

Drivers of Phytoplankton Bloom Interannual Variability in the Amundsen and Pine Island Polynyas

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Abstract

The Amundsen Sea Embayment (ASE) experiences both the highest ice shelf melt rates and the highest biological productivity in West Antarctica. Using 19 years of satellite data and modelling output, we investigate the long-term influence of environmental factors on the phytoplankton bloom in the Amundsen Sea (ASP) and Pine Island (PIP) polynyas. We test the prevailing hypothesis that changes in ice shelf melt rate could drive interannual variability in the polynyas' surface chlorophyll-*a* (chl*a*) and Net Primary Productivity (NPP). We find that the interannual variability and long-term change in glacial meltwater may play an important role in chl*a* variance in the ASP, but not for NPP. Glacial meltwater does not explain the variability in neither chl*a* or NPP in the PIP, where light and temperature are the main drivers. We attribute this to potentially greater amount of iron-enriched meltwater brought to the surface by the meltwater pump

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downstream of the PIP, and the coastal ocean circulation accumulating and transporting iron towards the ASP.

Short Summary

We investigate the phytoplankton bloom variability and its drivers in the Amundsen polynyas (areas of open water within sea ice). Between 1998 and 2017, we find that changes in melting ice shelves may have different impacts on biological productivity between the Amundsen Sea (ASP) and Pine Island (PIP) polynyas. While ice shelves melting seems to play an important role for phytoplankton growth variability in the ASP, light and warmer waters appear to be more important in the PIP.

1. Introduction

Coastal polynyas are open ocean areas formed by strong katabatic winds pushing sea ice offshore (Morales Maqueda, 2004). They are the most biologically productive areas in the Southern Ocean (SO) relative to their size (Arrigo et al., 1998). This high biological productivity contrasts sharply with the rest of the SO, where low iron and light availability generally co-limit phytoplankton growth (Boyd et al., 2007). In West Antarctica, the Amundsen Sea Embayment (ASE) hosts two of the most productive Antarctic polynyas: The Pine Island Polynya (PIP) and Amundsen Sea Polynya (ASP) (Arrigo and van Dijken, 2003).

The phytoplankton community in the ASE is generally dominated by *Phaeocystis antarctica* (Lee et al., 2017; Yager et al., 2016), which is adapted to low iron availability and variable light conditions, and forms large summer blooms (Alderkamp et al., 2012; Yager et al., 2016). Diatoms like *Fragilariopsis sp.* and *Chaetoceros sp.* are also present, often becoming more important near the sea-ice edge or under shallow, stratified mixed layers where silicic acid (Si) and iron (Fe) are more available (Mills et al., 2012). In exceptional years, such as 2020, diatoms like *Dactyliosolen tenuijunctus* replaced *P. antarctica* as the dominant taxon, driven by anomalously shallow mixed layers and sufficient Fe–Si supply (Lee et al., 2022). This dynamic balance highlights how light, nutrient supply, and stratification control community composition in these highly productive and complex Antarctic systems.

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65 The ASE is also the Antarctic region experiencing the highest mass loss from the Antarctic ice
66 sheet. It has been undergoing increased calving, melting, thinning and retreat over the past three
67 decades (Paolo et al., 2015; Rignot et al., 2013; Rignot et al., 2019; Shepherd et al., 2018). In the
68 ASE, this ice loss is mainly through enhanced basal melting of the ice shelves. This is attributed
69 to an increase in wind-driven Circumpolar Deep Water (CDW) fluxes and ocean heat content
70 intruding onto the continental shelf through deep troughs such as the Pine Island and Dotson-
71 Getz, and flowing into the ice shelves cavities (Dotto et al., 2019; Jacobs et al., 2011; Pritchard et
72 al., 2012). There, warm waters fuel intense basal melt of the Pine Island, Thwaites, and Getz ice
73 shelves, and returns as a fresher, colder outflow that can strengthen stratification (Jenkins et al.,
74 2010; Ha et al., 2014). The PIP and ASP differ in their exposure to CDW and in local
75 circulation: the ASP is more strongly influenced by upwelled modified CDW (mCDW) and
76 glacial meltwater inputs, whereas in the PIP, the deep mCDW retains more of its original
77 offshore characteristics, with vertical exchange only significantly occurring beneath the ice
78 shelves, leading to a more stratified and less directly ventilated surface layer (Assmann et al.,
79 2013; Dutrieux et al., 2014). These hydrographic contrasts can shape the timing and magnitude
80 of phytoplankton blooms and nutrient dynamics across the two polynyas.

81

82 Melting ice shelves can explain about 60% of the biomass variance between all Antarctic
83 polynyas, suggesting that they are the primary supplier of dissolved iron (dFe) to coastal
84 polynyas (Arrigo et al., 2015), and can directly or indirectly contribute to regional marine
85 productivity (Bhatia et al., 2013; Gerringa et al., 2012; Hawkins et al., 2014; Herraiz-
86 Borreguero et al., 2016). The strong melting of the ice shelves can release significant quantities
87 of freshwater at depth (Biddle et al., 2017), resulting in a strong overturning within the ice
88 shelves cavity, called the meltwater pump (St-Laurent et al., 2017). Modelling efforts have
89 identified both resuspended Fe-enriched sediments and CDW entrained to the surface by the
90 meltwater pump as the two primary sources of dFe to coastal polynyas, providing up to 31% of
91 the total dFe, compared to 6% for direct ice shelves input (Dinniman et al., 2020; St-Laurent et
92 al., 2017). Other drivers such as sea-ice coverage (and associated increases in light and dFe
93 availability when sea ice retreats), or winds have also been shown to impact primary productivity
94 in polynyas (Park et al., 2019; Park et al., 2017; Vaillancourt et al., 2003).

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97 The key question of how glacial meltwater variability may impact biological productivity in the
98 ASE has previously been raised during the ASPIRE program (Yager et al., 2012). During the
99 expedition, a significant supply of melt-laden iron-enriched seawater to the central euphotic zone
100 of the ASP was observed, potentially explaining why this area is the most biologically
101 productive in Antarctica (Randall-Goodwin et al., 2015; Sherrell et al., 2015). Other studies in
102 the Western Antarctic Peninsula and East Antarctica showed that the meltwater pump process
103 was also responsible for natural Fe supply to the surface, increasing primary productivity (Cape
104 et al., 2019; Tamura et al., 2023).

105
106 In this study, we investigate the long-term relationship between the main environmental factors
107 of the ASE and the surface biological productivity, with a focus on ice shelves melting. A
108 demonstrated relationship between glacial meltwater and phytoplankton growth would have far-
109 reaching consequences for regional productivity in coastal Antarctica, and possibly offshore,
110 over the coming decades under expected climate change scenarios (Meredith et al., 2019). We
111 test the hypothesis that changes in glacial meltwater are linked to the surface ocean primary
112 productivity variability observed over the last two decades. We use a combination of satellite
113 (ocean color and ice shelf melting rate), climate re-analysis, and model data spanning 1998 to
114 2017.

115 116 2. Material and Methods

117 118 2.1 Study area and polynya mapping

119

120 We focus on the PIP and ASP in the ASE in West Antarctica (Fig. 1). The ASE is comprised of
121 several ice shelves and glaciers, including: Abbot (Abb), Cosgrove (Cs), Pine Island (PIG),
122 Thwaites (Tw), Crosson (Cr), Dotson (Dt) and Getz (Gt). The PIG and Thwaites have received
123 significant attention in recent years due to their potentially large contribution to sea level rise
124 (Rignot et al., 2019; Scambos et al., 2017). Along with the Crosson and Dotson ice shelves, the
125 PIG and Thwaites are undergoing the highest melt rate, which is expected to increase under
126 climate change scenarios (Naughten et al., 2023; Paolo et al., 2023). The polynyas' boundaries

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were determined using a 15% sea-ice concentration (SIC) mask (Moreau et al., 2015; Stammerjohn et al., 2008) for every 8-day period from June 1998 to June 2017 to accurately represent the size of the polynyas through time.

