# Biophysical coupling of seasonal chlorophyll-a bloom variations and phytoplankton assemblages across the Peninsula Front in the Bransfield Strait

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Abstract. This study investigates the spatio-temporal variations of the <u>chlorophyll-a (chl-a)</u> blooms in the Bransfield Strait

- 10 (BS) at a climatological scale (1998-2018). We propose that a suitable monitoring of these blooms can be achieved through remotely-sensed observations only if the <u>BS</u> is divided following the Peninsula Front (PF), which ultimately influences the phytoplankton assemblage. Our analysis is based on characterising climatological fields of Sea Surface Temperature (SST), air temperature, <u>sea jce coverage</u>, chl-a concentrations and wind stress, guided by synoptic novel and historical *in situ* observations which reveal two niches for phytoplankton assemblage: the Transitional Bellingshausen Water (TBW) and
- 15 Transitional Weddell Water (TWW) pools. The TBW pool features stratified, less saline, warmer waters with shallow mixed layers, while the TWW pool features well-mixed, <u>saltier</u>, and <u>colder</u> waters. We identify that the 0.6°C isotherm corresponds to the <u>summertime</u> climatological <u>PF</u> location, effectively dividing the <u>BS</u> in two different scenarios. Furthermore, the 0.5 mg m<sup>-3</sup> chl-a isoline aligns well with the 0.6°C isotherm, serving as a threshold for chl-a blooms of highest concentrations around the South Shetland Islands. These thresholds enable for the first time the monthly climatological descriptions of the
- 20 two blooms developing in BS at both sides of the <u>PF</u>. We think this approach underscores the potential of combining SST and chl-a data to monitor the year-to-year interplay of the chl-a blooms occurring in the TBW and TWW pools contoured by the <u>PF</u>.

**Keywords**: Bransfield Strait, Chlorophyll-a bloom, Peninsula Front, Sea Surface Temperature, Biophysical coupling, Phytoplankton assemblage.

#### **1** Introduction

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Antarctic marine ecosystems are highly dependent on the seasonal cycle of the ocean-atmosphere interaction and associated sea ice dynamics (Schofield *et al.* 2010; Ducklow *et al.* 2013; Montes-Hugo *et al.* 2009; Sailley *et al.* 2013; Brown *et al.* 2019). Through this work we aim to characterise the seasonal variability of the biophysical coupling supporting the surface

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chlorophyll-a bloom in the Bransfield Strait (BS), which is located in the Southern Ocean (SO) between the South Shetland 45 Islands (SSI) and the Antarctic Peninsula (AP).

The BS is connected to the west with the Bellingshausen Sea and to the east with the Scotia and Weddell Seas (Figure 1). The confluence of water masses of different origin in this area leads to a highly dynamic system where different ocean properties interact. Most of the previous studies have described the ocean surface dynamics of the BS based on summertime 50 data (Figure 1a), when two inflows enter the strait and circulate cyclonically (Grelowski et al., 1986; Hofmann et al., 1996; Zhou et al., 2006; Sangrà et al., 2017). The horizontal and vertical structure of the summertime circulation and hydrography

in the BS, when the chlorophyll-a bloom develops, may be summarised as follows.

The western inflow is the Bransfield Current (BC; Niller et al., 1991; Zhou et al., 2002, 2006) which is a coastal jet flowing 55 to the northeast and transporting Transitional Zonal Water with Bellingshausen influence (TBW) along the southern slope of SSI. TBW is typically found within the first 300 m as a well-stratified and relatively warm ( $\Theta > -0.4^{\circ}$ C) and fresh (<34.45) water, (Sangrà et al., 2017), seasonally originated in the Bellingshausen Sea and Gerlache Strait due to summer heating and ice melting (Tokarczyk, 1987; García et al., 1994; Sangrà et al., 2011). The eastern inflow is the Antarctic Coastal Current (CC), which travels southwestward and transports Transitional Zonal Water with Weddell influence (TWW) in this area of

60 Antarctica, countering the northern AP coastline. TWW is distinguished by colder ( $\Theta < -0.4^{\circ}$ C) and saltier (>34.45) waters than TBW (Sangrà et al., 2017), coming from the Weddell Sea (Tokarczyk, 1987; García et al., 1994) and being rather homogeneous throughout the water column (Grelowski et al., 1986; Hofmann et al., 1996; García et al., 2002; Zhou et al., 2002). Between the BC and the CC, there is a street of mesoscale anticyclonic eddies (AEs) of TBW characteristics (Sangrà et al., 2011, 2017). Lastly, the BC recirculates around the islands, transporting TBW as a part of the summertime circulation (Sangrà et al., 2017).

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As will be analysed thoroughly in this work, at the encounter of TBW and TWW a key feature in the biophysical coupling of the chlorophyll-a bloom emerges, the Peninsula Front (PF; García et al., 1994; López et al., 1999). The PF is generally formed at about 20-30 km from the AP slope as a mesoscale shallow structure of 10 km wide (Sangrà et al., 2011)

- 70 confronting TBW and TWW, and expanding from the surface down to ~100 m. On the opposite side of the BS, closer to the SSI slope, one finds the subsurface Bransfield Front (BF) between 50 to 400 m (Niller et al., 1991; García et al., 1994; López et al., 1999), where TBW opposes TWW. The latter water mass widens its domain at depth over the whole strait. Generally, the BF extends between 10 to 30 km offshore from the SSI coastlines, being at its widest when approaching King George Island (Veny et al., 2022).
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As for the bio-chemical context, SO waters are characterised by high-nutrient low-chlorophyll (HNLC) conditions which are equivalent to high concentrations of inorganic macronutrients, but low phytoplankton abundance and rates of primary

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production (Mitchell and Holm-Hansen, 1991; Chisholm and Morel, 1991). Values of chlorophyll-a (<u>chl-a</u>) concentrations <u>in</u> the SO are frequently around 0.05-1.5 mg m<sup>-3</sup> (Arrigo *et al.*, 1998; El-Sayed, 2005; Marrari *et al.*, 2006).

- 85 However, inshore waters west of the Antarctic Peninsula (wAP) are among the most productive regions of the SO (El-Sayed, 1967; Comiso *et al.*, 1990; Sullivan *et al.*, 1993). Thus, the <u>chl-a</u> concentration in the wAP differs from that found in the <u>SO</u>, with values ranging more extensively from 0.16 to 7.06 mg m<sup>-3</sup> (Aracena *et al.*, 2018). Yet, Hewes *et al.* (2009) reported that concentrations in this region are generally not higher than 3 mg m<sup>-3</sup> based on satellite and *in situ* data. Previous studies, based mainly on summertime data (few during late-spring), have also characterised the spatial distribution of chl-a in the BS.
- 90 The distribution was described as patchy and related to the spatial domain of each characteristic water mass (Basterretxea and Aristegui, 1999) and the Upper Mixed Layer (UML) depth, which reflects vertical stability (Lipski and Rakusa-Suszczewski, 1990; Hewes *et al.*, 2009). Then, chl-a was found inversely correlated with UML depth and positively correlated with temperature, i. e. concentrations reach their maximums when UML depth is shallow, temperature is relatively high and surface waters are iron-replete (Hewes *et al.*, 2009). More recently, García-Muñoz *et al.* (2013) reported that the
- 95 highest phytoplankton concentrations along a cross-strait central transect in BS were correlated with relatively warm and stratified TBW waters. The authors also found that nanophytoplankton (2–20 μm) was predominant throughout the study area, which was dominated by small diatoms. However, haptophytes distribution co-varied with small diatoms and also appeared in well mixed TWW waters. As to diatoms, García-Muñoz *et al.* (2013) also found a shift from smaller to larger diatoms when closer to the AP. Sharply, cryptophytes were restricted to stratified TBW waters. These authors concluded, for
- 100 the first time in the literature, that phytoplankton assemblages around the SSI were strongly connected with the Bransfield Current System. This is the seed of our working hypothesis: the horizontal extent of the surface signal of chl-a bloom in the BS may vary monthly from spring to summer (months of bloom development) according to the spatial distribution of the <u>PF</u>, through which TBW and TWW interact and embed different phytoplankton assemblages. This being confirmed, one could long-term monitor the biophysical coupling between the surface chl-a bloom and the <u>PF</u> using remotely-sensed observations

105 of chl-a and Sea Surface Temperature (SST).

Nevertheless, the chl-a distribution in high latitudes has also been reported to be coupled to other biophysical factors such as sea ice formation and atmospheric forcing. It is known that the seasonal sea-ice extent and its timing are likewise determinant for the chl-a development (Garibotti *et al.*, 2003; Smith *et al.*, 2008). Furthermore, sea ice conditions are influenced by atmospheric forcing such as the regional wind stress magnitude and direction, which vary from year-to-year (Smith *et al.*, 2008). This manner, wind alterations significantly affect the sea-ice concentration around the West Antarctica (Holland and Kwok, 2012; Eayrs, *et al.*, 2019), although there are also seasonal and regional variations in the response (Kusahara *et al.*, 2018).

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Given the above scenario, one must confront that chl-a concentrations are not independently controlled by a single factor, and their temporal variations are complex, influenced by seasonal, intra- and interannual processes (Siegel *et al.*, 2002;
Stenseth *et al.*, 2003); e. g. sea ice-ocean interactions may even evolve differently from one season to the next one (Stammerjohn *et al.*, 2012; Holland, 2014).





In this work, we provide a comprehensive description of the seasonal variations of chl-a concentrations in the <u>PS</u> accounting for the biophysical coupling supporting its development. We hypothesize that this biophysical coupling is strongly conditioned by the spatio-temporal variability of the <u>PF</u>, as has been already argued.

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The structure of this manuscript is as follows. In Section 2, we describe the data and methods. In Section 3, we present and discuss the results distributed in four subsections. In Section 3.1, we set our working hypothesis by analysing observational data from two oceanographic cruises: CIEMAR (December 1999) and COUPLING (January 2010). In Section 3.2, we





construct satellite-based climatologies and examine the seasonally-varying horizontal distribution of \$ST, Sea Ice Coverage (SIC), chl-a concentrations, wind stress and Ekman pumping. In Section 3.3, we present the monthly evolution of the latter variables, along with air temperature, in order to characterise the spatio-temporal variability of the bloom according to four distinct regions (Figure 1b), which will be accounted for in the text. In Section 3.4, we address a review of works investigating the phytoplankton assemblage in the BS, bearing in mind the biophysical coupling previously described, in order to provide further insights based on the *state of the art* knowledge. Lastly, in Section 3.5, we construct satellite-based monthly climatologies of SST and chl-a along the same transects sampled during the CIEMAR and COUPLING cruises to

155 present a climatological context to our working hypothesis, through which the spatial distribution of the chl-a blooms in the <u>BS</u> varies according to the <u>PF</u> (monitored via SST), which contours the hydrographic area for TBW and TWW hosting different phytoplankton assemblages. Section 4 presents a summary of the main conclusions.

