oxygen saturation
INTRODUCTION

The Southern Ocean defined here as south of 40°S is well-known for its key role in the sequestration of CO$_2$ (Sabine et al., 2004; Takahashi et al., 2009) and accounts for about 43% of the global oceanic uptake of anthropogenic CO$_2$ (Frölicher et al., 2015). Colder surface water in the high latitudes enables it to absorb more CO$_2$ owing to increased solubility. Further, in the seasonal sea-ice zone, winter sea-ice cover reduces the CO$_2$ flux to the atmosphere, and during spring-summer when the sea-ice retreats; biologically driven CO$_2$ drawdown takes place (e.g., Hoppema et al., 1999; Bakker et al., 2008; Roden et al., 2016). The rapid uptake of CO$_2$ also takes place at regions of deep-water formation in the higher latitudes near the Weddell, Scotia, or Ross Seas and in the Mertz polynya (e.g., Bakker et al., 1997; Chierici et al., 2004; Fransson et al., 2004; Schmittner et al., 2007; Mattsdotter-Björk et al., 2014; Mu et al., 2014; Shadwick et al., 2014). CO$_2$ taken up from the atmosphere in the surface waters is sequestered into the deep ocean during deep-water formation and remained for a long time over several centuries (e.g., Kheshgi, 2004), climatically relevant. This deepwater is subsequently brought to the surface by upwelling; rich with CO$_2$ begins to equilibrate with the atmosphere driving ocean CO$_2$ outgassing but often counteracted by biological CO$_2$ drawdown (e.g., Fransson et al., 2004; Metzl et al., 2006; Gruber et al., 2009). Biological uptake of CO$_2$ in the surface Southern Ocean is enhanced at frontal structures as well in the Antarctic Circumpolar Current (ACC; e.g., Chierici et al., 2004; Ito et al., 2010) and the marginal sea ice zone (e.g., Froneman et al., 2004; Arrigo et al., 2008).

An annual mean uptake of $-0.42 \pm 0.07$ Pg C y$^{-1}$ was estimated for the region south of 44$^\circ$S between 1990 and 2009 from the integrated model and inversions method. This was consistent with the value calculated from surface ocean carbon dioxide observations ($-0.27 \pm 0.13$ Pg C y$^{-1}$) for the same period (Lenton et al., 2013). These values were also consistent with the contemporary mean uptake ($-0.34 \pm 0.02$ Pg C y$^{-1}$) determined from ocean inversions and $-0.30 \pm 0.17$ Pg C y$^{-1}$ from surface pCO$_2$ climatology (for the decade of 1990s to early 2000s) by Gruber et al. (2009) for the reference year 2000. More recently, new Southern ocean uptake of $-0.16 \pm 0.18$ was estimated in the south of 44$^\circ$S from mapped surface observations of combined float observations with shipboard data of 2015–2017 (Bushinsky et al., 2019). However, the area south of 58$^\circ$S shows different estimates of the flux over the models and observational results. For example, the model and inversions indicate a small annual sink of CO$_2$, whereas the observationally based estimate shows the area as a weak source of atmospheric CO$_2$ (Lenton et al., 2013). These discrepancies most likely are due to sparse observations since this part of the Southern Ocean (south of 58$^\circ$S) is poorly sampled (e.g., Monteiro et al., 2010). It could also be due to the limitation in time-space resolutions by model formulations, of the large seasonal variability in the various processes including temperature wind regimes, sea-ice conditions, and biological activity which govern atmosphere-ocean interactions (Takahashi et al., 2012). Thus, the current understanding of the seasonal drivers of sea surface CO$_2$ in the Southern Ocean is still limited, where more observations during different seasons are required to be used in models to accurately represent the seasonal cycle of CO$_2$ (Chierici et al., 2012; Lenton et al., 2013; Mongew et al., 2016). Further, the net annual CO$_2$ uptake, as well as the long-term trends in the seasonally ice-covered areas, are largely unknown (e.g., Takahashi et al., 2009; Long et al., 2013; Wanninkhof et al., 2013; Gregor et al., 2018). There is a need for a more comprehensive analysis of the individual regions and seasons (Hauck et al., 2010; Monteiro et al., 2015; McKinley et al., 2017). This study elucidates the drivers of variability for the sea surface CO$_2$ and sea-air CO$_2$ flux estimate in the South Atlantic Ocean with a focus on the Weddell gyre region and near the Antarctic coast given the importance of the region in the sequestration of atmospheric CO$_2$ and the climate system.

The Weddell Sea forms an important part of the Southern Ocean south of 58$^\circ$S because of the large cyclonic Weddell gyre which extends from the open ocean to the coastal region off the Antarctic continent such as the Dronning Maud Land (Figure 1). The Weddell gyre plays an important role in the CO$_2$ drawdown from the atmosphere (Hoppema et al., 1999; Bakker et al., 2008). Although, only weak annual CO$_2$ uptake occurs in the Weddell gyre (Hoppema et al., 1999; Brown et al., 2015) within the annual Southern Ocean sink of $-0.16$ to $0.34$ Pg C y$^{-1}$ (Gruber et al., 2009; Lenton et al., 2013; Bushinsky et al., 2019). A strong seasonal cycle exists in the sea surface CO$_2$ concentration and sea-air CO$_2$ flux (Vernet et al., 2019). High biological productivity (photosynthetic) during spring and summer in the Weddell gyre associated with seasonal sea ice edge dynamics (Smith and Barber, 2007) modulates the CO$_2$ variability with higher uptake (e.g., Hoppema et al., 1999; Vernet et al., 2019; Henley et al., 2020). Photosynthetic activity at Weddell and coastal polynyas (Arrigo et al., 2008; Cape et al., 2014) and the formation and export of deep water to the world's oceans (Grant et al., 2006; Brown et al., 2014) also contributes to the atmospheric CO$_2$ sequestration in this region. The surface water cooling that occurs during autumn leads to the uptake of CO$_2$ while in late autumn and during winter, there is the outgassing of CO$_2$ to the atmosphere (Brown et al., 2015). The deepening of the mixed layer depth associated with upwelling of CO$_2$-rich Circumpolar deep water (CDW) leads to increased surface ocean CO$_2$ concentration and results in the outgassing of CO$_2$ to the atmosphere which diminishes when the winter sea ice caps the surface ocean (Bakker et al., 2008; Brown et al., 2015).