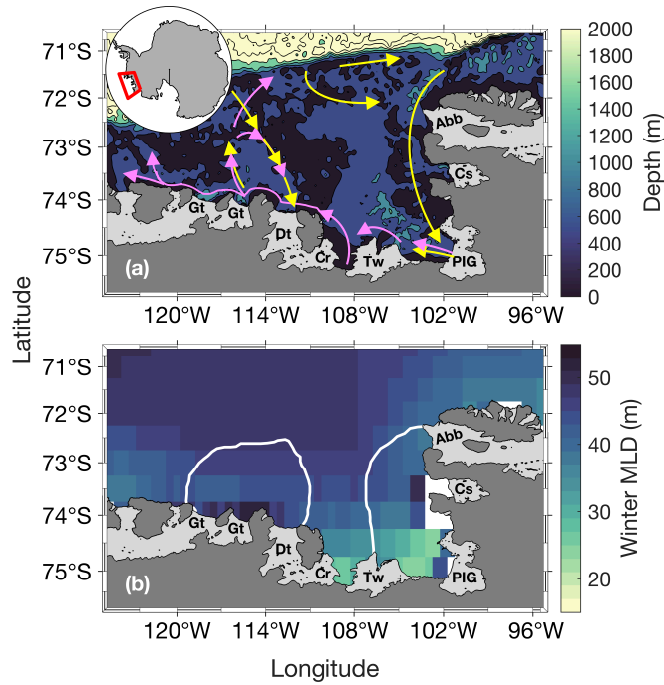


Fig. 1. Study area. Panel (a) shows the bathymetry (from ETOPO1; Amante & Eakins, 2009) and panel (b) shows the climatological April-September (that we call winter) mixed-layer depth (MLD) from 1998 to 2016 (n=114). Panel (a) shows a simplified schematic of the local deep ocean circulation (~ below 400m, yellow arrows) and upper glacial meltwater/sediments/circumpolar deep water sourced dFe pathways (magenta arrows), which follows the local upper ocean circulation. Schematic adapted from St-Laurent et al. (2017). The white lines in panel (b) represent the climatological summer polynyas' boundaries for the Amundsen (ASP; left) and Pine Island (PIP; right) polynyas. The dark grey area is mainland

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190 Antarctica. Light grey areas indicate floating ice shelves and glaciers: Abbot (Abb), Cosgrove
191 (Cs), Pine Island Glacier (PIG), Thwaites (Tw), Crosson (Cr), Dotson (Dt) and Getz (Gt).

193 2.2 Satellite ocean surface chlorophyll-*a* and net primary productivity

195 We obtained level-3 satellite surface chlorophyll-*a* concentration (chl_a) with spatial and
196 temporal resolution of 0.04° and 8 days from the European Space Agency (ESA) Globcolor
197 project. We used the CHL1-GSM (Garver-Siegel-Maritorena) (Maritorena and Siegel, 2005)
198 standard Case 1 water merged products consisting of the Sea-viewing Wide Field-of-view
199 (SeaWiFS), Medium Resolution Imaging Spectrometer (MERIS), Moderate Resolution Imaging
200 Spectroradiometer (MODIS-A) and Visible Infrared Imaging Suite sensors (VIIRS). We chose to
201 perform our analysis with the merged GlobColour product, which has been widely applied and
202 tested in Southern Ocean and coastal Antarctic studies (Ardyna et al., 2017; Sari El Dine et al.,
203 2025; Golder & Antoine, 2025; Nunes, Ferreira & Brito, 2025), to increase our spatial and
204 temporal coverage.

206 We estimated phytoplankton bloom phenology metrics following the Kauko et al. (2021)
207 method. Firstly, for a given 8-day period, we applied a spatial 3x3 pixels median filter to reduce
208 gaps in missing data. Then, if a pixel was still empty, we applied the average chl_a of the previous
209 and following week to fill the data gap. Data were smoothed using a 4-point moving median
210 (representing a month of data). For each pixel, the threshold for the bloom detection was based
211 on 1.05 times the annual median. The threshold method is frequently used (Racault et al., 2012;
212 Siegel et al., 2002) and proven reliable at higher latitudes (Marchese et al., 2017; Soppa et al.,
213 2016; Thomalla et al., 2023). We then determined 5 main bloom metrics. The bloom start (BS) is
214 defined as the day where chl_a first exceeds the threshold for at least 2 consecutive 8-day periods.
215 Conversely, the bloom end is the day where chl_a first falls below the threshold for at least 2
216 consecutive 8-day periods. The bloom duration (BD) is the time elapsed between bloom start and
217 bloom end. The bloom mean chl_a (BM) and bloom maximum chl_a are respectively the average
218 and maximum chl_a value calculated during the bloom. Each year is centered around austral
219 summer, from June 10th year *n* (day 1) to June 9th year *n*+1 (day 365 or 366). We also averaged
220 our 8-day data to monthly data to perform a spatial correlation analysis (see section 2.6).

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224 We note that satellite ocean-colour chl_a algorithms (including the GlobColour merged product
225 used here) are globally tuned and may underperform in optically complex waters (e.g., with
226 elevated dissolved organic matter or suspended sediments, ‘Case 2’). In the ASP, past work
227 (Park et al. 2017) showed that satellite chl_a climatologies reflect broad seasonal patterns that are
228 consistent with *in situ* measurements of phytoplankton biomass and photophysiology, but there is
229 limited data from regions immediately adjacent to glacier fronts or during times of strong
230 meltwater input. Thus, while we consider satellite chl_a to be useful for capturing spatial and
231 temporal variability at polynya scale, uncertainty likely increases in optically complex zones
232 near glacier margins or during low-light periods, and needs to be considered while interpreting
233 results.

234
235 Eight-day satellite derived Net Primary Productivity (NPP) data with 1/12° spatial resolution,
236 spanning 1998 - 2017 using the Vertically Generalized Production Model (Behrenfeld and
237 Falkowski, 1997) were obtained from the Oregon State University website. The VGPM model is
238 a chlorophyll-based approach and relies on the assumption that NPP is a function of chl_a,
239 influenced by light availability and maximum daily net primary production within the euphotic
240 zone. SeaWiFS-based NPP data span 1998 - 2009, MODIS-based data span 2002 - 2017. To
241 increase spatial and temporal coverage, we averaged SeaWiFS and MODIS from 2002 to 2009,
242 where there was valid data for both in a pixel. NPP data were also monthly averaged and used to
243 compare with chl_a spatial and temporal patterns.

244
245 We caution that our study focuses on surface productivity, and satellites cannot detect under-ice
246 phytoplankton, sea-ice algal blooms, or deeper productivity, therefore likely underestimating
247 total primary productivity (Ardyna et al., 2020; Boles et al., 2020; Douglas et al., 2024; McClish
248 & Bushinsky, 2023; Stoer & Fennel 2024).

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250 2.3 Ice shelves volume flux

251
252 We used the latest ice shelf basal melt rate estimates from Paolo et al (2023). These estimates are
253 derived from satellite radar altimetry measurements of ice shelves height, and produced on a 3

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258 km grid every 3 months, with an effective resolution of ~5 km. For this study, our basal melt
 259 record spans June 1998 to June 2017. We calculated ice shelves volume flux rate for every
 260 gridded cell by multiplying the basal melt rate by the cell area. Data were summed for each ice
 261 shelf for a 3-month period. A 5-point (15 months) running mean was applied to reduce noise,
 262 such as spurious effects induced by seasonality on radar measurements over icy surfaces (Paolo
 263 et al., 2016), and data were temporally averaged from October to March to match the SO
 264 phytoplankton growth season (Arrigo et al., 2015), providing yearly mean values. The Abbot,
 265 Cosgrove, Thwaites, [PIG](#), Crosson, Dotson and Getz ice shelves were used to calculate a single
 266 total meltwater volume flux (TVFall) for the ASE to investigate the link with surface chl_a and
 267 NPP. We also investigated the relationship between each polynyas' productivity and their closest
 268 ice shelf. The Abbot, Cosgrove, PIG and Thwaites ice shelves were used to calculate the flux
 269 rate in the PIP (TVFpip) while the Thwaites, Crosson, Dotson and Getz ice shelves were chosen
 270 for the ASP (TVFasp). The Thwaites was used in both due to its central position between the two
 271 polynyas. We thereafter use the term glacial meltwater which defines meltwater resulting from
 272 ice shelf melting.

274 2.4 Simulated dFe distribution

276 The spatial distribution of dFe from different sources in the embayment was investigated from
 277 Dinniman et al. (2020) model output. The model used is a Regional Ocean Modelling System
 278 (ROMS) model, with a 5 km horizontal resolution and 32 terrain following vertical layers and
 279 includes sea-ice dynamics, as well as mechanical and thermodynamic interaction between ice
 280 shelves and the ocean. The model time run spans seven years and simulates fourteen different
 281 tracers to understand dFe supply across the entire Antarctic coastal zone, with the last two years
 282 simulating biological uptake. For the purpose of this study, we only use four different dFe
 283 sources/tracers in the ASE: ice shelf melt, CDW, sediments and sea ice. Each tracer estimation is
 284 independent from each other, meaning that one source does not affect the other, and they have
 285 the same probability for biological uptake by phytoplankton. That is, dFe from all sources can
 286 equally be taken up by phytoplankton. This is parametrized in the model as all iron molecules
 287 being bound to a ligand and therefore remaining in solution in a bioavailable form (Gledhill &
 288 Buck, 2012). For a detailed and complete explanation of the model, see Dinniman et al. (2020).

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2.5 Other environmental parameters

We used SIC data spanning June 1998 to June 2017 from the National Snow and Ice Data Center (Cavalieri et al., 1996). The data are Nimbus-7 SMMR and SSMI/SSMIS passive microwave daily SIC with 25 km spatial resolution. We computed the sea-ice retreat time (IRT) and open water period (OWP) metrics using a 15% threshold (Stammerjohn et al., 2008). Daily data were monthly averaged to perform a spatial correlation analysis (see section 2.6).

We collected monthly level-4 Optimum Interpolation Sea Surface Temperature (OISST.v2) 0.25° high resolution dataset from the National Oceanic and Atmospheric Administration (Huang et al., 2021). Using this dataset compared to others has been proven to be the most suitable for our region of interest (Yu et al., 2023).