#### 2 Data and methods

*In situ* observations and remotely-sensed measurements are detailed in the following, separately, for clarity. Seasons are defined following Zhang *et al.* (2011) and Dotto *et al.* (2021), as: summer (January-February-March), autumn (April-May-June), winter (July-August-September) and spring (October-November-December).

#### 2.1 In situ observations: Antarctic cruises

The data inspiring the hypothesis that we address in this work, i. e. the spatial distribution of the <u>chl-a</u> bloom in the <u>BS</u> as strongly conditioned by the <u>PF</u>, rely to a great extent on Conductivity, Temperature, Depth (CTD) and fluorescence measurements collected from two interdisciplinary cruises: CIEMAR and COUPLING. The fluorescence measurements were collected with an ECO fluorometer, which measures fluorescence from <u>chl-a</u>, fDOM, uranine, rhodamine, and phycocyanin and phycoerythrin. In this work, we analyse the fluorescence from <u>chl-a</u>.

- On the one hand, the CIEMAR cruise was conducted in December 1999 (Corzo *et al.*, 2005; Primo and Vázquez, 2007;
  Sangrà *et al.*, 2011), and, on the other hand, the COUPLING cruise was conducted in January 2010 (Hernández-León *et al.*, 2013; Sangrà *et al.*, 2014; Sangrà *et al.*, 2017). Both cruises were carried out onboard the R/V BIO Hespérides. For further details about the CTD stations map of both cruises, the reader is referred to Sangrà *et al.* (2011) and Sangrà *et al.* (2017).
- Additionally, *in situ* surface and subsurface (10 m depth) temperature measurements were downloaded from PANGAEA
   (https://www.pangaea.de/) and the World Ocean Database (WOD; <a href="https://www.ncei.noaa.gov/products/world-ocean-database">https://www.ncei.noaa.gov/products/world-ocean-database</a>) in order to assess the goodness of available open-access remotely-sensed products of <u>\$ST</u> and support the choice of the product providing the best fit. In Table A1 of the Appendix <u>A</u>, a summary of the cruises and corresponding dates for the CTD measurements used to this aim is presented.

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# 2.2 Remotely-sensed products: Sea Surface Temperature and Sea Ice Coverage

We use satellite data of SST and SIC from the Operational Sea Surface Temperature and Ice Analysis (OSTIA; Good et al.,	
2020) downloaded from the Copernicus Marine Environment Monitoring Service (CMEMS; https://marine.copernicus.eu/)	$ \subset  $
and developed by the United Kingdom Met Office. The motivation behind this choice is supported by a quantitative	Z
intercomparison between available SST open-access products and in situ temperature measurements (see this analysis in the	(

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Appendix A).

OSTIA provides the <u>SST</u> free of diurnal variability and the sea ice concentration. It is a reprocessed dataset with a high grid resolution of 0.05°, which accounts for both in situ and satellite data, and presents a processing level L4. This work analyses OSTIA data from 1998 to 2018.

## 2.3 Remotely-sensed products: Chlorophyll-a

We compute monthly climatologies of surface chl-a concentrations based on multi sensors/algorithms. The product name is OCEANCOLOUR GLO BGC L4 MY 009 104, obtained from CMEMS (https://marine.copernicus.eu/). Chl-a data have a spatial and temporal resolution of 4 km and from 09/1997 to present, respectively, and a processing level L4. This work analyses concentrations from 1998 to 2018.

# 2.4 Remotely-sensed products: Wind and Air Temperature

We use the monthly averaged reanalysis of air temperature at 2 m and wind components at 10 m from ERA5 (Hersbach et al., 2020), which have a horizontal resolution of 0.25° over the period 1940 to present. From the wind components, we calculate the wind-stress and Ekman pumping.

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We calculate the wind stress ( $\tau$ ), and wind stress zonal ( $\tau_x$ ) and meridional ( $\tau_y$ ) components, following Equations 1-3 (Patel, 2023):

	$\tau = \rho \cdot U_{10}^2 \cdot C_{D,\bullet}$	(1)
	$\tau_x = \rho \cdot U_{10} \cdot u \cdot C_{D_{-}},$	(2)
225	$\tau_{\nu} = \rho \cdot U_{10} \cdot \nu \cdot C_{D},$	(3)

where  $\rho$  is the air density (1.2 kg m<sup>-3</sup>);  $U_{10} = \sqrt{u^2 + v^2}$  is the absolute value of the wind speed at 10 m above the surface (u and v are the eastward and northward wind speed components, respectively); x and y are the eastward and northward spatial coordinates; and,  $C_D$  is the drag coefficient, which is a function of wind speed,  $U_{10}$ . The equations used for wind stress computation are based on Gill (1982) formula and a non-linear  $C_D$  based on Large and Pond (1981), modified for low wind

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speeds (Trenberth *et al.*, 1990). We note that the mean  $C_D$  we obtained for our climatological maps in the BS is  $1.4 \times 10^{-3} \pm 0.16 \times 10^{-3}$ , analogous to the values reported by Kara *et al.* (2007) over the SO.

We also compute the Ekman vertical velocity as following:

$$w_E = \frac{1}{\rho_0} \operatorname{curl}\left(\frac{\tau}{f}\right)$$

(4)

where  $\rho_0$  is the water density (1025 kg m<sup>-3</sup>);  $f = 2\Omega sin\varphi$  is the Coriolis frequency ( $\Omega$  is the Earth rotation rate, 7.2921x10<sup>-5</sup> rad s<sup>-1</sup>, and  $\varphi$  is the latitude). Positive (negative)  $w_E$  values indicate upward (downward) velocities leading to upwelling (downwelling).

#### **3** Results and Discussion

Through this section we assess the major physical drivers potentially conditioning the vertical and horizontal structure of the chl-a bloom in the BS. To this aim, in Section 3.1 we analyse the vertical and horizontal structure of two chl-a blooms in the BS based on hydrographic measurements from two cruises (1999 and 2010) along three cross-strait transects (T-I, T-II, T-III). In Section 3.2 we analyse the seasonal variations of the horizontal structure of the chl-a bloom and the PF from a climatological perspective based on remotely-sensed observations over a 21-year period (1998-2018): SST, SIC, wind stress and Ekman pumping. Next, in Section 3.3, we examine in detail the monthly climatologies of selected boxes of study in the BS (adding air temperature to the analysis). In Section 3.4, we provide a summary review of the research on the

255 phytoplankton assemblage in the <u>BS</u>. Lastly, in Section 3.5 we construct satellite-based monthly climatologies of SST and chl-a along the same locations as the transects T-I, T-II and T-III to provide a statistically robust (i. e. climatological) context to our working hypothesis, through which the chl-a bloom extent varies according to the PF.

# 3.1 Vertical and horizontal structure along CIEMAR and COUPLING transects

We present the vertical structure of temperature, salinity, density, and fluorescence in the <u>BS</u> (Figures 2, 3 and 4) based on data collected during the multidisciplinary CIEMAR and COUPLING cruises through late-spring 1999 and earlysummertime 2010, respectively. The physical oceanographic aspects from these cruises were firstly presented in Sangrà *et al.* (2011) and Sangrà *et al.* (2017).

Previous studies based on these measurements and in line with biophysical phenomena focused, among other aspects, on turbulence as a driver for phytoplankton distribution as well as on mesoscale physical features as key players in determining phytoplankton assemblages (García-Muñoz *et al.*, 2013; Macías *et al.*, 2013; Sangrà *et al.*, 2014). In García-Muñoz *et al.* (2013), following COUPLING cruise measurements, the authors concluded that phytoplankton assemblages around the SSI were strongly connected with the Bransfield Current System. Furthermore, the authors suggested that, considering the

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 $τ_x = ρ C_p U_{10}u$ ;  $τ_y = ρ C_p U_{10}v$ , → → → → → → → → → (1)where  $τ_x$  and  $τ_y$  are the zonal and meridional wind stress components, respectively; ρ is the air density (1.225 kg m<sup>3</sup>);  $C_p$  is the drag coefficient (1.25x10<sup>-3</sup>; Kara *et al.*, 2007);  $U_{10} = \sqrt{u^2 + v^2}$  is the wind speed at 10 m above the surface; *u* and *v* are the eastward and northward velocity components; and, *x* and *y* are the eastward and northward spatial coordinates, respectively **Deleted:** 2