This sea ice dynamic area is characterized by an abundant and persistent sea-ice cover that has extreme seasonal variability. The maximum extent of sea-ice cover occurs in September and by the following April, it would have shrunk to the third of the maximum extent (e.g., Vernet et al., 2019). Sea ice drift in dense pack ice and with icebergs are transported northward by the gyre. These melt in warmer waters and return freshwaters to the central gyre carrying micronutrients especially iron and stimulate primary productivity (Atkinson et al., 2001). The wind-driven, hydrodynamic circulation of the Weddell gyre connects the promontories (Astrid Ridge and Maud Rise) with the Weddell Sea on the west (Figure 1). The Weddell gyre makes the region the Weddell Sea deep water (WSDW) formation zone, as well
as a region of upwelling (as a result of the interaction of the rise with the Weddell gyre) of warm deep water (WDW) also known as CDW (Hoppema et al., 1995; Fahrbach, 2006; Figure 1). The upwelling occurs particularly during austral autumn and winter periods (Gordon and Huber, 1990) through Ekman pumping. The biological productivity also can occur during sea ice cover due to the lowering of the sea ice concentration by upwelling of the WDW resulting in an ice-free ‘hole’ at the Maud Rise (Weddell polynyas) and near the Antarctic continent (coastal polynyas; Arrigo et al., 2008; Bakker et al., 2008; Cape et al., 2014; Vernet et al., 2019), which allows light to reach the surface ocean and stimulate primary productivity (Arrigo and Van Dijken, 2003). The upwelling of WDW tends to cause supersaturation of CO$_2$ in the surface, offset by biological drawdown decreases the surface ocean inorganic carbon levels (Fransson et al., 2004; Brown et al., 2015). The interplay between upwelling and biological production sets the source or sink characteristics in this region. Large seasonal changes in the surface CO$_2$ concentrations are observed due to the intense photosynthesis in summer and upwelling of deep waters during winter (Takahashi et al., 2014). Relative to other regions of the Southern Ocean, there are fewer sea surface CO$_2$ observations in the eastern and southern boundaries of the Weddell gyre. This is because of their remoteness and extreme conditions, especially during autumn/winter (e.g., Takahashi et al., 2009; Vernet et al., 2019). Figure 2 presents the sea surface fCO$_2$ observations available in SOCAT v-2020 (Bakker et al., 2016, 2020) from 1999 to 2019 showing the data available in the eastern and southern boundaries of the Weddell gyre are mostly just for one to 2 years. Enhanced observations of the nuanced interplay between physical and biological processes, at varying spatial and temporal scales in this data-sparse region of the Southern Ocean are required. Estimating the CO$_2$ fluxes will also require a thorough understanding of the processes controlling spatial and temporal variations.

This study presents a unique dataset of surface water fCO$_2$, dissolved oxygen (DO), chlorophyll a, sea surface salinity, and temperature, and estimates on the air-sea CO$_2$ exchange characteristics in a large part of the Southern Ocean during austral autumn (February to April). The overarching aim of this study was to investigate the surface ocean property–property relationship to explain the physical and biological
drivers controlling CO$_2$ fluxes in an area and season where few observations exist (see Figure 2).

**MATERIALS AND METHODS**

**Data and Calculations**

The South Atlantic Southern Ocean was sampled between the 28th of February and the 10th of April 2019 onboard the Norwegian RV Kronprins Haakon. The study transect spanned from Punta Arenas (Chile) across ACC through the Weddell gyre to offshore of the Dronning Maud Land and Kong Håkon VII Hav at the Antarctic Coast. From there, heading northward along the 6°E meridian in the ACC toward Cape Town, South Africa (Figure 1; yellow transect) forms the eastern part of the transect. Underway continuous measurements were made for sea-surface and atmospheric CO$_2$ molar fractions (xCO$_2$) and ancillary parameters [sea-surface temperature (SST), sea-surface salinity (SSS), chlorophyll a fluorescence (chl$_a$) and DO]. Discrete seawater samples from Niskin bottles on a CTD rosette and from the ships’ water intake were also collected to supplement the dataset.

The dataset was divided into three sub-regions, categorized as the Weddell Sea (WS) region, Astrid Ridge (AR) region, and the Maud Rise (MR) region (Table 1 and Figure 1). This delineation was corresponded to the split along the transect (Figure 1) to ease the data analysis. The WS region span across part of the Weddell Sea in its southern extent and the ACC north of 60°S (ACCwest) in its northern extent (Figure 1). The AR region consists of the coastal waters near the Antarctic coast (66°S–68°S). Finally, the MR region spans the Weddell gyre in its southern extent to the north of 55°S in the ACC (ACCeast; Figure 1). The 60°S and 55°S are the northern boundary of the Weddell Gyre in the west and east, respectively (Deacon, 1979).

Using hydrographic data from this study; oceanic fronts in the ACC were identified by characteristic property indicators based on criteria adopted from earlier works (Deacon, 1982; Orsi et al., 1995; Pollard et al., 2002; Chierici et al., 2004; Mattsdotter-Björk et al., 2014; Freeman, 2017; Strass et al., 2017). Four major fronts were indicated along the study transect (Figure 1, black solid triangles). The ACC fronts as defined by Orsi et al. (1995) are overlaid on the map (Figure 1, gray lines) with the study transect.

**Fugacity of Carbon Dioxide**

Sea surface and atmospheric CO$_2$ molar fractions (xCO$_2$) were measured onboard by an autonomous underway partial pressure of CO$_2$ (pCO$_2$) observation system (General Oceanics®, Inc., model 8050) which consists of a gas-water equilibrating chamber and an infrared analyzer (LICOR®, Model, 7000). The seawater was pumped from a side intake at 4 m below the sea surface, sprayed through the equilibration chamber; to equilibrate the CO$_2$ in the seawater with the air in the headspace of the chamber, and measured by the infrared analyzer, with an accuracy of ± 0.2 ppm. The analyzer was calibrated every 2.5 h using three standard gases in synthetic air with CO$_2$ molar fractions of 230, 400, and 550 ppm. The accuracy of the measurements by the General Oceanics system was estimated using secondary standards calibrated toward NOAA gases (traceable to WMO-x93 scale). Between calibrations, continuous measurements were made every third minute in a sequence of xCO$_2$ of standard gases, of air, and the seawater. The sea surface water and atmospheric

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**TABLE 1 | Cruise transect and the three sub-regions defined with their coordinates.**

| Transsects            | Latitude Range | Longitude Range | Period of Occupation |
|-----------------------|----------------|-----------------|---------------------|
| Entire cruise         | 68°S–43°S      | 62°W–13°E       | 1 March – 10 April, 2019 |
| Weddell Sea region (WS) | 55°S–67°S      | 62°W–22°W       | 1 – 7 March, 2019   |
| Astrid Ridge region (AR) | 67°S–68°S      | 10°W–12°E       | 10 – 27 March, 2019 |
| Maud Rise region (MR)     | 65°S–43°S      | 2°E–13°E        | 29 March – 7 April, 2019 |
fugacity of CO₂ (fCO₂ssw and fCO₂atm) were computed from the xCO₂ through pCO₂ corrected for non-ideal behavior using SST and SSS following the methods of Pierrot et al. (2009) for fCO₂ssw and with Lencina-Avila et al. (2016) for fCO₂atm. For an uncertainty in the measured standards of less than 1 ppm (reflecting the standard deviation of the difference between measured standards and certified values), with an accuracy in the equilibrator temperature of 0.01°C, (0.009°C at 0°C water temperature) and at the intake temperature of 0.001°C, the determined fCO₂ will be within 2 µatm as previously stated by Pierrot et al. (2009). This assumes that the pressure is determined within 0.2 hPa.