We obtained monthly Photosynthetically Available Radiation (PAR) from the same Globcolour project at the same spatial and temporal resolution (0.04° and 8 days) as chl_a.

We used monthly averaged ERA5 reanalysis of zonal (u) and meridional (v) surface wind speed at 10 m above the surface (Hersbach et al., 2020).

We investigated monthly mean MLD from the Estimating the Circulation and Climate of the Ocean (ECCO) ocean and sea-ice state estimate project (ECCO consortium et al., 2021). The dataset is the version 4, release 4, at 0.5° spatial resolution.

[Variability in the sea-ice landscape can be influenced by the Amundsen Sea Low \(ASL\) in West Antarctica \(Hosking et al., 2013; Turner et al., 2016\). We therefore finally looked at the impact of the ASL and its potential influence on sea-ice variability. Monthly ASL indices \(latitude, longitude, central and sector pressure\) derived from ERA5 reanalysis data were obtained from the ASL climate index page \(Hosking et al., 2016\).](#)

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2.6 Statistical analysis

Because some of our data were not normally distributed, we consistently applied nonparametric tests throughout our statistical analysis. A Mann-Kendall test was performed to detect linear trends in chl_a and NPP. A two-tailed non-parametric Spearman correlation metric (ρ , p) was calculated to investigate the relationship between chl_a, NPP, and environmental factors, as well as between the phytoplankton bloom and sea-ice phenology metrics. A two-tailed Mann-Whitney test was performed to detect any significant mean differences for chl_a, sea-ice phenology metrics, MLD, PAR and dFe sources between the two polynyas. Monthly spatial correlations were tested between SIC, winds, chl_a, NPP, SST, and PAR after removing the seasonality for each parameter. As well, a yearly spatial correlation between chl_a, NPP and TVFall was performed. The relationships between chl_a, NPP and environmental factors were explored using a Principal Component Analysis (PCA). No pre-treatment (mean-centering or normalization) was applied to the variables prior to PCA, as all variables are expressed in comparable units and ranges, consistent with common practice in marine biogeochemistry studies (Marchese et al., 2017; Liniger et al., 2020). The Spearman, Mann-Whitney and PCA analysis were conducted using the mean TVFs, MLD, SST, and PAR calculated over the October-March period for each year, with the associated bloom and sea-ice phenology metrics. Every statistical test was run with a 95% (p -value < 0.05) confidence level. Our study spans 1998-2017. We are constrained by the start of satellite ocean color data (1998) and the end of the ice shelf basal melt rate record (2017) from Paolo et al (2023).

3. Results

3.1 Glacial meltwater, chl_a, and NPP variability

The annual climatology maps reveal substantially higher chl_a and NPP in the ASP compared to the PIP (Fig. 2). Chl_a starts increasing in mid-November to reach its average peak earlier in the PIP than the ASP. At its peak, chl_a in the ASP is 6.49 mg m⁻³ and 4.94 mg m⁻³ in the PIP (Fig. 3a). During the bloom period, chl_a is also higher in the ASP on average compared to the PIP (ASP = 5.21 ± 1.29 mg m⁻³; PIP = 3.69 ± 1.11 mg m⁻³; p -value < 0.01; Fig. 3b; Supplementary

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Table T1). When looking at polynya area integrated values (concentration multiplied by area gives units of mg m^{-1}), chl a is significantly higher in the ASP than in the PIP, and increases with the polynya area (Supplementary Figs. S1 and S2). NPP is also significantly higher in the ASP than in the PIP ($1.88 \pm 1.12 \text{ TgC y}^{-1}$ vs $0.85 \pm 0.86 \text{ TgC y}^{-1}$, $p\text{-value} = 0.004$; Supplementary Fig. S3). No significant interannual trends in mean chl a and NPP during the bloom are observed for either polynya ($p\text{-value} > 0.1$; Fig. 3b; Supplementary Fig. S3). The climatological winter MLD in the ASP is deeper (MLD ASP = $45.8 \pm 8.0\text{m}$; MLD PIP = $36.4 \pm 7.3\text{m}$; $p\text{-value} < 0.01$; Fig. 1b), indicating that it may better entrain deeper sources of nutrients into the upper waters for the following phytoplankton growing season, resulting in higher productivity (Fig. 2).

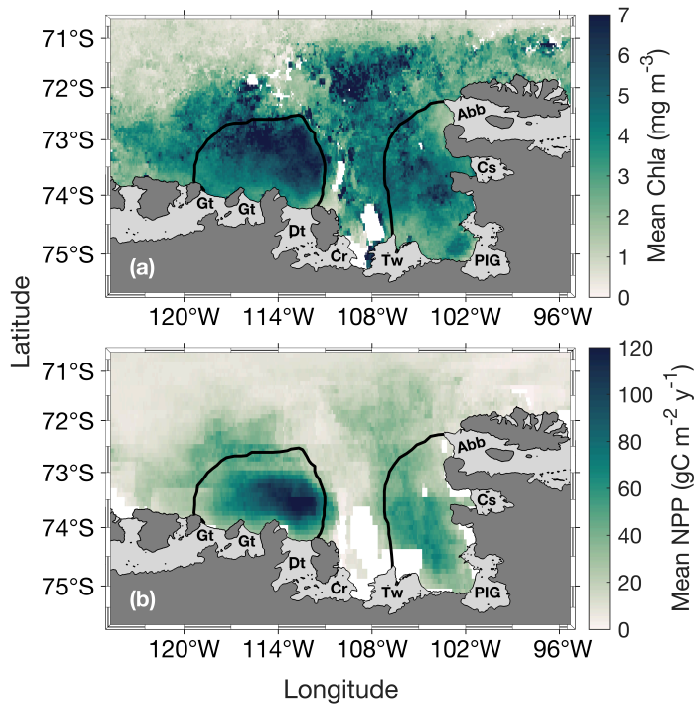


Fig. 2. Spatial distribution of (a) mean surface chlorophyll- a (chl a) concentration during the bloom and (b) net primary productivity (NPP) climatology (1998 – 2017) for the Amundsen

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(ASP; [left](#)) and Pine Island (PIP; [right](#)) polynyas. The black lines represent the climatological summer polynyas' boundaries.

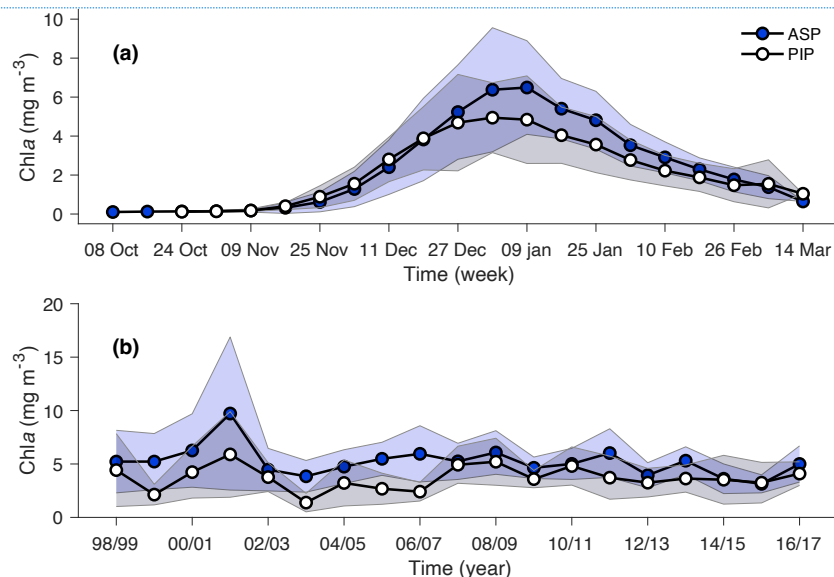


Fig. 3. (a) Weekly chlorophyll-*a* (*chl a*) [concentration](#) climatology (1998-2017) for the [Amundsen](#) (ASP; blue circles) and [Pine Island](#) (PIP; white circles) [polynyas](#). (b) Bloom mean *chl a* time series of ASP (blue circles) and PIP (white circles). Shaded areas represent the standard deviation for a given year. The relationship between *chl a* (in mg m⁻³ and mg m⁻¹) and the polynya size is shown in Supplementary Fig. S2.

The variability in TVFall is statistically uncorrelated with surface *chl a* and NPP in both polynyas from 1998 to 2017 (Fig. 4; Supplementary Fig. S4). However, the relationship becomes strongly significant in the ASP for both mean and [maximum](#) *chl a* when we remove the *chl a* outlier in 2001/02 (red data point; Figs. 4a-b), although not for NPP (Supplementary Figs. S4a-b). The positive relationship implies that surface *chl a* in the ASP is higher when more glacial meltwater

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is delivered to the embayment. No strong relationships are observed in the PIP between TVFall, surface chl *a* and NPP (Figs. 4c-d; Supplementary Figs. S4c-d).