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- 285 recurrence of the Bransfield Current System during the austral summer, the observed distribution of phytoplankton, which responded to this current system, should also be a quasi-permanent feature. In the following, we combine the measurements from the two cruises for the first time to address this working hypothesis, where we add and highlight that the key player appears to be the cross-strait gradient marked at surface by the <u>PF</u>, and that this may enable the long-term monitoring of the biophysical coupling between the surface chl-a bloom and the PF based on satellite measurements.
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  - The CIEMAR transects cover the region from Livingston Island and King George Island towards the northern tip of the <u>AP</u> (Figures 2 and 4, respectively). Both transects were conducted in December 1999. On the other hand, the COUPLING transect covers the region from Nelson Strait to the <u>AP</u> tip (Figure 3) and was carried out in January 2010.
- 295 For clarity, we name the three transects of study as T-I, T-II and T-III moving from west to east. Thus, transect T-I (Figure 2) and T-III (Figure 4) correspond to CIEMAR and present novel measurements of fluorescence (not previously published); while transect T-II (Figure 3), located in between T-I and T-III, corresponds to COUPLING. The three transects originate over the shelf of the SSI and extend towards the AP, running nearly perpendicular to the main axis of the strait. Notably, measurements from both sea trials agree well in showing a coherent vertical and horizontal structure of hydrography and 300 fluorescence. In Figures 2-4, two panels are always dedicated to show the TS diagram and the station map to help the reader in following the ocean property descriptions.
- Starting with T-I (Figure 2), this originates to the south of the Livingston Island and extends towards the AP. Along T-I, temperatures are above 0°C in the upper 40 m for its full extent (Figure 2a), being relatively higher in the proximity of the 305 SSI at a distance of 0-20 km and close to the AP, where they reach near surface values exceeding 0.6°C and 0.4°C, respectively. On the other hand, salinity is remarkably fresher and lighter near the SSI (S < 34 and  $\sigma_{\theta} < 27.3$  kg m<sup>-3</sup>), as opposed to saltier and denser waters towards the  $\Delta P$  (S > 34.3 and  $\sigma_0 \sim 27.6$  kg m<sup>-3</sup>; Figure <u>5</u> 2b and 2c, respectively). Fluorescence levels exceeding 2-3 (Figure 2d) peak in the warmest (>0.2°C) and lightest surface waters (upper 50 m) near the SSI (stations B1-B2) and near the AP (stations B5-B7). The isopycnal of 27.64 kg m<sup>-3</sup> is highlighted in black in all 310 vertical sections as a reference to the water mass boundary between TBW ( $\sigma_{\Theta} < 27.64 \text{ kg m}^3$ ) and TWW ( $\sigma_{\Theta} > 27.64 \text{ kg m}^3$ ) <sup>3</sup>; Sangrà et al., 2011, 2017). These observations indicate the presence of relatively warmer TBW flowing near the surface to the south of Livingston Island. Coastal signals of Bransfield Strait Shelf Water (BS Shelf Water), characterised by lower salinity values (Zhou et al., 2006; Polukhin et al., 2021), were also detected. In addition, relatively colder TWW flows closer to the AP at deeper levels (> 60 m depth), where fluorescence sharply diminishes (< 0.5) along the entire transect (Figure 315 2d). The PF is not visible along T-I given the basin-wide extent of TBW at surface, which prevents shoaling of TWW.

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- 340 T-II (Figure 3) originates to the south of the Nelson Strait and extends towards the AP. Largely, along T-II, TBW waters are visible as warmer (Θ > -0.4°C) and fresher (<34.45) waters than TWW (Sangrà *et al.*, 2017). The subsurface signal of TWW extends towards the SSI, confronting TBW between 60-120 m depth at around stations 2-3, where they form the <u>BF</u> (Sangrà *et al.*, 2011). Newly, the high fluorescence patch (> 1) extends within the warmest and freshest surface waters (Figure 3d) from the surface down to 60 m depth at its deepest, contouring the isotherm of 0.2°C from the SSI until stations 9-10. At this location, the 0.2°C isotherm reaches the surface, temperature decreases rapidly towards the AP (<-0.6°C; Figure 3a), and salinity and density increase (>34.3 and >27.64 kg m<sup>-3</sup>; Figures 3b and 3c). This gradient forms the PF, where TBW and TWW.
  - TWW confront each other close to the AP. Remarkably, near surface (0-30 m) fluorescence levels decrease below 0.5 (Figure 3d) on the TWW side of the <u>PF</u>.

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Figure 3. Same as in Figure 2 but for T-II, surveyed during the COUPLING cruise (January 2010) and running from Nelson Strait to the Antarctic 360 Peninsula. Additionally, the dashed black line represents the isopycnal of 27.55 kg m<sup>-3</sup> which is used as a reference more adjusted to our dataset to distinguish between TBW and TWW.

Lastly, T-III (Figure 4) originates to the south of King George Island and extends towards the AP. Generally, we observe an analogous vertical structure to that described for T-II, suggesting that a horizontal coherence exists between transects;
especially when accounting that differences with T-I are due to the latter is at a farther distance from the Weddell Sea and, hence, presents a weaker signal of TWW at surface. As observed in Figures 2 and 3, the chl-a bloom suggested by high fluorescence values (> 1) is again embedded within the pool of TBW closer to the SSI, where waters are relatively warmer and fresher as compared to TWW waters close to the AP. Newly, the PF (stations T10-T11) appears to delimit the surface easternmost reach of the patch with highest fluorescence. However, we must also note that, between the PF and the <u>AP</u>, a
less prominent and coherent patch of values higher than the baseline exists down to nearly 120 m depth, both in T-II and T-III (fluorescence > 0.5 and >1, respectively).

Remarkably, two other studies (Basterretxea and Arístegui, 1999; Gonçalves-Araujo *et al.*, 2015) have also captured a consistent cross-strait pattern where the highest chl-a concentrations are embedded within the TBW reservoir in the first ~60 m of the water column and the easternmost extent of this signal coincides with the location of the <u>PF</u>, through which TBW

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and TWW interact. In both cases, in spite of the sharp decrease of chl-a across the PF, the authors also found that chl-a 10

concentrations were not low on the TWW reservoir, but also relatively high although occupying a wider depth range (0-100 m). Their vertical sections were constructed from ship-based measurements collected along a transect parallel to T-III but farther north, departing from King George Island, in January 1993 and February-March 2009: Figure 6 in Basterretxea and Arístegui (1999); and, Figure 3 in Gonçalves-Araujo *et al.* (2015), respectively. This supports the existence of different phytoplankton assemblages occupying different niches according to the dominant water masses.







Figure 4. Same as in Figure 2 but for T-III, surveyed during the CIEMAR cruise (December 1999) and running from King George Island to the Antarctic Peninsula. Additionally, the dashed black line represents the isopycnal of 27.55 kg m<sup>-3</sup> which is used as a reference more adjusted to our dataset to distinguish between TBW and TWW.

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In all panels d) from Figures 2-4, a solid red line is added to indicate <u>LIML</u> depth. Roughly, this estimate of the UML depth, fits well with the depth of the high fluorescence patch embedded within the TBW reservoir, and which keeps phytoplankton under favourable light conditions, a better supply of dissolved iron (Prézelin *et al.*, 2000) and within a depth range with proper conditions for accumulation of phytoplankton biomass (Mukhanov *et al.*, 2021; Mendes *et al.*, 2023). Accordingly, relatively high fluorescence (>0.5) is accumulated along the entire BS in T-I, where UML depth is relatively low (<60 m), especially in stations B5-B7 (fluorescence above 2 and UML depths of ~15 m). The same pattern applies along T-II and T-III, with the high fluorescence patch embedded within the UML. In T-II, highest fluorescence (~2) is located near the PF, in

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station 9, where lowest UML depths occur (10 m). Analogously, in T-III fluorescence values of 2 are closer to the SSI

station 2 where UML depths are around 25 m. Moreover, near the AP, relatively low UML depths (10 m) are also observed in station T14 jointly with fluorescence values of 1.

- 405 Following results in García-Muñoz *et al.* (2013), the fluorescence observations presented here from the COUPLING cruise (T-II in Figure 3) can be attributed to different phytoplankton assemblages, as briefly introduced in Section 1, namely: cryptophytes in the upper 60 m of the TBW reservoir between the <u>BF</u> and the <u>PF</u>; and, nanophytoplankton along the full transect but at higher abundances for the largest fraction in the TWW reservoir, accounting for the weaker but deeper signal in fluorescence (from the surface down to 100 m). This suggests that the fluorescence signal measured by the ECO
- 410 fluorometer might be dominated by cryptophytes. Whether this is also the case for the CIEMAR transects (T-I and T-III in Figures 2 and 4) is a feature we cannot confirm in absence of a phytoplankton assemblage study for that cruise. However, the fluorescence distribution appears consistent a decade apart in showing highest and shallower values within the relatively warm and stratified TBW reservoir, and lower but deeper values within the cold and well-mixed TWW reservoir. The stronger signal in fluorescence during the CIEMAR could be then attributed to a higher abundance of cryptophytes within
- 415 the TBW reservoir, if we assume that the pattern observed by García-Muñoz *et al.* (2013) is recurrent over time. Supporting this, recent studies have also confirmed the preferred niche of cryptophytes in the BS is the relatively warmer, less saline and stratified waters of the TBW reservoir, where they also compete with diatoms (Mendes *et al.*, 2013; Gonçalves-Araujo *et al.*, 2015; Mukhanov *et al.*, 2021; Costa *et al.*, 2023; Mendes *et al.*, 2023).
- 420 Results from the *in situ* measurements collected during the CIEMAR and COUPLING cruises, occurring a decade apart, plus more recent evidence of phytoplankton assemblages following the ocean dynamics of the Bransfield Current System jointly support further the basis of our working hypothesis: the biophysical coupling between the spatial distribution of the surface chl-a bloom and the <u>PF</u> in the <u>BS</u> may be long-term monitor using remotely-sensed observations of chl-a and SST.
- 425 In the following section, we analyse a set of satellite-based climatologies with the aim to demonstrate that the horizontal variability of the <u>PF</u> (and hence the interaction between TBW and TWW) plays a major role in determining the spatial extent of the patch with highest surface chl-a bloom in the <u>BS</u>. We complete this analysis by considering the role of several physical drivers which also contribute to set the niche for phytoplankton assemblage through a biophysical coupling. We expect this joint climatological perspective of the seasonal variations of the chl-a bloom and the PF, unprecedented in the literature, provides the basis for their long-term monitoring. Counting with a robust long-term phytoplankton monitoring approach will enable a better understanding of the biophysical coupling setting the baseline of the marine food web in the <u>BS</u>.

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#### 445 3.2 Seasonal variations of the chl-a bloom and Peninsula Front coupling

The remotely-sensed climatologies of SST, SIC, wind stress, Ekman pumping and chl-a were computed for the period 1998 to 2018, and are presented in Figures 5-8, respectively.

Regarding the <u>SST</u> (Figure 5), the most outstanding feature governing the summer months in the <u>BS</u> is a strong cross-strait gradient, where high temperatures (>1°C) spread around the SSI and lower temperatures (<0°C) appear to enter into the basin from the Weddell Sea, turning around the AP and spreading southwestward along the peninsula shelf. This strong temperature gradient is the surface signal of the <u>PF</u>, where TBW confronts TWW. Previous studies, based on *in situ* summertime data, have used different thresholds for the isotherm characterising the location of the PF at surface, where TBW and TWW interact: Sangrà *et al.* (2017) used the isotherm of -0.4°C, while Catalán *et al.* (2008) used the isotherm of

455 1°C. The choice of these isotherms is not trivial and one must identify the isotherm embedding the water body flowing from the Weddell Sea into the <u>BS</u>, thus separating TWW from TBW. Looking at Figure 5a, we note the climatological isotherm characterising the PF location at surface during the summer months corresponds to 0.6°C isotherm. Through autumn and spring, the PF is also visible although a different isotherm rises as characteristic of this thermal front, being -1.2°C and -0.8°C, respectively. Lastly, during the winter months the surface signal of the PF vanishes, as one could expect, due to the

460 atmospheric forcing prevailing in the homogenisation of the upper ocean. Within the strait, surface temperatures are around -1.8°C ± 0.2°C.