**Sea-Air Carbon Dioxide Flux Calculations**

Sea-air CO₂ flux (FCO₂) was calculated using equation (1):

\[
F_{CO_2} = k \times s \times \Delta f_{CO_2}
\]

where

\[
\Delta f_{CO_2} = f_{CO_2} \text{ssw} - f_{CO_2} \text{atm}
\]

k is the CO₂ gas transfer velocity coefficient (cm hr⁻¹) estimated using the formulation of Wanninkhof (2014) with the coefficient factor of 0.251 and a quadratic wind speed. Co-location wind speed recorded at 36 m height from the ship’s scientific data and corrected to a 10 m standard height was utilized.

s is the CO₂ solubility coefficient (mol L⁻¹ atm⁻¹) calculated following the equation and the coefficients for temperature and salinity dependence of the solubility of CO₂ in Weiss (1974). ΔfCO₂ equals the sea-air fCO₂ gradient in µatm. The recalculated fugacity of CO₂ from the continuous underway measured mole fractions for the atmospheric CO₂ (Supplementary Figures 1A,B), was used in equation (2) for fCO₂atm in the calculations for all the regions. The flux values are reported in mmol m⁻² d⁻¹ using a conversion factor of 0.24, taking into account the µatm, the conversion of cm h⁻¹ to m d⁻¹ and mol L⁻¹ atm to mol m⁻³ atm.

Negative values of ΔfCO₂ represent the undersaturation of fCO₂ssw with respect to the atmospheric fCO₂ (fCO₂atm) and positive values represent supersaturation. The sea surface water is saturated with CO₂ when ΔfCO₂ is 0. Similarly, negative values of fCO₂ quantify the flux of CO₂ into the ocean while positives values indicate the degassing of CO₂ to the atmosphere. At equilibrium, fCO₂ is 0.

The fugacity of CO₂ saturation (fCO₂sat) was calculated using the following equation:

\[
f_{CO_2} \text{sat} = \frac{f_{CO_2} \text{ssw}}{f_{CO_2} \text{atm}} \times 100
\]

**Sea Surface Temperature and Salinity**

Continuous SST and SSS measurements were obtained by a thermostalinograph (SeaBird SBE21 TSG) and an additional temperature sensor (SBE38) at the seawater intake at the bottom of the ship, with an accuracy of ± 0.001°C. The accuracy of the salinity sensor of ± 0.02 psu was obtained by comparison with salinity measured using an Autosol® on discrete water samples collected from the intake and the drift corrections obtained from calibrations of the sensor.

**Dissolved Oxygen, Oxygen Saturation, and Apparent Oxygen Utilization**

Dissolved oxygen concentration ([O₂], µmol kg⁻¹) was measured with the Aandera optode (model 4330, the accuracy of < ± 2%) sensor attached to the seawater intake, corrected for SSS. The optode DO values were evaluated by Winkler titration of water samples collected at 5–10 m depth from Niskin-CTD rosette and water intake, resulting in uncertainty of 2 ± 1%. Percentage oxygen saturation (O₂sat) and apparent oxygen utilization (AOU) were estimated using Eqs (4) and (5; Garcia et al., 2013), respectively.

\[
O_2 \text{ sat} = \frac{[O_2]}{[O_2]_1} \times 100
\]

\[
\text{AOU} = [O'_2] - [O_2]
\]

[O₂] is the O₂ solubility concentration (µmol kg⁻¹) calculated as a function of *in situ* temperature, salinity, and one atmosphere of the total pressure. The values of [O₂] were calculated according to Garcia and Gordon (1992) based on the values of Benson and Krause (1984).

The AOU of a water sample is the difference between the concentration at oxygen saturation and the measured oxygen concentration in the water with the same physical and chemical properties. AOU is commonly used to investigate the sum of the biological activity that the sample has experienced since it was last in equilibrium with the atmosphere. At 100% O₂sat, AOU is 0. High oxygen utilization results in O₂sat < 100 and AOU > 0, while low oxygen utilization results in O₂sat > 100 and AOU < 0. Thus, positive AOU will depict the respiration/mineralization process and negative AOU will depict photosynthesis.

**Chlorophyll a**

Underway chlorophyll a fluorescence (chlfluor) was measured with the Wetstar fluorometer every minute. The fluorometer was calibrated against a total of 109 chl-a measurements (chlextr) obtained from discrete samples, which were collected and analyzed after extraction using the acetone-spectrofluorimetric method (Holm-Hansen and Riemann, 1978). Supplementary Figure 2A shows the diurnal (day and night) fluorescence derived chl-a which does not show a significant difference in the slope and intercept hence the combined data (Supplementary Figure 2B) was used for the calibration. Therefore, the linear correlation obtained between the combined diurnal Wetstar sensor fluorescence and the extracted chl a, (r² = 0.85, N = 109) was used to convert the voltage signal (mV) to *in situ* chl-a:

\[
\text{chl} - a = (0.09 \times \text{fluorescence}) + 0.06
\]

**RESULTS**

**Physical Properties and Hydrography**

Temperature ranges within the WS, AR, and MR regions were −0.9 to 6.19°C, −1.9–0.4°C, and −0.1–14.0°C, respectively (Table 2). SSS values were in the range of 33.26–34.46 psu, 33.45–34.25 psu, and 33.74–35.10 psu for the respective regions.
(Table 2), SST varied widely over the WS and MR regions with mean values and standard deviations of 1.1 (± 2.0)°C and 1.9 (± 3.1)°C, respectively, indicating the difference between warmer characteristics of ACC waters and the colder and more stratified water in the Weddell gyre. In the AR, temperature showed relatively small variability, the mean SST was -0.7 (± 0.7)°C (Table 2).