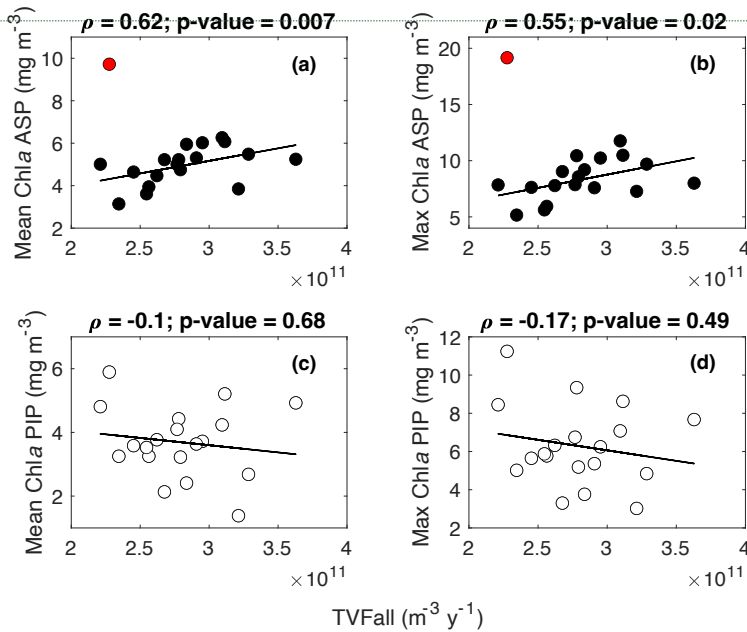


Fig. 4. Scatter plots of mean and maximum (max) surface chlorophyll-*a* (chl *a*) concentrations with the total volume flux (TVFall) for (a-b) the Amundsen (ASP) and (c-d) the Pine Island (PIP) polynyas from 1998 to 2017 (*n*=19). The fitted lines and statistics exclude year 2001/02 (red outlier) for the ASP regressions. If all data is considered, the relationships between mean chl *a*, max chl *a* and TVFall in the ASP are not significant. TVFall is an annual integral representing the sum of all ice shelves (see methods section) for the Amundsen Sea Embayment (ASE).

When fluxes from individual glaciers are considered, PIP chl *a* does not correlate with Abbot, Cosgrove, PIG, Thwaites or TVFpip fluxes (Table 1). On the other hand, ASP chl *a* shows strong relationships with TVFasp, the Dotson and Crosson ice shelves (Table 1), and all ice shelves

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become significantly correlated with mean and maximum chl_a when year 2001/02 is removed. There are no statistically significant relationships between individual ice shelves and NPP in either polynyas.

Table 1. Statistical summary (Spearman's rank correlation) of the relationships between ice shelves volume flux, mean and maximum (max) surface chlorophyll-*a* (chl_a) concentrations (n=19) in both polynyas. The * marks a significant (p-value < 0.05) relationship. All relationships between mean chl_a, maximum chl_a and ASP ice shelves become significant when year 2001/02 is removed.

| | Amundsen Sea polynya (ASP) | | | | Pine Island polynya (PIP) | | | |
|----------|----------------------------|--------------|----------------------|--------------|---------------------------|---------|----------------------|---------|
| | Mean chl _a | | Max chl _a | | Mean chl _a | | Max chl _a | |
| | rho | p-value | rho | p-value | rho | p-value | rho | p-value |
| Abbot | / | / | / | / | 0.09 | 0.73 | -0.04 | 0.88 |
| Cosgrove | / | / | / | / | -0.32 | 0.18 | -0.46 | 0.05 |
| PIG | / | / | / | / | -0.04 | 0.88 | -0.13 | 0.61 |
| Thwaites | 0.16 | 0.52 | 0.11 | 0.66 | 0.12 | 0.63 | 0.09 | 0.71 |
| Crosson | 0.43 | 0.07 | 0.50 | 0.03* | / | / | / | / |
| Dotson | 0.48 | 0.04* | 0.54 | 0.02* | / | / | / | / |
| Getz | 0.37 | 0.12 | 0.43 | 0.07 | / | / | / | / |
| TVFasp | 0.42 | 0.07 | 0.46 | 0.05* | / | / | / | / |
| TVFpip | / | / | / | / | 0.009 | 0.97 | -0.1 | 0.68 |

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Moved down [3]: chl_a are strongly correlated with TVFall in southern-eastern part of the ASP, in front of the Dotson ice shelf (Figs. 5a-b), where a positive relationship with NPP is also observed (Fig. 5c), although not significant.

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Fig. 4. Scatter plots

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Deleted: Statistical summary of the relationships between ice shelves volume flux and surface chlorophyll-*a* (chl_a).

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Spatially, the mean and maximum chl_a are strongly correlated with TVFall in southern-eastern part of the ASP, in front of the Dotson ice shelf (Figs. 5a-b), where a positive relationship with NPP is also observed (Fig. 5c), although not significant.

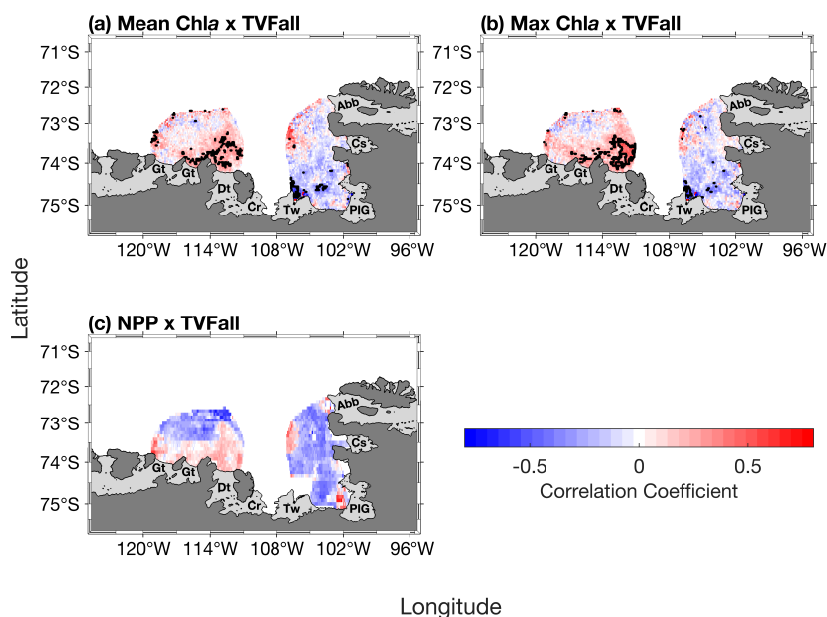


Fig. 5. Spatial correlation maps between total volume flux (TVFall) and (a) surface mean chlorophyll-*a* (chl_a) concentration, (b) surface maximum (max) chl_a concentration and (c) net primary productivity (NPP) (n=19). The black contour represents significant correlations at 95% confidence level. Data outside of the summer climatological polynyas' boundaries were masked out.

3.2 Simulated dFe sources distribution

The modelled spatial distribution of surface dFe sources is presented in Fig. 6. On average, the smallest dFe source in the embayment is from the ice shelves, with a maximum concentration

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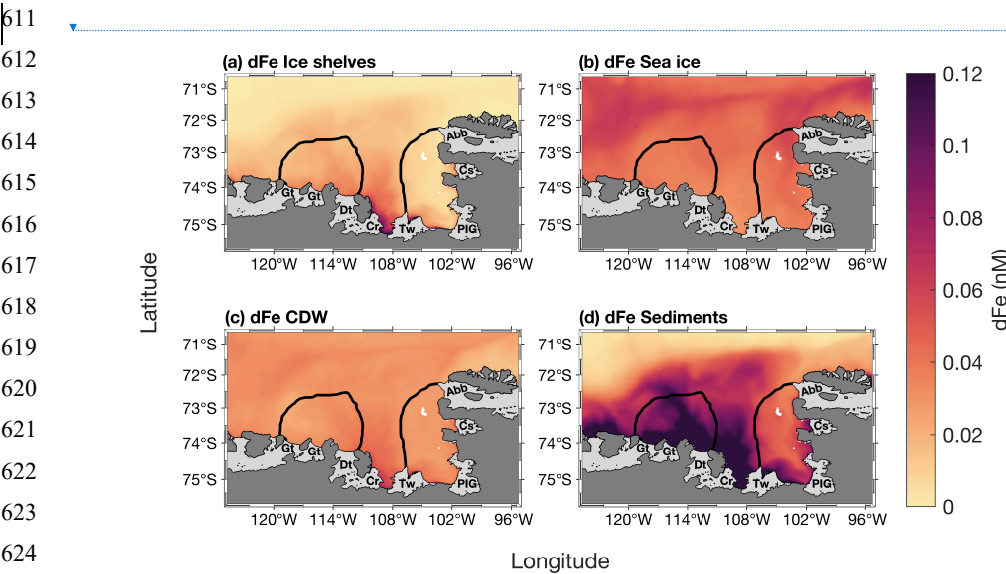
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Fig. 5

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600 between the Thwaites and Dotson ice shelves. The dFe from sea ice is slightly higher than from
 601 ice shelves and similar over the two polynyas, and is higher near the sea-ice margin (Fig. 6b). The
 602 dFe from CDW is also higher between the Thwaites and Dotson (Fig. 6c). Sediment is the
 603 dominant dFe source (Fig. 6d). Its distribution spreads from 108°W to the western part of the Getz
 604 ice shelf. The highest sediment-sourced dFe concentration is found along the coast and inside the
 605 ASP. On polynya-wide average basis, the sediment reservoir contributes significantly more to total
 606 dFe in the ASP (58.3%, 0.13nM) compared to sea ice (16.5%, 0.04nM), CDW (13.5%, 0.03nM)
 607 and ice shelves (11.7%, 0.03nM). In the PIP, the contribution of sediments is still significantly
 608 higher (41.2%; 0.08nM) but lower than the ASP and the contribution gap with the other sources
 609 decreases. The CDW and sea ice contribute 22.5% (0.04nM) and 18.9% (0.035nM) to the dFe pool
 610 respectively, while ice shelves are still the smallest sources at 14.5% (0.03nM) in the PIP.