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It is worthwhile noting that in Figures 3 and 4 we used the 0.2°C and 0.8°C isotherms, respectively, reaching the surface to define the location of the <u>PF</u>, and in Figure 5 we used <u>a</u> different isotherm. This is not in contradiction. One must keep in mind that the 0.2°C and 0.8°C characteristic isotherms worked well through synoptic transects, which took place in late December and January, while in Figure 5a a summertime climatological field is examined after time-averaging 3 data months over a period of 21 years. This accounts for the seasonally-varying values provided above, and which differ from the synoptic values.

- 470 To the best of our knowledge, this is the first time that a remotely-sensed SST seasonal climatology is shown with the focus in the BS. This prevents us from comparison with previous studies also using remotely-sensed observations. However, in the Appendix <u>A</u> we present an examination of the goodness of SST satellite measurements against concomitant *in situ* measurements, finding that a high correlation exists between the product we use (OSTIA), and *in situ* measurements (R<sup>2</sup> = 0.849). Also, the summertime field is in agreement with patterns reported in the literature for this season and based on *in situ*
- 475 hydrographic measurements (Sangrà et al., 2011, 2017). Additionally, we used a recently published seasonal climatology of hydrographic properties in the <u>BS</u> based on *in situ* measurements (Dotto et al., 2021), and produced an analogous figure (not shown) to our Figure 5 (with the same contour lines and colorbar). The comparison supports the major features of the

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490 seasonal patterns described above about the remotely-sensed SST signals driven by the ocean dynamics and atmospheric forcing governing the region. Exceptions occur north of the SSI in autumn and inside <u>BS</u> in spring, where the abundance of mesoscale features in the climatology based on *in situ* measurements (Dotto *et al.*, 2021) slightly hampers the view of the mean field pattern.

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Figure 5. Seasonal maps of Sea Surface Temperature (<u>SST</u>; in shades of colours) for: a) summer, b) autumn, c) winter and d) spring. The capital letters between brackets stand for the initial letter of the month. The SST climatologies are averaged from January 1998 to December 2018. The dashed isotherms are plotted at intervals of 0.2°C, while the solid line marks each 1°C interval.

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Figure 6 shows the seasonal <u>SIC</u> as a percentage of area covered by sea ice. A value of SIC about 15% is taken as indicative of the presence of sea ice. Thus, one can consider that during the summer and spring months the <u>BS</u> is generally free of sea ice with SIC < 15%. Through autumn, the atmospheric forcing starts leading the development of the SIC in the <u>BS</u>, which extends firstly over the colder waters of the Weddell Sea intrusion with SIC ranging from 15 to 25% (compare Figure 5b and Figure 6b). This is in agreement with a recent study developed over the western <u>AP</u>, which addresses the role of subsurface

505 Figure 6b). This is in agreement with a recent study developed over the western <u>AP</u>, which addresses the role of subsurface ocean heat on the modulation of the sea-ice seasonality, and highlights the importance of the upper ocean variability in

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setting sea-ice concentrations and thickness (Saenz *et al.*, 2023). Towards winter, the SIC is greater than 25% everywhere in the <u>BS</u> (Figure 6c), promoted by near-freezing sea surface temperatures around  $r1.8^{\circ}$ C ± 0.2°C (Figure 5c).

515 Accounting that the seasonal sea-ice retreat is complete from spring to summer in the entire **PS**, this suggests that the larger freshwater inputs reported in the literature over the TBW domain and contributing to the vertical stabilization of the water column, might be driven by a warmer oceanic forcing over coastal/glacial areas (Cook *et al.*, 2016) rather than by melting of the open ocean sea-ice.

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Figure 6. Same as Figure 5 but for Sea Ice Coverage (SIC). Solid black lines indicate a SIC percentage of 15%, which is the threshold to consider significant the presence of sea ice. Dashed grey lines represent SIC percentages of 25%, 50% and 75%.

525 Following seasonal panels in Figure 7, the development of the chl-a bloom in the <u>BS</u> is particularly revealing when using a logarithmic scale, which highlights spatial patterns otherwise slightly masked due to the strong signal of chl-a east of the <u>AP</u> in the Weddell Sea. West of the <u>AP</u>, chl-a bloom concentrations have been reported to range normally between 0.5-1 mg m<sup>-3</sup>

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(Ducklow *et al.*, 2008; Smith *et al.*, 2008). However, we must note that this threshold varies significantly depending on the study region given that there are areas with naturally either higher or lower phytoplankton concentrations.

Generally speaking, the chl-a bloom in the <u>BS</u> starts developing in spring, reaching its maximum horizontal extent with values above 0.5 mg m<sup>-3</sup> during the summer months, and still presenting patchy regions of high chl-a during autumn (Figure 7). We find that in the <u>BS</u> the isoline of 0.5 mg m<sup>-3</sup> works well as the threshold contouring the chl-a bloom around the SSI, as this appears to embed coherently in space the region with highest chl-a values during the summer months. Through winter, surface chl-a concentrations drop below 0.25 mg m<sup>-3</sup> everywhere in the <u>BS</u> except for the region adjacent to the northern shelf of the westernmost SSI (>0.25 mg m<sup>-3</sup>). We observe that although the <u>BS</u> receives inflows from the Weddell Sea to the east of the <u>AP</u>, the much higher chl-a concentrations present in the Weddell Sea do not extend into the <u>BS</u>. This is in spite of the fact that the waters from the Weddell Sea are continuously propagating around the northern tip of the climatologies covering at its highest values the entire <u>BS</u>. Differently, at its largest extent, with values higher than 0.5 mg m<sup>-3</sup>

(summer months), the chl-a bloom appears constrained to the domain of TBW waters sourced from the Bellingshausen Sea while the presence of TWW waters marks the boundary where chl-a concentrations drop sharply within BS. By comparison between Figure 5a and Figure 7a, it becomes evident that the spatial extent of the surface chl-a bloom surrounding the SSI
(chl-a > 0.5 mg m<sup>-3</sup>) aligns well with the surface signal of the PF in the BS, where TBW and TWW confront each other. To ease visualization of this coupling, the isotherms of 1°C and 0.6°C have been added over the summertime chl-a field (Figure 7a).

This bloom area where chl-a concentrations are higher than 0.5 mg m<sup>-3</sup> coincides in cross-strait direction with the chl-a
bloom boundaries reported by García-Muñoz *et al.* (2013) for cryptophytes and large nanophytoplankton surrounding the SSI (their Figure 4). On the Drake Passage side, the oceanward extent of their bloom ended at the subsurface Shetland Front (García-Muñoz *et al.*, 2013), embedding TBW over the northern shelf of the SSI, and accounting for the recirculation of TBW waters around the archipelago driven by the Bransfield Current (Sangrà *et al.*, 2017). In Figure 5a, the alignment of the subsurface Shetland Front to the north of the SSI is suggested by the isotherm of 1.6°C, which roughly follows the oceanward extent of the surface chl-a bloom (Figure 7a). On the BS side, the chl-a bloom investigated in García-Muñoz *et al.* (2013) also transitioned towards lower values across the PF in agreement with this study (Figures 5a and 7a), and

previous and later works (Basterretxea and Arístegui, 1999; Mendes *et al.*, 2013; Gonçalves-Araujo *et al.*, 2015; Mukhanov *et al.*, 2021). Lastly, we also note the resemblance of our summertime satellite-based climatologies of SST and chl-a (Figures 5 and 7) with those based on eighteen years of summertime hydrographic and chl-a measurements, through which the authors demonstrate that the distribution of high chl-a around the SSI corresponded to shallow UML depths in iron-rich

waters at salinities ~34 (Hewes et al., 2009; their Figure 4).

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Figure 7. Same as Figure 5, but for chlorophyll-a concentrations (chl-a). Solid black lines indicate chl-a concentrations of 0.5 mg m<sup>3</sup>, while solid grey lines represent chl-a concentrations of 0.25 and 1 mg m<sup>3</sup>. Solid and dashed red lines in panel a) indicate 0.6°C and 1°C summer isotherms, respectively (see Figure 5a). For the autumn season (panel b), only the mean of April month<sub>y</sub> is considered due to the absence of data during other months, which results from the presence of ice cover. Similarly, for the winter season (panel c), the mean of August and September months are solely considered for the same reason.

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Figure 8. Same as Figure 5 but for Ekman pumping. Positive (negative) vertical velocities are indicated in shades of red (blue) and represent upwelling (downwelling) processes. Solid black lines refer to zero-velocities. Solid red and blue lines represent vertical velocities of 10 cm day<sup>-1</sup> and -10 cm day<sup>-1</sup>, respectively. Black vectors depict the wind stress. Wind stress reference vector is displayed over the southern AP with a value of 0.02 N m<sup>-2</sup>.

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In Figure 8, the seasonal climatology of the wind forcing acting over the bloom domain is presented following the wind stress field (black vectors) and Ekman pumping (vertical velocity;  $w_E$ ). The dominant winds in the <u>BS</u> are the westerlies (Vorrath *et al.*, 2020), which flow across the strait with greater, basin-wide intensity during winter and spring months. In shades of colours, the Ekman pumping is shown with positive (red) and negative (blue) vertical velocity values implying that wind stress drives, respectively, either local upwelling or downwelling. Generally, upwelling is observed in the <u>BS</u> throughout the year, with a few spatial and temporal exceptions. Downwelling occurs mostly south of King George Island year-round. During autumn, this downwelling area south of King George Island expands towards the AP more extensively. This feature remains through the winter months although constrained to a smaller extent not reaching the AP. During winter and spring, the westerlies drive in the <u>BS</u> relatively strong upwelling vertical velocities especially along the shelf west of the 605 AP.