The frontal systems described for the studied region are identifiable from the sharp gradient created by the most rapid changes in SST and SSS. For instance, while going southwards across ACCwest in the WS region, at 57.5°S, the SST and SSS rapidly decreased from 6 to 3°C and from 34.1 to 33.75 psu, respectively (Figure 3A), indicating the location of the Antarctic Polar Front (APF) on the western flank of the cruise track. A decrease in SST to about 1°C with an increase in SSS from 33.8 psu to 34.0 psu at around 59°S appears to be an expression of Southern ACC Front (Figure 3A, SACCF). Moving northward from AR on the eastern flank of the transect, SACCF was also identified at 53.25°S by the SST increase from below 1 to about 4°C with the corresponding decrease in the SSS from 34.0 psu to 33.8 psu (Figure 3C, SACCF) in the MR region. A weak northward increase of SST from below 4°C to about 5°C between 51 and 50°S shows an expression of APF (Figure 3C) and a northward increase of SST from 5 to 8°C with an increase in SSS from 33.70 to 33.80 psu indicating the Sub-Antarctic Front at 48.2°S (Figure 3C, SAF). The large sharp increase of SST and SSS from 8.5 to 14°C and from 33.9 to 35.10 psu, respectively, with warmer (greater than 11.5°C) and saltier (greater than 34.9 psu) waters on the northern side indicated the Subtropical Front at 45.28°S (Figure 3C, STF). STF was further south than the other fronts indicated for this study compared with the Orsi et al. (1995) frontal positions along the transect.

**Spatial Distribution of fCO$_2$ssw, With the Physical and Biological Parameters**

**WS Region**
The fCO$_2$ssw distributions in the WS region ranged from 315 to 455 µatm (Table 2). The WS region showed the largest spatial variability of fCO$_2$ssw relative to other regions (Figure 3D) and included, the highest fCO$_2$ssw of 455 µatm recorded on the cruise in the WS region south of the SAACF, between 60 and 62°S, accompanied by a large increase in SSS, a small decrease in the SST and a small increase in chl-a (Figures 3A,D,G). Increasing fCO$_2$ssw toward saturation relative to the mean fCO$_2$atm also corresponded with increasing SSS in the SACCF (Figures 3A,D). The highest chl-a concentration (0.4 mg m$^{-3}$) in this region was found between 58 and 59°S (Antarctic zone; Figure 3G). South of 66°S the fCO$_2$ssw drastically decreased to the minimum value of 315 µatm, which coincided with a peak in chl-a of about 0.35 mg m$^{-3}$ (Figures 3D,G) and low temperatures. However, DO concentration variation only corresponded partly to the variability in the chl-a concentration. This was particularly clear in the APF and south of 66°S where no correlation between chl-a and DO was observed (Figures 3G,J). Opposite variation of AOU with DO saturation was observed as expected (Figures 3G,J), whereas there are deviations in the co-variation between the DO and oxygen saturation in the APF and SACCF region (Figure 3F).

**AR Region**
Along coastal longitudes of the AR region (Figure 1, Astrid Ridge region), all parameters showed less variability, except for
FIGURE 3 | Spatial distributions of sea surface fugacity of CO$_2$ with physical and biological parameters. Column 1: (A,D,G,J) WS region, Column 2: (B,E,H,K) AR region, and Column 3: (C,F,I,L) MR region. The red color represents the distributions of the atmospheric fugacity of CO$_2$ in each of the defined regions.

chl-a which showed several peaks (Figure 3H) with an uneven fCO$_2$/chl-a relationship for photosynthesis (Figures 3E,H). For example, a chl-a concentration of 1.1 mg m$^{-3}$ coincided with a fCO$_2$ssw value of about 320 µatm as well as chl-a concentrations between 0.2 and 0.6 mg m$^{-3}$ as observed at 5°W and 2°W. Moreover, some values of fCO$_2$ssw were lower than 320 µatm and coinciding with chl-a concentrations of about 0.2–0.4 mg m$^{-3}$ at 7°E (Figures 3E,H). Low values of fCO$_2$ssw, SST, and SSS with high values of chl-a were recorded in this region (Figures 3B,E,H). AOU also showed the opposite variation to DO saturation (Figures 3H,K) and DO and oxygen saturation generally co-varied in this region (Figure 3K).

MR Region
In the MR region, fCO$_2$ssw showed large variability as indicated by the standard deviation of ± 23 µatm (Table 2). The lowest fCO$_2$ssw values in the south coincided with high chl-a and the lowest SST (Figures 3C,I). The fCO$_2$ssw increased northward and reached oversaturation at about 55°S, coinciding with increased SST and fCO$_2$ssw variability that coincided partly with the fronts (Figures 3C,F,I). The northward warming at the fronts corresponded to increasing fCO$_2$ssw, leading to fCO$_2$ oversaturation with an exception at the STF where warming corresponded to a decrease in fCO$_2$ssw and increased chl-a. AOU showed opposite variation to DO saturation (Figures 3I,L) and
DO and oxygen saturation co-varied except for north of 54°S in the frontal waters (Figure 3I).

**Spatial Distribution of FCO$_2$, ΔfCO$_2$, and Wind Speed**

The spatial distribution of the sea-air CO$_2$ flux (FCO$_2$, mmol m$^{-2}$ d$^{-1}$), ΔfCO$_2$ (µatm), and the co-location wind speed (m s$^{-1}$) from the ship data corrected to 10 m standard height, for the three defined regions are presented in Figure 4. Negative FCO$_2$ denotes a CO$_2$ flux into the ocean (CO$_2$ influx; sink) and positive values denote CO$_2$ flux out of the ocean to the atmosphere (CO$_2$ outgassing; source). The FCO$_2$ spatial distribution along the entire cruise varied from near zero wind speed of 0.1 m s$^{-1}$ to a maximum wind speed of 25.4 m s$^{-1}$ (Table 2) with mean values for each region being 8.5 (± 2.3) m s$^{-1}$, 9 (± 4) m s$^{-1}$ and 10 (± 5) m s$^{-1}$ (Table 2) for WS, AR, and MR, respectively. Despite occasions of CO$_2$ outgassing along the transect in the WS and MR region (Figures 4A,C), both regions showed an average ocean CO$_2$ uptake of −1.5 mmol m$^{-2}$ d$^{-1}$ and −6.1 mmol m$^{-2}$ d$^{-1}$ (Table 2) respectively and as well as the AR region where CO$_2$ influx was observed throughout the region (Figure 4B). The AR region showed the largest average CO$_2$ influx for atmospheric CO$_2$ with an average flux estimate of −10.0 mmol m$^{-2}$ d$^{-1}$ (Table 2).