626 **Fig. 6.** Two-years top-100m averaged spatial distribution of surface dissolved iron (dFe)
 627 contribution from (a) ice shelves, (b) sea ice, (c) circumpolar deep water (CDW) and (d) sediments
 628 simulated by the model from Dinniman et al. (2020). The black lines represent the climatological
 629 summer polynyas' boundaries.

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3.3 Environmental parameters, chl_a and NPP variability

During the phytoplankton growth season (October-March), SIC is spatially significantly anticorrelated to the meridional winds speed in both polynyas (Fig. 7a). Chl_a is significantly positively correlated with SST in the eastern ASP, and the whole PIP (Fig. 7b), but weakly with PAR in both polynyas (Fig. 7c). Finally, PAR and SST are positively related in both central polynyas, albeit not significantly (Fig. 7d). We note that similar spatial relationships are observed when NPP is correlated with SST and PAR (Supplementary Fig. S5).

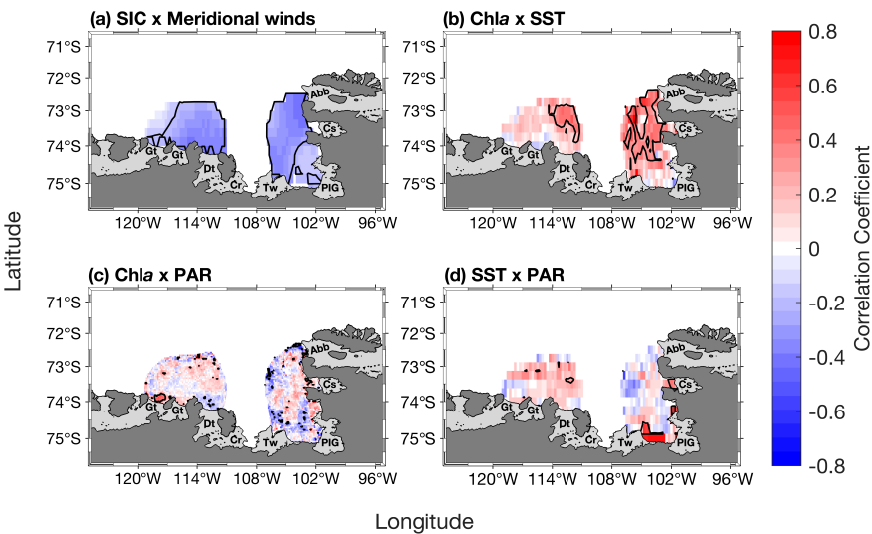


Fig. 7. Spatial correlation map between sea-ice concentration (SIC) and (a) meridional winds. Spatial correlation maps between mean chlorophyll-*a* (chl_a) concentration and (b) sea surface temperature (SST), (c) photosynthetically available radiation (PAR). (d) Spatial correlation map between PAR and SST. Data span 1998 – 2017 from October to March ($n=114$). The black contour represents significant correlations at 95% confidence level. Seasonality was removed before performing the correlation. Data outside of the summer climatological polynyas' boundaries were masked out.

666 Regarding the phenology, the bloom start is positively correlated to IRT and negatively with
667 OWP in the ASP, although not significantly with the OWP (Table 2). This means that the bloom
668 starts earlier and later as IRT does, and that longer OWP and earlier bloom starts are correlated
669 with earlier ice retreat. The bloom mean and bloom [maximum \(max\)](#) chl_a are not correlated with
670 either IRT and OWP in the ASP. IRT and OWP are significantly related ($p = -0.93$; p-value <
671 0.001). When year 2001/02 is removed, no significant changes in the relationships between all
672 parameters are detected. In the PIP, all metrics are significantly related to each other, except for
673 PAR and OWP (Table 2). That is, the bloom start is positively correlated with IRT and
674 negatively with OWP, while the bloom duration, mean chl_a, max chl_a and NPP are negatively
675 linked to the IRT and positively with OWP. SST and PAR are negatively correlated with IRT,
676 and positively with SST. IRT and OWP are significantly related in the PIP ($p = -0.88$; p-value <
677 0.001).

678
679 **Table 2.** Statistical summary ([Spearman's rank correlation](#)) of the relationships between the [sea-](#)
680 [ice phenology](#) metrics and environmental parameters (n=19) [in both polynyas](#). The * marks a
681 significant (p-value < 0.05) relationship. IRT = ice retreat time, OWP = open water period, NPP =
682 net primary productivity, SST = sea surface temperature, PAR = photosynthetically available
683 radiation. Removing year 2001/02 for the ASP slightly changes the strength of the relationships
684 between parameters (i.e., rho) but not the significance.

| | Amundsen Sea polynya (ASP) | | | | Pine Island polynya (PIP) | | | |
|----------------|-------------------------------|--------------|-------|---------|------------------------------|---------------|--------------|--------------|
| | IRT | | OWP | | IRT | | OWP | |
| | rho | p-value | rho | p-value | rho | p-value | rho | p-value |
| Bloom start | 0.51 | 0.03* | -0.43 | 0.07 | 0.56 | 0.02* | -0.48 | 0.04* |
| Bloom duration | -0.12 | 0.63 | 0.09 | 0.71 | -0.56 | 0.02* | 0.59 | 0.01* |
| Bloom mean | 0.19 | 0.44 | -0.33 | 0.17 | -0.67 | 0.003* | 0.50 | 0.04* |
| Bloom max | 0.24 | 0.32 | -0.35 | 0.14 | -0.65 | 0.005* | 0.52 | 0.03* |
| NPP | -0.55 | 0.02* | 0.45 | 0.05 | -0.72 | 0.001* | 0.54 | 0.02* |

| | | | | | | | | |
|-----|-------|------|-------|------|--------------|---------------|-------------|--------------|
| SST | -0.09 | 0.72 | -0.01 | 0.96 | -0.57 | 0.02* | 0.52 | 0.03* |
| PAR | -0.09 | 0.72 | 0.05 | 0.84 | -0.62 | 0.007* | 0.38 | 0.12 |

We explore the relationships between phytoplankton bloom phenology metrics and their potential environmental drivers by conducting a multivariate PCA for both polynyas (Fig. 8). The PCA reduces our datasets (11 variables) and breaks them down into dimensions that capture most of the variability and relationships between all variables. Arrows indicate the contribution of each variable to the dimensions, with longer arrows representing stronger influence. Observations (in our case, years) positioned in the direction of an arrow are more influenced by that variable. In the ASP (Fig. 8a), the first two principal components explain 55.3% of the total variance (Dim1: 35%, Dim2: 20.3%). NPP in the ASP is closely associated with BD, indicating that the bloom duration is the primary driver of production. On the other hand, environmental vectors such as TVFall and TVFasp project more strongly onto Dim2 with the bloom mean chl_a, indicating that meltwater input may influence surface chl_a interannual variability, and is less directly tied to NPP. We note that when year 2001/02 is removed, the relationship between TVFasp and TVFall becomes much stronger with [BM](#) (Supplementary Fig. S6a) and is slightly anticorrelated to SST and MLD. In the PIP (Fig. 8b), the first two components account for 63.7% of the total variance (Dim1: 46.7%, Dim2: 17%). Compared to the ASP, both NPP and BM cluster strongly with BD and PAR on Dim1. Additionally, IRT, OWP and SST and MLD aligned along Dim1, which explains 46.7% of the total variance compared to 35% for the ASP, suggesting that physical conditions might play a stronger structuring role in PIP compared to the ASP. In contrast, TVFall and TVFpip stand alone and align more strongly with Dim2, suggesting a less dominant influence of meltwater on [BM](#) and NPP variability in the PIP. [The summer](#) polynya-averaged PAR and MLD are significantly stronger and deeper, respectively, in the ASP compared to the PIP during the bloom season (MLD ASP = 28.5 ± 5.7 m; MLD PIP = 24.9 ± 3.7 m; p-value = 0.03 and PAR ASP = 31.5 ± 5.4 Einstein m⁻² d⁻¹; PAR PIP = 26.5 ± 6.7 Einstein m⁻² d⁻¹; p-value = 0.02).