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Importantly, during spring and summer (months of chl-a development; Figures 7a and 7d), the westerlies appear slightly stronger along the southern shelf of the SSI (over the domain of TBW) as compared to westerlies acting over the shelf west of the <u>AP</u> (over the domain of TWW). Following this, one could reasonably expect deeper mixed layers over the domain where the wind stress forcing is stronger; however, along the southern shelf of the SSI winds favour the Bransfield Current transport of TBW via downwelling-favourable Ekman transport while, along the shelf west of the <u>AP</u>, winds exert a moderate counterforcing to the entrance of the Antarctic Coastal Current driving upwelling-favourable Ekman transport. We find this asymmetry may be contributing to maintain the two distinct niches across the <u>PF</u>: warmer, less saline and stratified waters transported by the Bransfield Current on the TBW side; and colder, saltier and well-mixed waters transported by the Antarctic Coastal Current on the TWW side.

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## 3.3 Monthly variations of the chl-a bloom and Peninsula Front coupling

Following results from previous subsections, we note two areas in the <u>BS</u> are distinctive not only regarding their ocean dynamics as previously known (Figure 1), but also regarding the nature of the chl-a bloom. The first one is where the chl-a bloom spreads with highest concentrations over the relatively warmer and more stratified TBW water, flowing northeastward along the southern shelf of the SSI. The second one is the relatively colder and more homogeneous TWW waters flowing

southwestward along the western shelf of the <u>AP</u>.

For further study of the monthly evolution of ocean and atmospheric conditions influencing the development of the surface chl-a bloom over each area, we divided the <u>BS</u> into four boxes of study (Figure 1b). These boxes were designed to capture, respectively, the northern and southern domain of the surface chl-a bloom embedded in TBW waters south of the SSI, and northern and southern domain of the surface chl-a bloom embedded in TWW waters west of the <u>AP</u>. The resulting climatologies of SST, air temperature, SIC, chl-a, along-shore wind stress and Ekman pumping over the period 1998-2018 are presented in Figure 9 and reveal several spatio-temporal similarities, and differences, which stand out and provide further insights. To this aim the wind stress was decomposed into its along-shore ( $\tau_{xr}$ ) and cross-shore ( $\tau_{yr}$ ) components through rotation of the cartesian components 36.25° in counterclockwise sense.

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The monthly climatologies of SST and air temperature (Figures 9a and 9b) present a coherent seasonal cycle where warmer (colder) temperatures are found within all the regions for summer (winter) months. Spatially, SST within the boxes south of the SSI are generally warmer than those along the west  $\Delta P$ . This is more prominent during summer months, when cross-strait temperature gradients are higher with differences between boxes at opposites of the strait about 0.6°C to 1.4°C (Figure 9a). These differences decrease towards the winter months, when all regions approach at surface near-freezing temperatures

about -1.8°C from July to August. Evolving through the spring months, temperature differences start to increase again but are 19

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655 not higher than 1°C when comparing boxes along the southern shelf of the SSI and along the shelf of the AP. Because boxes south of the SSI, sourced by TBW, depart from higher temperatures and all regions reach near-freezing temperatures during winter, their seasonal amplitudes are larger (and the slopes are more pronounced) as compared to boxes along the shelf of the west <u>AP</u>, sourced by TWW. Thus, the seasonal amplitude of the SST is more than 1.5 times larger for the southern shelf of the SSI (~3°C) than along the west <u>AP</u> shelf (~1.8°C).

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The seasonal amplitude of the air temperature cycle (Figure 9b) is larger than that displayed in SST. However, warmer temperatures are once again observed in the boxes situated along the southern shelf of the SSI, where temperatures evolve from 1°C (summer) towards -5° to -6°C (winter), in contrast to the boxes situated along the west <u>AP</u> shelf, where temperatures evolve from 0°C (summer) towards -8°C (winter). As compared to the SST annual cycle, we observe the air

665 temperature is more homogeneous during the summer and spring months (temperature differences among boxes are < 1.25°C) than during autumn and winter (temperature differences among boxes are > 2.5°C). This is the reverse pattern as shown in SST, where more homogeneous temperatures among regions were found through the winter months. The reason behind the more homogeneous pattern in SST during the winter months may be due to sea water approaching near-freezing temperatures, what sets a threshold which homogenizes the ocean surface under an extreme cooling atmospheric forcing.

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The SIC monthly climatology (Figure 9c) follows an inverse relationship with SST and air temperature (Figures 9a and 9b), where higher values of SIC are found during late autumn, winter and early spring months and absence of sea ice is found through late spring, summer and early autumn months (<15% SIC). Through these latter seasons, the sea ice retreat is driving melting waters into the environment. This is a key factor in pytoplankton biomass accumulation since it allows upper ocean

- 675 stratification during spring/summer, leading to favourable sunlight conditions for phytoplankton to grow (Ducklow *et al.*, 2013). Then, the SIC peaks in July at about 50% closest to the AP tip, at about 40% farther south along the AP and south of the northernmost SSI. One month later the SIC peaks in August at about 30% south of the southernmost SSI.
- Remotely-sensed chl-a observations enable the visualization of the monthly evolution from August to April (Figure 9d) with a data gap due to sea ice coverage from May to July. Yet, a seasonal cycle is visible with higher chl-a concentrations through spring and summer months; lower, and declining, chl-a concentrations through early autumn; and, lower, and increasing, chla concentrations through late winter. This latter increasing trend is concomitant to the decrease in SIC, when sea ice starts melting in August (same month when SST and air temperature also start increasing). Following the literature, the date of the bloom initiation is determined as the first day at which chlorophyll levels rise a 5% above the climatological median (Siegel
- 685 et al., 2002) and stays above this value for at least two consecutive weeks (Thomalla et al., 2011). This threshold was computed assuming linear interpolation over winter to get the climatological median. Bearing these criteria in mind, our climatologies indicate the chl-a bloom in the <u>BS</u> starts through mid-October, departing from a baseline value for chl-a concentrations ~0.2 mg m<sup>-3</sup> in August. Since mid-October (early spring) onwards, chl-a concentrations start increasing in the

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entire BS although more steeply along the southern shelf of the SSI, and slightly delayed in the northern box of the western shelf of the AP.

Chl-a peaks through December and February at 0.68 mg m<sup>-3</sup> south of the southernmost SSI and in February at 0.63 mg m<sup>-3</sup> south of the northernmost SSI (Figure 9d). Along the shelf of the west <u>AP</u>, chl-a peaks to the south in December at 0.43 mg m<sup>-3</sup>; and, one month later, to the north in January at 0.37 mg m<sup>-3</sup>. Generally, although standard deviations are large and
overlap each other cycles, these monthly climatologies suggest a northward development for the chl-a peaks with about 1-2 months of delay.

The pattern described above for the four boxes of study supports the likely existence of two different chl-a blooms developing simultaneously but of different *nature* (i. e. phytoplankton assemblage) in the <u>BS</u>, as suggested by their different intensity and timing (month of initiation and rate of increase). This is in agreement with former results in a series of studies which reported that cryptophytes compete in the <u>BS</u> primarily with diatoms and other nanophytoplankton groups (Mura *et al.*, 1995; García *et al.*, 2013; Mendes *et al.*, 2013; Gonçalves-Araujo *et al.*, 2015; Mukhanov *et al.*, 2021; Costa *et al.*, 2023; Mendes *et al.*, 2023), following strategies to adapt better to water mass distribution in the basin, what ultimately controls the time and space variability of <u>BS</u> phytoplankton communities.

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Only two former studies have reported monthly climatologies of the surface chl-a bloom in the <u>BS</u>; however, none of them framed the boxes of study such that the two blooms were simultaneously, and distinctively, captured. In the first study, the authors placed a rectangular box embedding at the same time both margins of the <u>BS</u>, with no distinction between the TBW and TWW domains (Gonçalves-Araujo *et al.*, 2015). The resulting time-series (2002-2010) displays a strong interannual variability, with summertime values ranging from ~ 1.1 mg m<sup>-3</sup> (2006) to 0.37 mg m<sup>-3</sup> (2003; their Figure 9). In the second

- (15) variability, with summertaine values ranging from ~ 1.1 mg m (2000) to 0.57 mg m (2000, uten Figure 9). In the second study, the authors placed a slanted rectangular box parallel to the SSI coastline, and similar to our two boxes south of the SSI, but in their case extended towards Elephant Island (La *et al.*, 2019). The resulting monthly climatology of the chl-a over the period 2002-2014 (12-year mean) displays the cycle from October to April. The chl-a bloom develops then from baseline concentrations below 0.2 mg m<sup>3</sup> in October to peak concentrations ranging from ~ 1.75-1.95 mg m<sup>3</sup> (their Figure 2) through
- 720 February to March. We attribute the higher climatological values in La *et al.* (2019), occurring about one month later than in our boxes along the SSI, to the different choice of the study area. In their case the northward extension of the box may be including dynamics out of the <u>BS</u>, from the confluence zone with the Weddell Sea. Also, the latter peak in time for this extended region is in agreement with our results in Figure 9d, which suggests the maxima in chl-a develops later as one moves northward along the <u>BS</u>.

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Finally, the along-shore wind stress (Figure 9e) displays year-round downwelling-favourable winds along the southern shelf of the SSI and upwelling-favourable winds along the shelf of the west <u>AP</u>. In all cases, a quarterly cycle stands out with

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740 maxima values (in descending order) in September, December, February and May (i. e. winter, spring, summer and autumn). A similar cycle is found along the shelf of the west AP for the Ekman pumping (Figure 9f), where vertical velocities are upwelling favourable (positive) year-round with a quarterly cycle (same maxima time variability). Along the southern shelf of the SSI, vertical velocities are also upwelling favourable (positive) year-round but less intense and more homogenous through seasons. Peak vertical velocities are 20 cm day<sup>-1</sup>, 17.5 cm day<sup>-1</sup> and 5 cm day<sup>-1</sup> for boxes along the shelf of the west AP, south of the southernmost SSI and south of the northernmost SSI, respectively.

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Figure 9. Monthly climatology over the period 1998 to 2018 of a) Sea Surface Temperature (SST), b) air temperature, c) Sea Ice Coverage (SIC), d) chlorophyll-a (chl-a) concentration, e) along-shore wind stress, and f) Ekman pumping (vertical velocity) for each study box, as delimited in Figure 1b. The horizontal dashed lines in panel c) indicate the threshold (15%) to consider significant the presence of sea ice, while in panel d) indicate the threshold set to identify the initiation of the bloom following Siegel et al. (2002) and Thomalla et al. (2011). The mean monthly values are represented by the solid line, while the corresponding standard deviation is shown in coloured shades.