In the WS region, FCO$_2$ showed a similar spatial variation pattern with ΔfCO$_2$ (Figure 4A). The large CO$_2$ outgassing in the WS coincided with the highest positive ΔfCO$_2$, with little change in the wind speed (Figures 4A,D). However, the largest undersaturation (largest negative ΔfCO$_2$) in the southernmost part at 67°S did not result in the expected higher CO$_2$ influx, a consequence of the relatively low wind speed (Figure 4D).

The AR region showed uptake of CO$_2$, which generally increased with increasing wind speed (Figures 4B,E). Also, here between 3°W and 1°E, the CO$_2$ influx did not correspond with the large undersaturation of fCO$_2$ssw (large negative ΔfCO$_2$, Figure 4B) where the CO$_2$ influx was nearly zero regardless of the large undersaturation of fCO$_2$ssw of about −60 µatm (Figure 4B). This also corresponded to the relatively low wind speed values recorded along this longitude (Figure 4E).

Moving northward in the MR region to 60°S, increasing wind speed as well as the magnitude of undersaturation of fCO$_2$ssw (large negative ΔfCO$_2$) resulted in larger CO$_2$ flux. However, between 60 and 57°S, the FCO$_2$ showed little change although both ΔfCO$_2$ and wind speed increased (Figures 4C,F). North of 57°S, FCO$_2$ showed a similar spatial variation pattern with ΔfCO$_2$ and higher wind speeds seemed to favor CO$_2$ outgassing between 52 and 48°S (Figures 4C,F). The highest wind speed of 25.4 m s$^{-1}$ recorded on the cruise was found along the latitudes of 52°S and around 65°S (Figure 4F). An increasing CO$_2$ influx with larger negative gradients of ΔfCO$_2$ and increasing wind speed was observed north of STF in the subtropical waters (Figures 4C,F).

**DISCUSSION**

This study presents a recent sea-air CO$_2$ flux estimate from direct CO$_2$ observations in the Atlantic sector of the Southern Ocean and within the Weddell gyre region. From the observations performed in autumn, the whole region along the study transect acted as an average ocean CO$_2$ sink for atmospheric CO$_2$ with an average FCO$_2$ of −6.2 (± 8) mmol m$^{-2}$ d$^{-1}$ (Table 2). The Weddell gyre region showed a strong uptake of atmospheric CO$_2$ at the Maud Rise feature and near the Antarctic coast at the Astrid Ridge region and strong outgassing around 60°S. Integrating the CO$_2$ uptake for the Weddell gyre region, a
The biological CO$_2$ drawdown continues into autumn with ongoing primary production at the late stage (Brown et al., 2015). This could explain the high chl-a concentrations within the AR and MR regions for this study (Section “Spatial Distribution of fCO$_2$ssw, With the Physical and Biological Parameters”). Also, the autumn bloom was observed in the upper water column west of the Astrid Ridge during the cruise for this study (Kauko et al., in review).

The WS region acted both as a weak sink and a weak source with an average fCO$_2$ of $-1.5 \pm 3$ mmol m$^{-2}$ d$^{-1}$. The large variability was associated with the large fCO$_2$ssw supersaturation, fCO$_2$ outgassing between 60 and 62$^\circ$S, and the low CO$_2$ influx found in the SACCf (Figure 4A). This may result from the upwelling in the region indicated by an increase in the SSS (Figure 3A). Upwelled deep waters are older and relatively rich in CO$_2$ from the remineralization of organic carbon in the deep (Gordon and Huber, 1990; Hoppema et al., 2000; Fransson et al., 2004). Once at the surface, they begin to equilibrate with the atmosphere driving CO$_2$ outgassing (Metzl et al., 2006; Gruber et al., 2009). Interestingly, the high fCO$_2$ssw is also seen at the same latitude north of the Peninsula in the SOCAT v-2020 for the number of available fCO$_2$ observations (Supplementary Figures 4A,B). This is likely a permanent local event as the area is known for intense upward mixing of cold deep waters (Heywood et al., 2002, 2004) which may be linked with the bathymetry interaction with the high rising South Scotia Ridge in the Scotia Sea in this region. The large CO$_2$ influx observed south of 67$^\circ$S (Figure 4A) can be attributed to upwelling-induced primary production. This is because the large undersaturation of fCO$_2$ssw relative to the atmospheric CO$_2$ in Figure 3D corresponded to an increase in SST and high SSS (upwelling) (Figure 3A), and increased chl-a (primary production; Figure 3G). The course of the WDW from the western part of the Weddell gyre (Figure 1), supports the upwelling of WDW (increase in SST and high SSS) in the region.

The largest average regional CO$_2$ sink of $-10.0$ mmol m$^{-2}$ d$^{-1}$ (Table 2) was estimated for the AR region in the far south along the Antarctic coast. The sink was associated with the high chl-a concentrations recorded in this region (Figure 3H) which likely caused the undersaturation of fCO$_2$ssw (Figure 4B; ΔfCO$_2$). Therefore, the high ocean uptake of CO$_2$ could be attributed to primary production in combination with the effect of wind speed; since CO$_2$ influx also increased once the wind speed increases as much as to 10$^8$E (Figures 4B,E). Maud Rise region was a stronger average CO$_2$ sink than the WS region and has an average fCO$_2$ of $-6.1 \pm 8$ mmol m$^{-2}$ d$^{-1}$ (Table 2). The MR region spans from the Weddell gyre at the Maud Rise to the ACC and subtropical waters north of the STF (Figure 1). The net sink in this region was greatly influenced by the large CO$_2$ uptake at the Maud Rise and in the subtropical waters (Figure 4C). The Maud Rise has been identified by many other authors as an area of high primary productivity which results in a large biological CO$_2$ drawdown. Formation and upwelling of WDW (Gordon and Huber, 1990; Hoppema et al., 1999), interacting with the Maud Rise brings nutrients and CO$_2$ to the surface and drives productivity
(Holm-Hansen et al., 2005). Indeed, the observed increase in SSS and SST (upwelling of WDW) at the Maud Rise latitude (65°S; Figure 3C) and the corresponding increase in the chl-a concentration (Figure 3I) supports the observed net CO₂ sink. The CO₂ outgassing in the ACC component of the MR region is associated with high wind speeds at the fronts and enhanced deep upwelling, driven by the strong westerly winds (e.g., Fransson et al., 2004; Brown et al., 2015). At the most northern extent of the MR region (north of the STF; Figure 3, column 3), the low values of DO, chl-a, and CO₂ssw are characteristics of oligotrophic subtropical waters. Enhanced transport of dissolved organic carbon from the surface to the deep ocean has previously been observed in the oligotrophic subtropical oceans (Roshan and DeVries, 2017). This could potentially translate to the CO₂ssw observed in the region (Figure 3F) and which results in the large undersaturation of CO₂ssw (ΔfCO₂) and subsequent high fCO₂ (Figure 4C). Hoppema et al. (2000) also recorded undersaturated surface water for CO₂ and observed that the surface water in the region tends to flow northwards to participate in the formation of Antarctic Intermediate Water (AAIW) and so constitute a conduit for CO₂ uptake from the atmosphere. Other sea-surface fCO₂ observations in this region during April/May of the same year (2019) as this study also show low fCO₂ for the same region near the STF and similar fCO₂ distributions overall for the region of study (Supplementary Figure 5). This corroborates the autumn distribution of sea surface fCO₂ in the region for 2019 as presented in this study.