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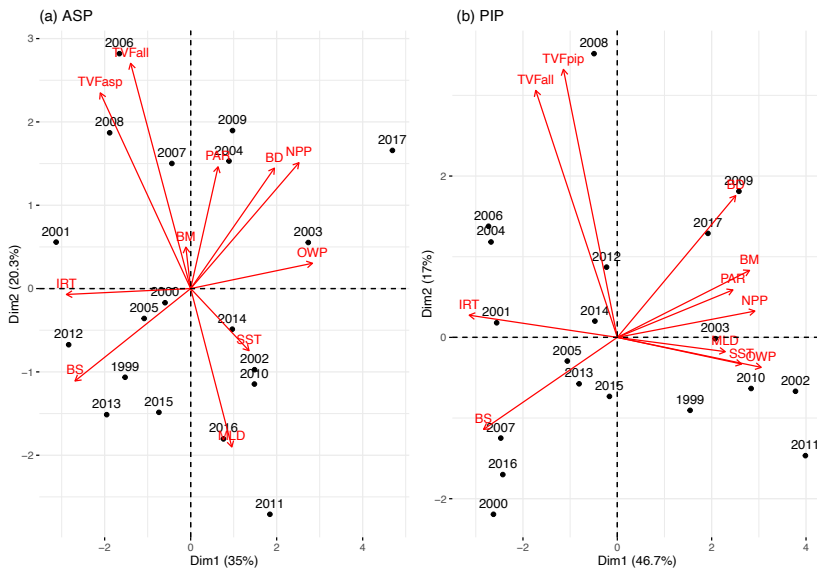


Fig. 8. Principal component analysis biplot of environmental parameters (red) and years (black) for (a) the [Amundsen \(ASP\)](#) and (b) the [Pine Island \(PIP\) polynyas](#). TVFasp = total volume flux for ASP; TVFpip = total volume flux for PIP; TVFall = total volume flux for all ice shelves; BM = bloom mean; PAR = photosynthetically available radiation; BD = bloom duration; NPP = net primary productivity; OWP = open water period; SST = sea surface temperature; MLD = mixed-layer depth; BS = bloom start; IRT = ice retreat time. The same plot is presented in supplementary Fig. S6, but removing year 2001/02 for the ASP, emphasizing the relationship between total volume flux (TVFall, TVFasp) and BM in the ASP.

Finally, we find on average weak spatial negative relationships between SIC and ASL latitude, longitude, mean sector and actual central pressure in both polynyas during the growing season (Supplementary Fig. S7), and only slightly significant in the eastern PIP.

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813 4. Discussion

814

815 4.1 Effect of glacial meltwater on phytoplankton *chl**a* and NPP

816

817 The relationship between glacial meltwater, surface *chl**a* and NPP observed over the last two
818 decades was distinctly different between the two polynyas. In the ASP, we found that enhanced
819 glacial [meltwater](#) translates into higher surface *chl**a*, but not with NPP (when removing year
820 2001/02; Figs. 4a-b; Supplementary Fig. S6a). Modelling results (Fig. 6) suggest that sediment
821 from the seafloor is the main source of dFe in the ASP, but this source is also linked to glacial
822 melt. Ice shelf glacial meltwater drives the meltwater pump, which brings up mCDW and fine-
823 grained subglacial sediments to the surface. This result is in agreement with previous research:
824 Melt-laden modified CDW flowing offshore from the Dotson ice shelf to the central ASP
825 (Sherrell et al., 2015), and resuspended sediments (Dinniman et al., 2020; St-Laurent et al., 2017;
826 2019) have been identified as significant sources of dFe to be used by phytoplankton.
827 Interestingly, both dFe supplied from ice shelves and CDW are most important in front of the
828 Thwaites and Crosson ice shelves, where the area averaged basal melt rate, and thus likely the
829 area averaged meltwater pumping (Jourdain et al., 2017), are typically strongest in observations
830 (Adusumilli et al., 2020; Rignot et al., 2013) and modelling (Fig. 6). The year 2001/02 does not
831 stand out as being influenced by any specific parameter in the ASP compared to other years (Fig.
832 8a, Supplementary Fig. S6a). The anomalously high surface *chl**a* observed during this year, as
833 also reported by Arrigo et al. (2012), may result from exceptional conditions that were not
834 captured by the parameters analysed in our study, for instance, an imbalance in the grazing
835 pressure. Interestingly, surface *chl**a* and NPP exhibit contrasting trends when averaged across the
836 polynya. While TVFall may explain some of the variance in surface *chl**a*, it does not account for
837 the variance in NPP, whether assessed through direct or multivariate relationships. This
838 decoupling between *chl**a* and NPP in the ASP suggests that glacial meltwater, while enhancing
839 surface phytoplankton biomass through nutrient delivery, may also promote vertical mixing. This
840 mixing deepens the mixed layer, reducing light availability and constraining photosynthetic
841 rates. These rates are influenced by fluctuations in the MLD, even in the presence of high
842 biomass and sufficient macronutrients. The summer MLD is deeper in the ASP, which would
843 decrease light availability, despite higher PAR compared to the PIP. Previous studies report that

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847 the small prymnesiophyte *P. antarctica*, a low-efficiency primary producer (Lee et al., 2017), is
848 better adapted to deeper mixed layers and therefore lower light conditions (Alderkamp et al.,
849 2012; Mills et al., 2010) and could contribute to high surface chl*a* decoupled from NPP, as
850 observed in the ASP. This is consistent with past *in situ* studies showing systematic differences
851 in mixed-layer structure between the two polynyas. The PIP commonly exhibits a shallow,
852 strongly stratified surface mixed layer while the ASP is more variable and has been observed to
853 host deeper MLD (Alderkamp et al., 2012; Park et al., 2017; Yager et al., 2016; Mills et al.,
854 2012). Near glacier and ice-shelf fronts, entrainment of iron-rich deep waters rising to the surface
855 through the meltwater pump can also produce surface chl*a* maxima (high biomass from *P.*
856 *antarctica*) without proportional increases in depth-integrated productivity due to self-shading
857 (Twelves et al., 2021). Further from the coast, meltwater spreading at neutral buoyancy
858 strengthens stratification, limiting vertical nutrient fluxes and thereby suppressing NPP despite
859 elevated chl*a*. These dual mechanisms are consistent with observational and modelling studies of
860 meltwater entrainment and dispersal (Randall-Goodwin et al., 2015; St-Laurent et al., 2017;
861 Dinniman et al., 2020; Forsch et al. 2021), and suggest that spatial heterogeneity in plume
862 dynamics could explain the observed chl*a* and NPP mismatch.

863
864 In the PIP, we did not find any long-term relationships between the phytoplankton bloom, NPP
865 and glacial meltwater. Variability in ice shelf glacial meltwater may not have the same effect on
866 the surface chl*a* and NPP in the PIP compared to the ASP. Iron delivered from glacial melt
867 process related in the PIP and west of it could accumulate and follow the westward coastal
868 current, towards the ASP (St-Laurent et al., 2017). These sources would include dFe from
869 meltwater pumped CDW, sediments and ice shelves, all of which are higher in front of the
870 Crosson ice shelf, west of the PIP (Fig. 6). With the coastal circulation, this would make dFe
871 supplied by glacial meltwater greater in the ASP, thereby contributing to the higher productivity
872 in the ASP. Recently, subglacial discharge (SGD) was shown to have a different impact on basal
873 melt rate in the ASE polynyas (Goldberg et al., 2023), where PIG had a lot less relative increase
874 in melt with SGD input than Thwaites or Dotson/Crosson. Thus, assuming a direct relationship
875 between melt rate, SGD and dFe sources, the signal in the PIP (fed by PIG melt) will be much
876 weaker than in the ASP (fed by upstream Thwaites, Crosson and local Dotson due to the
877 circulation), which might also explain the discrepancies between the PIP and ASP. A stronger

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878 meltwater-driven stratification may also dominate in the PIP, reducing vertical nutrient
879 replenishment and thereby limiting biomass growth (Oh et al., 2022), even where TVFall is high,
880 hence leading to a direct negative relationship observed compared to the ASP (Fig. 4;
881 Supplementary Fig. S4). The model outputs used here are critical to understand the spatial
882 distribution of dFe in the embayment. They strongly suggest, but do not definitively demonstrate,
883 the role of dFe in influencing the phytoplankton bloom interannual variability.