# 3.4 Spring-Summertime phytoplankton assemblages of the chl-a bloom: a summary review

755 Through the previous section, we have learned that a proper design of study boxes aligned with the climatological summertime position of the PF enables the identification of two distinct chl-a blooms in the BS based on satellite

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measurements according to two water mass scenarios: TBW and TWW. In this section, we review more than three decades of previous studies and indicate their main findings in Table 1, so that we can discuss them thoroughly and identify common patterns observed in the past.

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  - In summary, existing observations listed in Table 1 support that the phytoplankton community in the <u>BS</u> responds to a variety of factors which may vary from year to year, thus introducing high interannual variability in the phytoplankton assemblage. The factors primarily driving the nature of the chl-a bloom are: (1) vertical stability of the water column; (2) the depth of the UML, which influences the penetration of light into the depth range where biomass may accumulate near the
- 970 surface; (3) the existence of sea-ice retreat, supplying relatively cold freshwater to the environment; and, (4) the grazing pressure of herbivorous zooplankton. Berdalet *et al.* (1997) already accounted for these four factors and reported that the combination of those appear to play a major role in the development, accumulation and spatial variability of microplankton biomass. After reviewing the most recent studies, we confirm this statement still holds in water and applies also to at least the nanophytoplankton size (there is a scarcity of works investigating the picophytoplankton along cross-strait transects in
- 775 the BS so that we cannot extend here the statement robustly to this phytoplankton size). Interestingly, these physical factors may also condition the phytoplankton succession through a given bloom season and, thus, small cells appear to dominate the phytoplankton community structure during spring as large cells develop to form blooms in summer months (Petrou *et al.*, 2016).
- 780 In this context, it is worthwhile highlighting the results from two studies employing multi-year datasets of *in situ* observations of phytoplankton assemblage in the <u>BS</u> through four (Gonçalves-Araujo *et al.*, 2015) and nine (Mendes *et al.*, 2023) different bloom seasons, respectively.

On the one hand, in the first study, Gonçalves-Araujo *et al.* (2015) investigated microplankton (20-200 µm) and
nanoplankton (2-20 µm) through summertime of 2003, 2004, 2008 and 2009, identifying three main taxonomic groups within the study area: diatoms, flagellates and cryptophytes. From year to year, the authors show that the surface distribution of phytoplankton size was dominated by nanoplankton in 2003, 2004 and 2008 (>80% of the total chl-a) with no clear cross-strait gradient. Differently, in 2009 the surface distribution of chl-a presented two distinct domains: (1) in the TBW pool, a mixed community of microplankton and nanoplankton at high (~50-70%) and low (~30-50%) percentages of the total chl-a,
respectively; and, (2) in the TWW pool, a reversed mixed community of nanoplankton and microplankton at high (~80%) and low (~20%) percentages of the total chl-a, respectively. Regarding the taxonomic groups, the authors found that interannual variability in species composition resulted from an alternation between diatom-dominated and flagellate-dominated assemblages: 2003 and 2004 were dominated by cryptophytes nearly everywhere in the <u>BS</u>, 2008 by flagellates; and, 2009 by a mixture of diatoms close to the SSI and flagellates close to the AP (Gonçalves-Araujo *et al.*, 2015).

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Interestingly, in the second study, Mendes *et al.* (2023) investigate a subsequent period of time (2008-2018) based on measurements from nine different years (2011 and 2012 are absent) and demonstrated a strong coupling between biomass accumulation of cryptophytes, summer upper ocean stability, and the mixed layer. Through 2008-2018, Mendes *et al.* (2023)

- 805 report that <u>cryptophytes</u> present a competitive advantage in environments with significant light level fluctuations, normally found in confined stratified upper layers, and supported that observational finding with laboratory experiments where <u>cryptophytes</u> revealed a high flexibility to grow in different light conditions driven by a fast photo-regulating response. These results provided the basis to understand why the environmental conditions promoted the success of cryptophytes in coastal regions, particularly in shallower mixed layers associated with lower diatom biomass, and highlighted a distinct
- 810 competition or niche separation between diatoms and cryptophytes. Over the long-term variability, Mendes *et al.* (2023) concluded that <u>cryptophytes</u> are gradually outgrowing diatoms along with a decreased size spectrum of the phytoplankton community. This is in agreement with recent results supporting that the increasing melt-water input in the <u>BS</u> can potentially increase the spatial and temporal extent of cryptophytes (Mukhanov *et al.*, 2021), which benefit from the higher stabilization of the water column driven by the freshwater input.

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This reported shift towards a higher abundance of <u>cryptophytes</u> over diatoms is not trivial and, if persists in time, it will eventually impact the biogeochemical cycling in Antarctic coastal waters due to a shift in trophic processes (Mukhanov *et al.*, 2021). The latter work poses the scenario as follows. The replacement of large diatoms with small cryptophytes <u>favours</u> consumers like salps over Antarctic krill. Salps, a food competitor of Antarctic krill, can feed a wide range of taxonomic and size compositions of phytoplankton prey. Thus, salps present a much lower feeding selectivity (Haberman *et al.*, 2003) than Antarctic krill, which presents positive selectivity for diatoms (large prey) and avoid cryptophytes (smaller prey) when feeding on complex prey mixture (Haberman *et al.*, 2003). The shift towards an increasing role of cryptophytes in <u>BS</u> waters would then lead to constraints in food supply for krill, strengthening the abundance of its competitor. This would not only threaten Antarctic krill populations, but also higher consumers including penguins, seals, and whales, which feed on krill

825 (Loeb et al., 1997).

Based on the above review, we find that the biophysical coupling between the chl-a blooms at both sides of the <u>PF</u> is largely the result of interannually varying physical properties determined by the TBW and TWW pools, and that some of those physical properties could be easily monitored via remotely-sensed observations such as: (1) SST to control the extent of the 830 TBW and TWW pools; and, (2) SIC to monitor the sea-ice budget and sea-ice retreat as source of vertical stability to the water column. Through the last section of this study, we attempt to highlight that monitoring the spatio-temporal distribution of the chl-a blooms in the <u>BS</u> according to satellite measurements of SST and chl-a may represent a pivotal knowledge in future studies about the potential causes driving the long-term variability of the phytoplankton assemblage across the <u>PF</u>. 

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Reference	Methodology	PFTs in TBW	PFTs in TWW	PFTs in Bransfield	Date	
				Strait		
Mura et al., 1995	Fluorometric method and	Highest relative	Highest relative	The highest abundance	1993 (summer).	
	microscopy analysis.	contribution to	contribution to	across the PF was		
		community biomass by	community biomass	attributed to CPs.		
		eukaryotic picoplankton	by eukaryotic			Deleted: eukariotic
		and DTs.	picoplankton and DTs.			Deleted: enkariotic
Berdalet et al.,	Fluorometric and	The highest values of	The lowest values of	The degree of	1994. January	Deleted. curanolic
1997	biochemical methods to	MP biomass indicators	MP biomass indicators	stabilization of the water	(summer)	
	determine microplankton	(chl-a, ATP and protein)	(chl-a, ATP and	column, the depth of the		
	biomass.	were found in ice-	protein) were found in	UML and the grazing		
		melting waters and	TWW.	pressure of herbivorous		
		TBW.		zooplankton play a		
				major role in the		
				development.		
				accumulation and spatial		
				variability of MP		
				biomass		
				oroniuss.		
García-Muñoz et	Flow cytometry,	Highest abundance of	Higher abundance of	High abundance of NP	2010, January	
al., 2013	FlowCAM,	CPs and relatively high	NP (large size) and	(medium size) across the	(summer)	
	HPLC/CHEMTAX	abundance of NP (large	lower abundance of	PF.		
	pigment analysis.	size).	NP (small size).			
Mendes et al.,	HPLC, CHEMTAX,			Dominance of DTs in	2008-2009 (late	
2013	microscopy analysis.			deeper UML, higher	summer).	
				salinity and warmer SST.		
	HPLC, CHEMTAX,			Dominance of CPs in	2010 (late	
	microscopy analysis.			shallower UML, less	summer).	
				salinity and colder SST		
				(cold summer with late		
				ice retreat). Low diatom		Deleted: I
				biomass in presence of		
				high nutrient		
				concentrations		
				(particularly silicate) and		
				low chl-a.		
Gonçalves-Araujo	Either fluorometric or	Dominance of	Dominance of	Interannual variability of	2003, 2004, 2008,	
et al., 2015	spectrofluorometric	microplanktonic DTs	nanoplanktonic	chl-a bloom is governed	2009	
1	method, and microscopy	associated with higher	flagellates (CPs, HPs)	by alternation between	(summer).	Deleted: to

	analysis.	chl-a in shallower UML.	associated with lower	diatom-dominated		Deleted: to
			chl-a in deeper UML.	and flagellate-dominated		
				assemblages.		
Mukhanov et al.,	Flow cytometry,	Presence of CPs (9µm)	CPs were scarce or		2020, January	
2021	fluorescence.	and other NP (<3µm).	undetectable.		(summer).	
		Highest CP abundance				
		and biomass are found in				
		the photic layer around				
		the jet of the Bransfield				
		Current.				
Costa et al., 2023	HPLC, CHEMTAX,			Equivalent proportion	2013-2014, 2014-	
	microscopy analysis.			and abundance of	2015, November	
				smaller nanoflagellates	(spring) .	
				(CPs, DNs, P. antarctica		
				and green flagellates),		
				and centric and pennate		
				DTs. CPs prefer low		
				salinities, and centric		
				DTs prefer higher		
				salinities (>34). DNs and		
				centric DTs prefer		
				deeper UML.		
	HPLC, CHEMTAX,			Low diatom biomass	2013-2014,_2014-	
	microscopy analysis.			accumulation. Higher	2015,	
				proportion of CPs, DNs	(spring/summer).	
				and/or pennate DTs with		
				background presence of		
				mixed flagellates. CPs		
				and pennate DTs prefer		
				shallow UML <sub>2</sub> but CPs		
				occupy colder waters		
	UDI C CHENTAY			than pennate D1s.	2015 2017	
	nple, CHEMTAX,			riign diatom biomass	2015-2016,	
	meroscopy analysis.			by contria DTc	(spring/summer).	
Mandas at al	HDLC CHEMTAX SEM			CPc are gradually	2008 2018	
2023	DNA sequencing and			outgrowing DTs slong	2000-2018	Formatted: Don't adjust space between Latin and Asian text, Don't adjust space between Asian text and numbers
2023	phylogenetic inference			with a decreased size	(suillilei).	Deleted: 1
	phylogenetic interence.			spectrum of the		
				phytoplankton		
				community.		Deleted:
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Table 1. Summary review of studies investigating the chl<u>orophyll</u>-a bloom in the Bransfield Strait and reporting a description of the phytoplankton assemblage either by water mass domain (TBW or TWW) or without distinction. We must note that in none of the studies the full spectrum of Phytoplankton Functional Types (PFTs) is covered, and so this review attempts to provide a general overview of the existing knowledge. Acronyms for PFTs sizes are as follows: microphytoplankton (MP; 20-200 µm), nanophytoplankton (NP; 2-20 µm), picophytoplankton (PP; 0.2-2 µm). Other acronyms for PFTs are: diatoms (DTs), cryptophytes (CPs), haptophytes (HPs), dinoflagellates (DNs). Lastly, acronyms for methodology are: High Performance Liquid Chromatography (HPLC), Chemical taxonomy (CHEMTAX) software v1.95 (Mackey *et al.*, 1996), Scanning Electron Microscopy (SEM).