Regional Drivers of fCO₂ssw Using a Correlation Model

Dissolved O₂ can help to constrain understanding of the drivers of surface ocean carbon dynamics. To explain the nuanced interplay of the various drivers influencing the CO₂ssw variability, a correlation model between the saturation of fCO₂ (fCO₂sat) and O₂ saturation (O₂sat) relative to the atmosphere as utilized by Carrillo et al. (2004) was explored (Figure 5). In the model, the calculated fCO₂sat and O₂sat (see Sections Sea-Air Carbon Dioxide Flux Calculations and Dissolved Oxygen, Oxygen Saturation, and Apparent Oxygen Utilization) were correlated and used as an index to derive the controlling or dominant chemical, physical and biological processes influencing fCO₂ssw variability.

The correlative exercise segregated the fCO₂ and O₂ data into four quadrants (Carrillo et al., 2004; Moreau et al., 2013) depicting four case waters (Figure 5). The distribution of observations in each quadrant helps infer the dominating processes controlling fCO₂ssw distribution in surface waters. Quadrant I depicted surface waters of simultaneous fCO₂ undersaturation (below 100% saturation level) and O₂ supersaturation (above 100% saturation level), implying photosynthesis dominantly driving the spatial distribution of CO₂ssw. In Quadrant II, supersaturation of both O₂ and fCO₂ suggests warming of the surface waters as the dominant process increasing the saturation of both fCO₂ and O₂. Quadrant III depicted surface waters experiencing simultaneous fCO₂ supersaturation and O₂ undersaturation and implies dominant respiration/remineralization or upwelling of respired subsurface waters. Lastly, quadrant IV depicted surface waters of simultaneous fCO₂ and O₂ undersaturation which implies the cooling effect of temperature dominantly driving both fCO₂ and O₂ saturation in the surface water lower with respect to gas exchange. Consequently, if biological processes (primary production and respiration/remineralization) and upwelling are the dominant drivers controlling fCO₂ssw and O₂ saturation, a negative correlation will be expected with the O₂sat and fCO₂sat values distributed linearly through quadrant I and III in Figure 5. Also, if temperature-driven processes (warming and
cooling) are the dominant control factor for fCO$_2$ssw and O$_2$ saturation, a positive correlation between O$_2$sat and fCO$_2$sat with the values distributed linearly through quadrant II and IV will be expected in Figure 5. The process vectors (correlation lines) in Figure 5 show deviations in correlative relationships, potentially indicating a more complex interplay between the processes driving the observed distribution of fCO$_2$sat and O$_2$sat. This is illustrated in Sections “Surface Water Property–Property Relationship With fCO$_2$sat/O$_2$sat Correlations for Case Waters QI and QIII” and “Surface Water Property–Property Relationship With fCO$_2$sat/O$_2$sat Correlations for Case Waters QII and QIV” as the spatial distributions of the fCO$_2$sat/O$_2$sat for the quadrants are described relative to the surface water properties of controlling variables (chl-a, AOU, SST). For example, increasing chl-a, a proxy for primary production, results in the consumption of CO$_2$ with the release of oxygen in the surface waters (Sigman and Hain, 2012) affecting their respective saturation levels. Negative values of AOU indicating low oxygen utilization corresponds to the photosynthetic process, while positive values of AOU indicate respiration/remineralization and upwelling. Likewise, an increase in temperature produces an increase in both the O$_2$ and fCO$_2$ saturation state. SSS is used as an index for upwelling since upwelled waters are more saline. Deviations in the correlative relationships could also be due to the difference in sea-air gas exchange rates for O$_2$ and fCO$_2$ (Broecker and Peng, 1983) or due to the formation and dissolution of calcium carbonate (Dieckmann et al., 2008). Since fCO$_2$ssw is a component of the ocean’s buffer system, the sea-air CO$_2$ exchange has a slower response than the case for oxygen with timescales ranging from days to weeks for O$_2$ and months for fCO$_2$ [details of the sea-air exchange model for O$_2$sat and fCO$_2$sat in Carrillo et al. (2004)]. Thus, the processes of differential sea-air gas exchange will affect the O$_2$sat/fCO$_2$sat ratio. The case waters depicted by the quadrants are hereafter referred to as QI, QII, QIII, and QIV.

Surface Water Property–Property Relationship With fCO$_2$sat/O$_2$sat Correlations for Case Waters QI and QIII