884
885 Satellite algorithms commonly estimate NPP from surface chl_a, but the approach and
886 assumptions vary across models. The VGPM relates chl_a to depth-integrated photosynthesis
887 through empirical relationships with light and temperature (Behrenfeld & Falkowski, 1997). In
888 contrast, the Carbon-based Productivity Model (CbPM) emphasizes phytoplankton carbon
889 biomass and growth rates derived from satellite optical properties, decoupling productivity
890 estimates from chl_a alone (Westberry et al., 2008). The CAFE model (Carbon, Absorption, and
891 Fluorescence Euphotic-resolving model) integrates additional physiological parameters such as
892 chl_a fluorescence and absorption to better constrain phytoplankton carbon fixation (Silsbe et al.,
893 2016). In the Southern Ocean, where light limitation, iron supply, and community composition
894 strongly influence the relationship between chl_a and productivity, these algorithmic differences
895 can yield substantial variability in NPP estimates (Ryan-Keogh et al., 2023), with studies
896 showing that VGPM-type models often outperform CbPM in coastal Southern Ocean regions
897 (Jena et al., 2020). Because the VGPM algorithm does not explicitly incorporate the MLD, but
898 instead estimates primary production integrated over the euphotic zone based on surface chl_a,
899 PAR, and temperature, it may not fully capture the influence of variable MLD or subsurface
900 processes related to glacial melt, which could contribute to the observed decoupling between
901 chl_a and NPP. Therefore, while the observed decoupling between chl_a and NPP in the ASP
902 might also come from our choice of dataset, the VGPM model may be more appropriate for
903 coastal polynya environments, such as those in the Amundsen Sea. We finally note as a
904 limitation that satellite-derived chl_a and VGPM NPP estimates lack the vertical resolution
905 needed to resolve sub-plume stratification and mixing processes (e.g., fine-scale vertical
906 gradients in iron or nutrient fluxes), so our mechanistic interpretations of surface chl_a vs. depth-
907 integrated productivity decoupling must be taken with caution.

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Moved up [5]: Further from the coast, meltwater spreading at neutral buoyancy strengthens stratification, limiting vertical nutrient fluxes and thereby suppressing NPP despite elevated chl_a. These dual mechanisms are consistent with observational and modelling studies of meltwater entrainment and dispersal (Randall-Goodwin et al., 2015; St-Laurent et al., 2017; Dinniman et al., 2020; Forsch et al. 2021), and suggest that spatial heterogeneity in plume dynamics could explain the observed chl_a and NPP mismatch.

Moved down [6]: note as a limitation that satellite-derived chl_a and VGPM NPP estimates lack the vertical resolution needed to resolve sub-plume stratification and mixing processes (e.g., fine-scale vertical gradients in iron or nutrient fluxes), so our mechanistic interpretations of surface chl_a vs. depth-integrated productivity decoupling must be taken with caution.

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934 Direct observations from Sherrell et al. (2015) showed higher chl_a in the central ASP while
 935 surface dFe was low weeks before the bloom peak. This suggests a continuous supply and
 936 consumption of dFe in the area, most likely from the circulation, as mentioned earlier.
 937 Considering the long residence time of water masses in both polynyas (about 2 years (Tamsitt et
 938 al., 2021)), and the daily dFe uptake by phytoplankton (3-196 pmol l⁻¹ d⁻¹ (Lannuzel et al.,
 939 2023)), we also hypothesise that any dFe reaching the upper ocean from external sources is
 940 quickly used and unlikely to remain readily available for phytoplankton in the following spring
 941 season.

942

943 In recent model simulations with the meltwater pump turned off, Fe becomes the principal factor
 944 limiting phytoplankton growth in the ASP (Oliver et al., 2019). However, the transport of Fe-rich
 945 glacial meltwater outside the ice shelf cavities and to the ocean surface depends strongly on the
 946 local hydrography. While Naveira Garabato et al. (2017) suggested that the glacial meltwater
 947 concentration and settling depth (neutral buoyancy) outside the ice shelf cavities is controlled by
 948 an overturning circulation driven by instability, others suggest that the strong stratification plays
 949 an important role in how close to the surface the buoyant plume of said meltwater can rise
 950 (Arnscheidt et al., 2021; Zheng et al., 2021). Therefore, high melting years and greater TVFall
 951 might not necessarily translate into a more iron-enriched meltwater delivered to the surface
 952 outside the ice shelf cavities, close to the ice shelf edge, as rising water masses may be either
 953 prevented from doing so, or be transported further offshore in the polynyas where the
 954 phytoplankton bloom occurs, before they can resurface (Herraiz-Borreguero et al., 2016).

955

956 Although several Fe sources can fuel polynya blooms, and they depend on processes mentioned
 957 above, Fe-binding ligands may ultimately set the limit on how much of this dFe stays dissolved
 958 in the surface waters (Gledhill and Buck, 2012; Hassler et al., 2019; Tagliabue et al., 2019).
 959 Models of the Amundsen Sea (Dinniman et al., 2020, 2023; St-Laurent et al., 2017, 2019) did not
 960 include Fe complexation with ligands and assumed a continuous supply of available dFe for
 961 phytoplankton. Spatial and seasonal data on Fe-binding ligands along the Antarctic coast remain
 962 extremely scarce and their dynamics are poorly understood (see Smith et al. (2022) for a
 963 database of publicly available Fe-binding ligand surveys performed south of 50°S). Field
 964 observations in the ASP and PIP suggest that the ligands measured in the upwelling region in

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967 front of the ice shelves had little capacity to complex any additional Fe supplied from glacial
968 melt. As a consequence, much of the glacial and sedimentary Fe supply in front of the ice
969 shelves could be lost via particle scavenging and precipitation (Thuróczy et al., 2012). This was
970 also observed by van Manen et al. (2022) in the ASP. However, within the polynya blooms,
971 Thuróczy et al. (2012) found that the ligands produced by biological activity were capable of
972 stabilising additional Fe supplied from glacial melt, where we observed the highest productivity.
973 The production of ligands by phytoplankton would increase the stock of bioavailable dFe and
974 further fuel the phytoplankton bloom in the polynyas, potentially highlighting the dominance
975 of *P. antarctica*, which uses iron-binding ligands more efficiently than diatoms (Thuróczy et al.,
976 2012), even under low light conditions. Model development and sustained field observations on
977 dFe availability, including ligands, are needed to adequately predict how these may impact
978 biological productivity under changing glacial and oceanic conditions, now and in the future.

980 Overall, the discrepancies observed between the ASP and PIP point to a complex set of ice-
981 ocean-sediment interactions, where several co-occurring processes and differences in
982 hydrographic properties of the water column influence dFe supply and consequent primary
983 productivity.

985 4.2 Possible drivers of the difference in phytoplankton surface chl a and NPP between the 986 two polynyas

988 The biological productivity is higher in the ASP than the PIP, consistent with previous studies
989 (Arrigo et al., 2012; Park et al., 2017). In section 4.1, we mentioned the suspected underlying
990 hydrographic drivers of these differences. We related the higher biological productivity in the
991 ASP to a potentially greater supply of iron from melt-laden dFe-enriched mCDW and sediment
992 sources, but this difference in productivity could also be attributed to other local features. The
993 Bear Ridge grounded icebergs on the ASP's eastern side (Bett et al., 2020) could add to the
994 overall meltwater pump strength. They can enhance warm CDW intrusions to the ice shelf cavity
995 (Bett et al., 2020), increasing ice shelf melting and subsequent stronger phytoplankton bloom
996 from the meltwater pump activity. These processes are weaker or absent in the PIP. Few sources
997 other than glacial meltwater may influence the bloom in the PIP. For instance, dFe in the

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1001 euphotic zone can also be sustained by the biological recycling, as shown in the PIP by Gerringa
1002 et al. (2020).

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1004 Sea ice could also partly explain the difference in chl_a magnitudes, NPP, and variability between
1005 the ASP and PIP. The strong spatial correlation between SIC and meridional winds (Fig. 7a)
1006 indicates that southerly winds can export the coastal sea ice offshore and play a significant role
1007 in opening the polynyas. In the ASP compared to the PIP, sea ice retreats earlier (IRT = Jan 1st ±
1008 14d vs Jan 18th ± 17d, p-value = 0.003), the open water period is longer (OWP = 61 ± 16d vs 44
1009 ± 22d, p-value < 0.001), and the SIC is lower (Supplementary Fig. S8). In the ASP, an early sea-
1010 ice retreat leads to an earlier bloom start, but the longer open water period is not significantly
1011 associated with greater bloom mean [or maximum](#) chl_a (Table 2). On the other hand in the PIP,
1012 an early sea-ice retreat also triggers an early bloom start, but the longer open duration is
1013 associated with warmer water, higher bloom mean chl_a, [maximum](#) chl_a, and NPP. These results
1014 suggest that different processes might drive phytoplankton growth variability in the two
1015 polynyas. In the ASP, it is likely the replenishment of dFe mentioned above that mostly
1016 influences the bloom. In the PIP, higher SIC can delay the retreat time and shorten the open
1017 water season (Table 2, Supplementary Fig. S8), leading to lower chl_a and NPP compared to the
1018 ASP. The significant negative relationships between IRT, PAR, chl_a and NPP in the PIP (Table
1019 2, Supplementary Fig. S6) suggests a strong light limitation relief in the polynya. This light
1020 limitation hypothesis is further supported by the high correlation between polynya-averaged chl_a
1021 mean with PAR and SST in the PIP across the 19 years of study, compared to the lack of
1022 correlation in the ASP (Supplementary Table T2; p-value < 0.01 for all relationships in the PIP).
1023 [While *P. antarctica*](#) is usually the main phytoplankton species dominating in both polynyas, the
1024 combination of light-limitation relief and higher SST may create better conditions for a stratified
1025 and warmer environment that would favor diatom (Arrigo et al., 1999; van Leeuwe et al., 2020),
1026 as recently observed in the ASP (Lee et al., 2022). The positive association of PAR, SST and
1027 chl_a with MLD likely reflects conditions around sea-ice retreat (all negatively associated with
1028 IRT), when enhanced wind mixing deepens the mixed layer and replenishes surface nutrients
1029 while light availability and SST increases. This nutrient-light co-limitation phase supports high
1030 biomass accumulation, likely from diatoms. Similar results have been reported by Park et al.
1031 (2017). They found that the PIP was dFe [was not limited by dFe](#), potentially from biological

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recycling (Gerringa et al., 2020), compared to an iron-limited ASP. We hypothesise that the connection between glacial meltwater and chl_a that we found in the ASP is a response to iron input (also observed by Park et al. (2017) during incubation experiments) compared to the PIP, where light and temperature seem to play a more significant role in driving the phytoplankton bloom variability. Hayward et al. (2025) showed a decline in diatoms from 1997 to 2017 in the PIP. However, they observed an increase in diatoms after 2017, linked to regime shift in sea ice. Their study also indicates that diatoms are competitively disadvantaged under iron-depleted conditions. *P. antarctica*, which relies on dFe supplied by ocean circulation, would then tend to dominate in the ASP. Such shifts in phytoplankton composition are likely to affect carbon export, grazing, and higher trophic levels. Additional long-term data on inter-annual variability in phytoplankton composition and physiology will be essential to fully understand these relationships.