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### 3.5 Monthly variations of SST and chl-a along the CIEMAR and COUPLING transects

As a closure to our analyses we return to the synoptic transects which motivated our working hypothesis, based on *in situ* hydrographic and fluorescence measurements (T-I, T-II and T-III; Figures 2-4), and construct spatio-temporal climatologies of remotely-sensed SST and chl-a along the same transects (a series of black dots denote for reference the spatio-temporal position of the hydrographic stations along the Hovmöller diagrams in Figure 10). The aim is to highlight that the monthly

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position of the hydrographic stations along the Hovmöller diagrams in Figure 10). The aim is to highlight that the monthly variability of the easternmost extent of the chl-a bloom in the TBW pool and the westernmost extent of the chl-a bloom in the TWW pool responds closely to the monthly variability of the <u>PF</u>. We think this approach supports further the potential of long-monitoring the observed biophysical coupling via remotely-sensed measurements when study boxes are properly placed according to governing ocean dynamics.

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From early spring (October) to early autumn (April), the <u>PF</u> emerges prominently along transects T-II and T-III (Figure 10a), where relatively warm TBW, richer in chl-a along the southern shelf of the SSI (SST >  $1.4^{\circ}$ C; chl-a ~ 0.7-0.8 mg m<sup>-3</sup>) opposes relatively colder TWW and poorer in chl-a (SST ~  $-0.2^{\circ}$  to  $-0.6^{\circ}$ C; and chl-a < 0.3-0.4 mg m<sup>-3</sup>) along the western shelf of the <u>AP</u> (Figure 10b).

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It is worthwhile noting that the PF delineated along the synoptic transect T-II, found between stations 9-10 (Figure 3), corresponds closely to the climatological location of the PF (0.6°C and 0.5 mg m<sup>-3</sup>) observed between stations 8-9 (Figure 10a). Similarly, along T-III, both the hydrographic and the climatological PF are found at the same position, between stations T10 and T11.

#### 880

As it occurred along the synoptic transect T-I (Figure 2), the <u>PF</u> is not visible along the climatological transect T-I (Figure 10a), where relatively warm TBW invades the strait, and the TWW signal is absent from early spring (October) to early autumn (April) with SST values ~  $0.2^{\circ}$ C. The absence of a strong cross-strait temperature gradient along T-I is in agreement with an elongated patch of high chl-a concentrations which expands towards the western shelf of the AP, reaching values ~

885 0.5 mg m<sup>-3</sup> as far east as 84 km offshore the SSI (Figure 10b). This is analogous to the basin-wide, high fluorescence signal shown along the synoptic transect T-I in Figure 2.

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Throughout the remainder of the year, both SST and chl-a values follow similar patterns along the three climatological transects (T-I, T-II and T-III), displaying basin-wide, lower and more homogeneous values.

895 Notably, the highest chl-a concentrations are always found offshore along the three climatological transects, embedded in patches of warmest TBW (SST > 1.2°-1.4°C; chl-a ~ 0.6-0.8 mg m<sup>-3</sup>). These climatological transects (Figure 10b) confirm an earlier suggestion based on Figure 9, where the northward spatio-temporal migration of the chl-a bloom is apparent. Here we note that the highest chl-a concentrations along the three climatological transects occur around December in T-I, through December to February in T-II and around February in T-III.

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In summary, the remotely-sensed observations of SST and chl-a concentrations have proven to be of great potential to monitor major features of the chl-a blooms in the <u>BS</u> accounting for a biophysical coupling between two hydrographic scenarios (TBW and TWW pools) confronted along the <u>PF</u>. Importantly, we recall these two hydrographic scenarios embed different phytoplankton assemblages, as it has been discussed based on previous literature, and results from this study.





Figure 10. Monthly climatology from 1998 to 2018 of a) Sea Surface Temperature (SST), and b) chlorophyll-a (chl-a) concentration for each study transect (see Figures 2-4) from the South Shetland Islands (SSI) to the Antarctic Peninsula (AP). The black markers (dots) situated at the top of the subplots represent the station's positions along the transects. These same stations are also displayed during the summer months when the cruises were carried out. Additionally, the position of the Peninsula Front(PF), as identified during the cruises and located along the isopycnal of 27.55 kg m<sup>-3</sup>, is also indicated.

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Lastly, it is worthwhile noting that the alignment of the chl-a spatial distribution along an oceanic front is not a novel feature in the world's oceans, and has been already investigated in the literature (Moore and Abbott, 2002; Baird *et al.*, 2008; Von

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Bodungen et al., 2008). Thus, the novelty of our work lies in demonstrating through in situ observations and remotely-sensed measurements that such a biophysical coupling has the potential to be used to monitor the chl-a blooms and phytoplankton assemblages occurring seasonally in BS. This aspect is particularly relevant because BS is a key region for the sustainability

920 of marine Antarctic ecosystems, which is challenging to monitor due to the hazardous prevailing conditions in polar regions. In future studies, we expect the calculation of the frontal probability (Yang et al., 2023) of the PF through a multi-year timeseries of SST data may be beneficial to assess and co-locate interannually the alignment of the thermal front and the chl-a bloom domains using an automated algorithm for the Bransfield Strait study case (see Appendix B for further insights).

#### 925 4 Conclusions

In this study, we address the hypothesis that the spring-to-summertime biophysical coupling controlling the chl-a bloom in the BS could be monitored through a combination of remotely-sensed observations of chl-a and SST, which strongly conditions the spatio-temporal variability of the phytoplankton assemblage across the PF. Our approach is based on the characterisation of climatological fields, following the motivation from novel and historical synoptic in situ observations 930 (discussed in Section 3.1) which reveal that the PF may be used as a guideline to contour two distinctive niches for phytoplankton assemblage in the BS, both horizontally and vertically.

Through Section 3.2, we conclude that the surface distribution of the seasonal variation of the SST in the BS enables the identification of two environmentally different scenarios for the phytoplankton, which then grow under different strategies following the revised literature.

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The first scenario is the pool of Transitional Bellingshausen Water, relatively warmer and less saline waters in a stratified water column with shallow mixed layers as compared to the second scenario. The second scenario is the pool of Transitional Weddell Water, relatively colder and more saline waters in a well-mixed water column with deeper mixed layers. We identify that the climatological isotherm characterising the PF location at surface during the summer months corresponds to

940 0.6°C isotherm, which divides the BS in two domains. This division is further supported when we show that the isoline of 0.5 mg m<sup>-3</sup> concentration of chl-a aligns with the 0.6°C isotherm, which works well as a threshold contouring the chl-a bloom around the SSI and embedding coherently in space the region with highest chl-a values during the summer months.

Following the seasonal climatology of the SIC, we conclude that the larger freshwater inputs reported in the literature over 945 the TBW domain and contributing to the vertical stabilization of the water column might be driven by a warmer oceanic forcing over coastal/glacial areas of the SSI (Cook et al., 2016; Saenz et al., 2023) rather than by melting of the open ocean

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sea-ice. On the other hand, the seasonal climatology of the wind stress forcing suggests that the westerlies may play a major role in contributing to (1) stratified waters on the TBW domain via downwelling-favourable Ekman transport along the southern SSI shelf, through which the Bransfield Current flows; and (2) well-mixed upper layers on the TWW domain via upwelling-favourable Ekman transport along the west AP shelf.

- Through Section 3.3, and based on *ad hoc* study boxes located according to the spatial distribution of the remotely-sensed of chl-a concentrations, we conclude that two different climatological chl-a blooms developing simultaneously but of different *nature* (i. e. phytoplankton assemblage) can be identified in the <u>BS</u>, as suggested by their different intensity and timing (month of initiation and rate of increase). This is in agreement with former results in a series of studies which reported that <u>cryptophytes</u> compete in the <u>BS</u> primarily with diatoms and other nanophytoplanton groups (Mura *et al.*, 1995; García *et al.*, 2013; Mendes *et al.*, 2013; Gonçalves-Araujo *et al.*, 2015; Mukhanov *et al.*, 2021; Costa *et al.*, 2023; Mendes *et al.*, 2023),
- 965 following strategies to adapt better to the physical environment present through that year, and which display differently by zones in the monthly climatologies of SST, air temperature, SIC and wind stress forcing. Generally speaking, these studies have reported that TBW chl-a concentrations are commonly characterised by <u>cryptophytes</u> and small diatoms, while TWW chl-a concentrations are more frequently characterised by large diatoms.
- 970 In Section 3.4, we revised our results against the literature about phytoplankton assemblage in the BS, and conclude that the biophysical coupling between the chl-a blooms at both sides of the PF is largely the result of interannually varying physical properties determined by the TBW and TWW pools. This suggests that the combined analysis of remotely-sensed observations of chl-a and SST (as presented in this study) may be of help in elucidating the spatio-temporal variability of the two blooms occurring in the BS during the summer months from year to year. Nevertheless, we must note that a given uncertainty will still exist about knowing which phytoplankton assemblage products are further validated in the future. We have explored such products (not shown), but the lack of a product detecting only cryptophytes hampers the assessment of their year-to-year competition with diatoms (a product which actually exists) in the BS. We find this is of paramount importance for a more comprehensive understanding of the marine ecosystem composition in the BS.