The case waters QI representing photosynthesis were found only in parts of the WS region; in the ACCwest on its northern extent and a small section on the southern extent in the Weddell gyre.
The effect of photosynthesis on the \( f\text{CO}_2 \text{sw} \) distribution in this region confirms previous studies within the same region especially in the east of Drake Passage (Munro et al., 2015). QIII case waters representing respiration/remineralization and upwelling were found predominantly in the MR region (in the ACCeast) and some part of the WS region (in the Weddell gyre; Figure 6, deep blue). The \( f\text{CO}_2 \text{sat}/\text{O}_2 \text{sat} \) correlation for the QI and QIII in the WS region showed a significant negative correlation of \(-0.84, p < 0.001 \) with a slope of \(-0.78 \) (Figures 5A, I, III). This strong association between \( f\text{CO}_2 \text{sat} \) and \( \text{O}_2 \text{sat} \) indicated the combined influence of the biological and upwelling processes driving the observed \( f\text{CO}_2 \text{sat}/\text{O}_2 \text{sat} \) variability. Figure 7A presents the \( f\text{CO}_2 \text{sat}/\text{O}_2 \text{sat} \) variability for the case waters in QI and QIII. North of 60\(^\circ\)S in the ACC and between 61 and 62\(^\circ\)S in the Weddell gyre (Figure 3G), the variability in the chl-a concentrations, and the AOU (negative AOU) highlighted the photosynthetic relationship of \( f\text{CO}_2 \text{sat}/\text{O}_2 \text{sat} \) in the same region (Figure 7A). Between 60 and 61\(^\circ\)S and South of 63\(^\circ\)S along the same transect, upwelling of high saline (Figure 3A) and \( \text{CO}_2 \)-rich waters (Figure 3D) with respiration/remineralization (positive AOU, Figure 3G) were evident from the corresponding supersaturation of \( f\text{CO}_2 \text{sat} \) and the undersaturation of \( \text{O}_2 \text{sat} \) (Figure 7A). The positive values of AOU and increase in SSS in this region, support the respiration/remineralization and upwelling processes driving the observed \( f\text{CO}_2 \text{sat}/\text{O}_2 \text{sat} \) here. The transition to supersaturation and a local maximum of \( f\text{CO}_2 \text{sw} \) between 60 and 61\(^\circ\)S along the WS transect also previously observed near the same region (Stoll et al., 1999; Hoppema, 2004) was attributed to possible \textit{in situ} remineralizations of organic matter. Besides, the area is known for intense upward mixing of cold deep waters (Heywood et al., 2002, 2004). Southward of 63\(^\circ\)S, the observed values of chl-a concentration (Figure 3G) suggest an interplay of photosynthesis with the upwelling process shown for the region by the \( f\text{CO}_2 \text{sat}/\text{O}_2 \text{sat} \) distributions (Figure 7A). It is thought that photosynthesis offsets the \( f\text{CO}_2 \text{sat} \) while enhancing the \( \text{O}_2 \text{sat} \) and was evident in the slight supersaturation of \( f\text{CO}_2 \text{sat} \) and the near-saturation level (close to 100% saturation) of \( \text{O}_2 \text{sat} \) (Figure 7A) in the region. In the ACC, north of the APF the SST and SSS show a similar variation pattern (Figure 3A) to the \( f\text{CO}_2 \text{sat} \) (Figure 7A) while at the APF both \( f\text{CO}_2 \text{sat} \) and \( \text{O}_2 \text{sat} \) decreases with decrease in SST and SSS (Figures 3A, 7A).

This also shows the interplay of temperature with photosynthesis in QI case waters.

For the QIII case waters in the MR region found in the ACCeast (the northern extent) in the eastern sector of the study transect (Figure 6, deep blue), the correlation value is \(-0.27, p < 0.001 \) (Figures 5C, III). The \( f\text{CO}_2 \text{sat}/\text{O}_2 \text{sat} \) variability for the case waters in QIII along the section of the MR is shown in Figure 7B. Although the positive AOU values in Figure 3I along this section generally indicates the respiration/remineralization process derived for the QIII case waters as shown by the \( f\text{CO}_2 \text{sat}/\text{O}_2 \text{sat} \) distribution (Figure 7B), the correlation values indicate no clear correlation between \( f\text{CO}_2 \text{sat} \) and \( \text{O}_2 \text{sat} \) for the process in this region. South of 54\(^\circ\)S, the higher SST (Figure 3C) with increasing \( \text{O}_2 \text{sat} \) (Figure 7B) is indicative of the interplay of temperature with the respiration/remineralization process. Moreover, this effect of temperature is not observed correspondingly on the \( f\text{CO}_2 \text{sat} \) as the supersaturated \( f\text{CO}_2 \text{sat} \) was slightly above saturation level and almost constant while the undersaturated \( \text{O}_2 \text{sat} \) was increasing further north (Figure 7B). It is therefore deduced that temperature and photosynthesis (seen in the variation of chl-a along the section (Figure 3I), as well as respiration/remineralization processes, combine to drive the observed \( f\text{CO}_2 \text{sat}/\text{O}_2 \text{sat} \) variability in QIII for this region.

The overall mechanism above shows that the undersaturation and supersaturation of \( f\text{CO}_2 \text{sat} \) each in QI and QIII case waters were generally driven by photosynthesis and respiration/remineralization and upwelling, respectively as derived for the quadrants. However, along the ACCeast on the MR section, the influence of temperature on the spatial distributions of \( f\text{CO}_2 \text{sat}/\text{O}_2 \text{sat} \) affected the \( f\text{CO}_2 \text{sat}/\text{O}_2 \text{sat} \) correlation in the ACC.

**Surface Water Property–Property Relationship With \( f\text{CO}_2 \text{sat}/\text{O}_2 \text{sat} \) Correlations for Case Waters QII and QIV**

The dominating process derived for the case waters QII (Figure 6, red) and QIV (Figure 6, sky blue) is the effect of temperature on the variability of \( f\text{CO}_2 \text{sat}/\text{O}_2 \text{sat} \). These case waters (QII and QIV) were found in parts of the WS region while the waters of AR were completely distributed in the QIV. Waters along the southern extent and some parts of the ACCeast in the MR region were also observed as QIV case waters (Figure 6, sky blue).

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**Figure 7** | Spatial distribution of the \( f\text{CO}_2 \text{sat}/\text{O}_2 \text{sat} \) correlations for the case waters in QI and QIII. (**A**) WS region and (**B**) MR region.
The sections of the WS transect distributed in QIV were located south of 60°S in the Weddell gyre region (Figure 6, sky blue), with a considerably smaller section between 58 and 60°S (in the ACCwest) in the QII (Figure 6, red). A significant but weakly positive correlation of 0.33, \( p < 0.001 \) between fCO\(_2\)sat and O\(_2\)sat with a slope of 0.1 (Figures 5A,II,IV) was observed, indicating a weak effect of temperature on the fCO\(_2\)sat/O\(_2\)sat relationship. The low fCO\(_2\)sat/O\(_2\)sat correlation could be attributed to the near-saturation level of the supposed undersaturated fCO\(_2\) and O\(_2\) for most of the observations (Figure 8A), and the process vector (green regression line, Figure 5A) not passing through QII. As very few observations are in QII, this could indicate that the influence of temperature on the supersaturation of fCO\(_2\)sat and O\(_2\)sat in the QII is small. Figure 8A shows fCO\(_2\)sat/O\(_2\)sat variability for the case waters in QIV and the few observations for QII along the WS region. The cooling effect of temperature is expected as the dominant driver in QIV. However, the upwelling of rich-CO\(_2\) indicated by the higher SSS values between 61 and 64°S and at 67°S (Figure 3A) potentially influence the fCO\(_2\)sat in Figure 8A and enhance fCO\(_2\) to near-saturation level. Similarly, the near saturation of O\(_2\)sat (Figure 8A) could also potentially be the influence of photosynthesis along the same section (Figure 3G) enhancing the saturation of O\(_2\). This is particularly clear around 67°S where O\(_2\)sat was enhanced with fCO\(_2\)sat greatly reduced (Figure 8A). Furthermore, the positive values of AOU, albeit low (< 20 µmol kg\(^{-1}\); Figure 3G), indicate the influence of respiration/remineralization. Therefore the interplay between upwelling and respiration/remineralization and primary production (photosynthesis) enhanced the saturations of fCO\(_2\)sat and O\(_2\)sat for most of the observation for QIV in the WS. This indicates a more complex dynamic in the processes driving the observed fCO\(_2\)sat/O\(_2\)sat variability than the simplified representation of the cooling effect of temperature for the observations distributed in QIV.