Finally, the weak relationships between the ASL indices and SIC might be owing to the seasonal variation of the ASL, where its position largely varies during summer, and its impact in shaping coastal sea ice is also greater during winter and autumn in the Amundsen-Bellingshausen region (Hosking et al., 2013). The lack of strong significant relationships overall does not allow us to conclude that the ASL plays an important role in shaping the coastal polynyas landscape and influencing chl_a variability.

4.3 Limitations and future directions

We acknowledge that elevated concentrations of suspended sediments (and non-photosynthetically active particles in general) near the ocean surface can impart optical signatures that bias satellite-derived chl_a high in coastal waters. Consequently, the higher chl_a observed in the ASP relative to the PIP, as well as the weak correspondence between chl_a and NPP in ASP, may reflect some sediment-driven optical effects rather than enhanced phytoplankton biomass or productivity alone. While our results are consistent with known differences in iron supply and mixed-layer dynamics between the two polynyas, the potential contribution of sediment-related bias cannot be ruled out and should be acknowledged when interpreting spatial contrasts in satellite chl_a on the Antarctic shelf.

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Deleted: Our results suggest potential long-term changes in the phytoplankton community, specifically a shift towards diatoms in the ASE coastal regions during phytoplankton bloom. Hayward et al. (2025) reported

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Deleted: Variability in SIC and sea-ice retreat can be influenced by the Amundsen Sea Low (ASL). We therefore also investigated its potential role on sea-ice variability. We found on average weak spatial negative relationships between SIC and ASL latitude, longitude, mean sector and actual central pressure in both polynyas during the growing seasons (Supplementary Fig. S8), and only slightly significant in the eastern PIP. The weak relationships

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1083 While it seems reasonable that the higher ASP productivity could be driven by more iron
1084 delivered through a stronger meltwater pump downstream of the PIP, our data cannot confirm
1085 this hypothesis. To accurately understand the role of iron through the meltwater pump process,
1086 we would need to quantify the fraction of meltwater and glacial modified water (mix of CDW
1087 and ice shelf meltwater) reaching the ocean surface, together with the iron content. Obtaining
1088 this information is challenging over the decadal time scales considered and the method used in
1089 our study. Here, our intention was to provide valuable insights into the potential drivers of our
1090 results, and highlight the benefit of remote sensing, in this poorly observed environment. Our
1091 work directly aligns with Pan et al. (2025), who investigated the long-term relationship between
1092 sea surface glacial meltwater and satellite surface chl_a in the Western Antarctic Peninsula, and
1093 found a strong relationship between the two parameters, highlighting the importance of glacial
1094 meltwater discharge in regions prone to extreme and rapid climate changes.

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1096 In multimodel climate change simulations, Naughten et al (2018) showed an increase of ice
1097 shelves melting up to 90% on average, attributed to more warm CDW on the shelf, due to
1098 atmospherically driven changes in local sea-ice formation. More recently, Dinniman et al. (2023)
1099 also highlighted the impact of projected atmospheric changes on Antarctic ice sheet melt. They
1100 showed that strengthening winds, increasing precipitation and warmer atmospheric temperatures
1101 will increase heat advection onto the continental shelf, ultimately increasing basal melt rate by
1102 83% by 2100. Compared to present climate simulations, their simulation showed a 62% increase
1103 in total dFe supply to shelf surface waters, while basal melt driven overturning Fe supply
1104 increased by 48%. The ice shelf melt and overturning contributions varied spatially, increasing in
1105 the Amundsen-Bellingshausen area and decreasing in East Antarctica. This implies that, under
1106 future climate change, phytoplankton productivity could show stronger spatial asymmetry
1107 around Antarctica. The increasing melting and thinning of ice shelves will eventually result in
1108 more numerous calving events and drifting icebergs (Liu et al., 2015). Model simulations
1109 stressed the importance of ice shelves and icebergs in delivering dFe to the SO (Death et al.,
1110 2014; Person et al., 2019), increasing offshore productivity. As Fe will likely be replenished and
1111 sufficient from increasing melting in coastal areas, it is possible that the system will shift from

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1112 Fe-limited to being limited by nitrate, silicate, or even manganese (Anugerahanti and Tagliabue,
1113 2024), while offshore SO productivity will likely remain Fe-dependent (Oh et al., 2022).

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1115 5. Conclusions

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1117 Using spatial and multivariate approaches, our study explored the variability of surface chl_a and
1118 NPP in the Amundsen Sea polynyas over the last two decades, with a focus on the main
1119 environmental characteristics of the ASE. We found a strong relationship between ice shelf
1120 melting and surface chl_a in the ASP, when year 2001/02 was removed, a result in agreement with
1121 the ASPIRE field studies and previous satellite analyses. On the other hand, we did not find clear
1122 evidence of such a relationship in the PIP, where light, sea surface temperature and open water
1123 availability seem more important. The differences between the polynyas may lie in hydrographic
1124 properties, or the use of satellite remote sensing itself, which cannot tell us about processes such
1125 as Fe supply, bioavailability and phytoplankton demand. To gain greater insight, we referred to
1126 model simulations that showed the spatial variability in the magnitude of iron sources. Our
1127 results call for sustained *in situ* observations (e.g., moorings equipped with trace-metal clean
1128 samplers, and physical sensors to better understand year-to-year water mass meltwater fraction
1129 and properties) to elucidate these long-term relationships. Satellite observations are a powerful
1130 tool to investigate the relationship between glacial meltwater and biological productivity on such
1131 time scales, which has until now relied almost exclusively on field observations and modelling.
1132 Using such tools, we showed how the relationship between phytoplankton and the environment
1133 varies spatially and temporally across 19 years.

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1135 Appendices

1136 No appendices are related to the manuscript.

1137

1138 Data availability

1139 Bathymetry data (Amante & Eakins, 2009) was taken from the NOAA website
1140 (<http://www.ngdc.noaa.gov/mgg/global/global.html>). Mixed-layer depth (ECCO Consortium et
1141 al., 2021) can be accessed here:
1142 https://podaac.jpl.nasa.gov/dataset/ECCO_L4_MIXED_LAYER_DEPTH_05DEG_MONTHLY

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1145 V4R4. Satellite surface chlorophyll-*a* and photosynthetically available radiation were
1146 downloaded from <https://www.globcolour.info/>. Sea surface temperature (Huang et al., 2021)
1147 can be found here <https://psl.noaa.gov/data/gridded/data.noaa.oisst.v2.highres.html>. Wind re-
1148 analysis data (Hersbach et al., 2020) are available at
1149 [https://cds.climate.copernicus.eu/datasets/reanalysis-era5-single-levels-monthly-](https://cds.climate.copernicus.eu/datasets/reanalysis-era5-single-levels-monthly-means?tab=download)
1150 [means?tab=download](https://cds.climate.copernicus.eu/datasets/reanalysis-era5-single-levels-monthly-means?tab=download). Sea-ice concentration (Cavalieri et al., 1996) was obtained from
1151 <https://nsidc.org/data> and Net Primary productivity (Behrenfeld and Falkowski, 1997) was
1152 downloaded from <http://sites.science.oregonstate.edu/ocean.productivity/index.php>. Circumpolar
1153 surface model output from Dinniman et al (2020) can be found at [https://www.bco-](https://www.bco-dmo.org/dataset/782848)
1154 [dmo.org/dataset/782848](https://www.bco-dmo.org/dataset/782848). The Amundsen Sea Low index (Hosking et al., 2016) data are available
1155 at http://scotthosking.com/asl_index.

1156 1157 **Author contributions**

1158 GL conceptualised and led the study; MSD provided the dissolved iron model output. All authors
1159 were involved in the interpretation of the results, the revision, and the writing of the final version
1160 of the paper.

1161 1162 **Competing interest**

1163 We declare having no competing interests.

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1182

1183 **References**

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