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Lastly, through Section 3.5 we conclude that combined analyses of remotely-sensed observations of SST and chl-a concentrations have a great potential to capture major features of the chl-a blooms in the <u>BS</u> accounting for a biophysical coupling between two hydrographic scenarios (TBW and TWW pools) confronted along and across the <u>PF</u>. We think that these results highlight the importance of long-term monitoring of the spatio-temporal distribution of the chl-a blooms in the BS using satellite measurements of SST and chl-a. Such monitoring may prove pivotal for future studies investigating the

forcings driving the long-term variability of the phytoplankton assemblage in the BS,

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#### Appendix A: Assessment of the SST products

We assess the goodness of three available SST products by comparison with near-surface (0-1 m and 10 m depth) *in situ* temperature measurements collected from a variety of Antarctic cruises (Table <u>A</u>1). By linear regression, the coefficient of determination (R-squared) is used to evaluate the performance of the three satellite products, which are: (1) Optimally
 Interpolated SST (OI SST; <u>https://www.remss.com/</u>); (2) European Space Agency Climate Change Initiative (ESACCI; <u>https://marine.copernicus.eu/</u>); and, (3) Operational Sea Surface Temperature and Ice Analysis (OSTIA; <u>https://marine.copernicus.eu/</u>). For brevity, hereafter, we refer to the, as OI SST, ESACCI and OSTIA, respectively.

The grid spacing for OI SST is ~0.1°, but for both ESACCI and OSTIA is 0.05°. Meanwhile, temporal extent is also 1315 different, OI SST time range is from 01 June 2002 to present, ESACCI time range is from 01 September 1981 to 31 December 2016, and OSTIA time range is from 01 October 1981 to 31 May 2022.

For a fair comparison, because the OI SST product starts globally in 2002, we first compare the three products to cruises GOAL (GOAL03, GOAL04 and GOAL05) and BREDDIES (2003), at two different depths (0-1 m and 10 m; see Table A2 to learn about the amount of profiles by depth level used in the analysis, and Table A3 to learn about the results). Based on the low coefficients found for OI SST as compared to ESACCI and OSTIA (Table A3), we decide to discard OI SST for further analysis.

- Lastly, we use the entire dataset of available hydrographic observations (Table <u>A</u>1) in the study region, making a distinction between whether we use only data falling within Bransfield Strait (BS), Gerlache Strait (GS) or within both (<u>full domain</u>). These data are summarized in Table <u>A</u>4, where there are indications to the depth levels involved in the analysis: 0-1 m (8 cruises with 539 stations from 1990 to 2010) and 10 m (21 cruises with 1,133 stations from 1990 to 2011).
- Results in Table  $\Delta 5$  show the lowest coefficients are found in Gerlache Strait for both ESACCI and OSTIA, while these values increase when including measurements from Bransfield Strait. To some extent, this agrees with the expectation provided the narrow nature of Gerlache Strait (~10 km at its narrowest part and ~50 km at its widest part). This implies the ocean in Gerlache Strait is in close proximity to land nearly everywhere, leading the discrepancies between remotely-sensed and *in situ* observations (Zhang *et al.*, 2004; Xie *et al.*, 2008; Lee and Park, 2021).
- 335 Notably, *in situ* measurements at 10 m present a higher correlation with satellite SST everywhere (Table A5). Given the similarity of correlation coefficients for ESACCI and OSTIA, we select OSTIA given its longer time record, which is from 1981 to 2020 as opposed to ESACCI, which ends earlier in 2016 at the moment this analysis was performed in 2022.

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	CRUISE	DATES			
		Year	Month	Day Season	
	M11_4 (*)	1990	Jan	01-10, 16	Summer
1345	ANT-XII (PS33)	1994	Nov	26-30	Spring
			Dec	01-05	
	FRUELA (*)	1995	Dec	03-23, 26-31	Spring
		1996	Jan	02-05, 18-31	Summer
			Feb	01-05	
1350	ALMIRANTE IRIZAR	1996	May	08-09, 12-14	Autumn
	ANT-XIV (PS42)	1996	Nov	15-28, 30	Spring
			Dec	01, 03-06, 08-24	
	DOVETAIL	1997	Aug	04-23, 25-31	Winter
			Sep	01-05	
1355	ANT-XV (PS49)	1998	Mar	31	Summer
			Apr	01-06, 17-22	Autumn
	ALBATROSS (*)	1999	Mar	23-26, 31	Summer
			Apr	01-02	Autumn
	CIEMAR	1999	Dec	15, 18-30	Spring
1360	BREDDIES	2002	Dec	30-31	Spring
		2003	Jan	02-06, 11-14, 17-18, 21	Summer
	GOAL03 (*)	2003	Jan	23-27, 29, 31	Summer
			Feb	21-23	
	GOAL04 (*)	2004	Jan	18-29	Summer
1365	GOAL05 (*)	2005	Jan	19, 21, 24-26, 28-31	Summer
			Feb	01-05, 07	
	ANT-XXII (PS67)	2005	Mar	12-13, 15-19, 24, 31	Summer
			Apr	01-02	Autumn
	SOS1 (*)	2008	Feb	21-29	Summer
1370			Mar	01-04	
	ANT-XXIV (PS71)	2008	Mar	25-31	Summer
			Apr	01-05	Autumn
	JC-031	2009	Feb	13-14, 16-18, 20-21	Summer
	SOS2	2009	Feb	17-28	Summer
1375			Mar	01	
	COUPLING	2010	Jan	08-26	Summer
	SOS3 (*)	2010	Feb	16-24	Summer
	ANT-XXVI (PS77)	2011	Jan	05-17	Summer
				1	1

1380 Table A1. Overview of the cruises used to calculate the coefficient of determination (see Section 2) and the dates they were carried out. Only cruises marked with an asterisk (\*) provide data at depths of 0-1 m.

Cruise	Depth (m)	Profiles
GOAL03-04-05	0-1	190
	10	205
BREDDIES	10	61

Table A2. Number of profiles available in each cruise and depth used to analyse the goodness of available open-access remotely-sensed products of sea surface temperature.

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Cruise	Depth (m)			
		OISST	ESACCI	OSTIA
	0-1	0.360	0.704	0.707
GOAL03-04-05 (BS	10	0.258	0.574	0.512
and GS)				
BREDDIES (BS)	10	0.299	0.785	0.773

Table A3. R-squared coefficients for each SST satellite product (OI SST, ESACCI, OSTIA) compared to *in situ* SST data from four cruises' data at 0-1 m and 10 m depths (see amount of profiles in Table A2). The analysis is performed for different study regions: at the entire study region (BS and GS) and the 1395 Bransfield Strait (BS) surroundings (excluding the GS region).

	Depth (m)	Profiles
	0-1	539
BS and GS	10	1133
	0-1	417
Gerlache Strait	10	905
	0-1	122
Bransfield Strait	10	228

1400 Table A4. Same as Table A2, but here extended to eight cruises for depths of 0-1 m and twenty-one cruises for depths of 10 m. Regarding to Gerlache Strait region, there are only five cruises available for depths of 0-1 m (ALBATROSS, FRUELA, GOAL05, M11\_4) and ten cruises for depths of 10 m (ALBATROSS, FRUELA, GOAL03, GOAL04, GOAL05, M11\_4, ALMIRANTE IRIZAR, ANT\_XXVII\_PS77, CIEMAR, JC-031).

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Study Region	Depth (m)		
		ESACCI	OSTIA
	0-1	0.715	0.719
BS and GS	10	0.787	0.784
	0-1	0.431	0.487
Gerlache Strait	10	0.515	0.546
	0-1	0.815	0.787
Bransfield Strait	10	0.857	0.849

Table A5. R-squared coefficients for each SST satellite product (ESACCI, OSTIA) compared to in situ SST data from eight cruises' data at depths of 0-1 m

and twenty-one cruises' data at depths of 10 m (see amount of profiles in Table A4). The analysis is performed for different study regions: at the entire study region (BS and GS), only the Gerlache Strait (GS), and the Bransfield Strait (BS) surroundings (excluding the GS region)

# Appendix B: Frontal probability of the Peninsula Front



415 Figure B1. From left to right, the upper panels (a,b,c) show the Frontal Probability (FP) based on daily, monthly-averaged and seasonally-averaged data for sea surface temperature through 21 years of summertime. Lower panels (d,e,f) show the same as upper panels but based on chlorophyll-a (chl-a) concentrations. The climatological summertime isotherm of 0.6°C (dashed red line) and the isoline of 0.5 mg m<sup>-3</sup> chl-a concentrations (solid black line) as

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obtained for Figures 5 and 7 highlight the goodness of our methodology to select them as characteristic environmental values contouring the Peninsula Front 420 in Bransfield Strait.

Figure B1 presents the Frontal Probability (FP; Yang *et al.*, 2023) from the SST and chl-a fields in Bransfield Strait for the period 1998-2018. The Canny edge-detection algorithm (Canny, 1986) is applied to identify coherent frontal segments.

425 Then, the summertime FP is calculated based on three different cases, using fronts detected on daily data, monthly-averaged data, and seasonally-averaged data over a period of 21 years (in all cases the information corresponds solely to summertime). The FP is defined at each pixel as a percentage where the times that the pixel is identified as a front is referred to the number of total valid pixels for a given time interval.

Results support the choice of the characteristic isotherms and isoline of chl-a used in this study to distinguish in the BS two
 different pools of chl-a development. Additionally, we note that the signal of the Peninsula Front increases in FP, especially in SST, when based on time-averaged fields (panels b,c,e,f) as compared to daily fields (a,d). We attribute this to the recurrence of the Peninsula Front, which gets better defined when a time-average procedure is followed before applying the Canny edge-detection algorithm. Simultaneously, a noisier signal emerges regarding other non-recurrent fronts which are present only occasionally in time-average fields, thus leading to their presence only in a few fields when computing the FP.

We suggest that the FP may be used in future studies to code an automated algorithm capable of monitoring the chl-a blooms in Bransfield Strait based on remotely-sensed SST and chl-a data, using the South Shetland Islands and the Antarctic Peninsula as physical boundaries, and the Peninsula Front location as the oceanographic frontier contouring the TBW and TWW pools. Thus, co-locating interannually the alignment of the thermal front and the chl-a spatial distribution will enable the computation of accurate areas of integration for the assessment of the surface blooms acting in the Bransfield Strait.

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