The entire distribution of observations in the AR region are in the QIV (Figure 6, sky blue). The intricate interplay of different processes driving the relationship of fCO\(_2\)sat and O\(_2\)sat in the QIV is also observed in the AR region. An unexpectedly weak negative correlation of −0.19 (\( p < 0.001 \)) was estimated (Figure 5B) instead of the positive correlation expected for the fCO\(_2\)sat/O\(_2\)sat relationship in QIV. Visualizing the distribution of the observations in the quadrant indicates two data populations: one, a positive correlation for fCO\(_2\)sat and O\(_2\)sat on the lower part and secondly, a negative correlation for the upper part of the distribution (Figure 5B). Figure 8B, shows the variability of fCO\(_2\)sat/O\(_2\)sat for the case waters in QIV along the AR transect. Longitudinal distributions of each of the two visualized groups (Figure 9) reveals the effect of temperature (the positive correlation) on the fCO\(_2\)sat/O\(_2\)sat at the east of 9°E, between 5 and 8°E, and at 3°E for the first group (Figures 9A,C). This is observed along the sections as both fCO\(_2\)sat and O\(_2\)sat decrease (Figure 8B) with decreasing SST and SSS (Figure 3B) which becomes significant at the east of 9°E. This could indicate cooling and freshening of the surface waters from glacial and sea-ice melt near the continent which decreases the fCO\(_2\) and O\(_2\) saturations (Klinck, 1998; Ohshima et al., 1998 Dierssen et al., 2002; Carrillo et al., 2004). On the other hand, the longitudinal extent of the second data population distribution shows the effect of photosynthesis (negative correlation) across the whole region west of 10°E (Figure 9B,D). This corresponds with the variable chl-a concentrations associated with the fCO\(_2\)/chl-a relationship for photosynthesis along the region (see section “AR region”) and Figure 3H. The variable photosynthetic process could potentially be related to the patchy plankton blooms characteristics of Antarctic coastal waters (Carrillo et al., 2004). Furthermore, the increasing positive values of AOU along the region in the east of 9°E (Figure 3H) indicate also the influence of respiration/remineralization on the fCO\(_2\)sat/O\(_2\)sat relationship. Thus, the interplay of photosynthesis and respiration/upwelling combined with the cooling effect of temperature influence the undersaturation of fCO\(_2\) and O\(_2\) for AR in the QIV. This highlights the complex interaction between physical-chemical and biological processes setting the balance in the sea-air CO\(_2\) in the Southern Ocean (Marinov et al., 2006; Henley et al., 2020).

Finally, the distribution of observations in QIV for the MR region was located between 43 and 66°S (Figure 6, sky blue) in the eastern sector of the transect. The significant positive correlation between fCO\(_2\)sat and O\(_2\)sat (0.63, \( p < 0.001 \); Figures 5C,IV) indicates the dominance of the temperature effect in QIV. The
fCO$_2$/O$_2$ variation for the case waters in QIV along the MR transect is presented in Figure 8C. Despite the high positive correlation value of 0.63, indicative of temperature dominated influence, respiration/remineralization process (positive values of AOU, Figure 3I) and photosynthesis (Figure 3I) also contribute to the observed CO$_2$/O$_2$ saturation states. In the northern reaches of the MR transect near the STF and the subtropical oligotrophic waters, the undersaturation of fCO$_2$ and O$_2$ (Figures 3C,C) indicated the influence of other processes (see Section “Regional Drivers of FCO$_2$”) with the surface water properties of the controlling variables defined in this study.

Given the above, it can be said that the cooling effect of temperature derived for QIV was not spatially determined except in the part of the colder and fresher waters of the Antarctic coast. The complex interplay between temperature, photosynthesis, and respiration influences the undersaturation of fCO$_2$ and O$_2$ (Figures 3C,C) indicated the influence of other processes (see Section “Regional Drivers of FCO$_2$”) with the surface water properties of the controlling variables defined in this study.

CONCLUSION

This study investigates the drivers of variability for fCO$_2$ and estimates the FCO$_2$ during autumn in the Atlantic sector of the Southern Ocean, spanning across the Weddell gyre and ACC. The net CO$_2$ flux to/from the atmosphere is driven by the concentration gradient between the atmosphere and the ocean and agitated by the windspeed. During the period of occupation of the Southern Ocean for this study, the whole region along the study transect acted as a net CO$_2$ sink for atmospheric CO$_2$ with an average FCO$_2$ of $-6.2$ ($\pm 8$) mmol m$^{-2}$ d$^{-1}$. The highest net uptake was observed for the Astrid Ridge region near the Antarctic continent, mainly driven by elevated biological productivity. Using fCO$_2$/O$_2$ correlations, the underway-surface observations were partitioned into four quadrants, driven predominantly by (I) photosynthesis, (II) warming, (III) respiration/remineralisation, and upwelling, and (IV) cooling. Describing the spatial distributions of the fCO$_2$/O$_2$ correlations for the quadrants relative to the surface water properties of controlling variables (chl-a, AOU, SST, and SSS) shows the complex interplay of the different processes driving the fCO$_2$ distributions. Overall, the observations illustrated the complex interaction between physical-chemical and biological processes setting the balance in the sea-air CO$_2$ flux in the Southern Ocean and explains the variation in the sea surface CO$_2$. For instance, the upwelling of CO$_2$-rich waters is offset by CO$_2$ uptake through photosynthesis as observed in the MR and WS region.

Finally, this work contributes to increasing the spatial and temporal coverage of observational fCO$_2$ data with surface layer properties in the Weddell gyre and Antarctic coastal area, given the importance of the region to the Southern Ocean’s role in the global ocean CO$_2$ uptake.

DATA AVAILABILITY STATEMENT

The datasets used for this study are available at the Norwegian Polar Institute Data Center (NPDC) at the following website https://doi.org/10.21334/npolar.2020.cc4fb5f and in the Surface Ocean CO$_2$ Atlas (SOCAT).

AUTHOR CONTRIBUTIONS

AF, MC, and AR conceptualized the project. AF and MC collected the data and contributed specialist knowledge toward
constructing and finalizing the manuscript. MO did the data analysis, interpretation, and compiled the manuscript. AR and WJ as the graduate adviser of MO also contributed specialist knowledge towards constructing and finalizing the manuscript. All authors contributed to the manuscript revisions, read, and approved the submitted version.

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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.2020.614263/full#supplementary-material

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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