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Resolving sea ice dynamics in the north-western Ross Sea during the last 2.6 ka: From seasonal to millennial timescales

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29 Abstract

Time-series analyses of satellite images reveal that sea ice extent in the Ross Sea has experienced 30 31 significant changes over the last 40 years, likely triggered by large-scale atmospheric anomalies. However, resolving how sea ice in the Ross Sea has changed over longer timeframes has until now 32 33 remained more elusive. Here we used a laminated sediment piston core (14.6 m) collected from the 34 Edisto Inlet (Western Ross Sea) to reconstruct fast ice dynamics over the last 2.6 ka. Our goal was to 35 first understand the climate expression of selected well-defined sediment laminae and then use these 36 characteristics for reconstructing past sea ice behaviour across the whole sedimentary sequence. We used the recently established sea ice diatom biomarker proxy IPSO₂₅ in combination with diatom 37 census counts and bulk analyses. Analyses performed on a suite of discrete laminae revealed 38 statistically significant differences between dark and light laminae reflecting different depositional 39 conditions. Based on their respective biogeochemical fingerprints, we infer that dark laminae 40 accumulated during sea ice thaws in early summer. Under these conditions, laminae contain relatively 41 high concentrations of IPSO₂₅ and display an enriched $\delta^{13}C$ composition for the bulk organic matter 42 (OM). While diatom assemblages in dark laminae are relatively homogenous, as the thaw continues 43 44 later in the summer, Corethron pennatum becomes the dominant diatom species, resulting in the 45 formation of light laminae characterized by low IPSO₂₅ concentrations. Since C. pennatum can 46 migrate vertically through the water column to uptake nutrients and avoid competition in oligotrophic 47 waters, its high concentration likely reflects stratified and ice-free surface waters typical of late 48 summer.

49 Down-core trends show that the correlation between sediment brightness and geochemical fingerprint (i.e., IPSO₂₅ and δ^{13} C) holds throughout the record. Based on the knowledge gained at lamina level, 50 51 our down-core high-resolution reconstruction shows that the summer fast ice coverage changed dramatically during the late Holocene. Specifically, we conclude that the Edisto inlet experienced 52 53 regular early summer opening between 2.6 ka, and ca. 0.7 ka, after which, coastal fast ice persisted 54 during summer months and ice-free conditions became less frequent. Comparison with previous 55 regional ice core data suggests that the sudden cooling recorded over the Victoria Land Coast region 56 since 0.7 ka might potentially explain our observation of persistent summer fast ice in the Western

- Ross Sea. Our study has shown that multi-proxy data derived from laminated sediments can provide
 hitherto unknown detail regarding past summer sea ice dynamics in coastal Antarctic regions.
- 59

60 **1. Introduction**

Since multichannel passive-microwave satellite records began in the late 1970's, it has become evident 61 62 that Antarctic sea ice extent has exhibited sub-regional patterns. Various mechanisms have been proposed to explain such spatial variability, including the El Niño-Southern Oscillation (Stammerjohn 63 64 et al., 2008), the Interdecadal Pacific Oscillation (Meehl et al., 2016), the Amundsen Sea Low (Turner et al., 2009), as well as basal melting of Antarctic ice shelves (Bintanja et al., 2013). Despite these 65 66 diverse explanations, the collective picture indicates that Antarctic sea ice distribution is expressed by large-scale climate dynamics; in turn, sea ice regulates fundamental aspects of global climate (e.g. 67 thermohaline circulation, ocean-atmosphere heat/gas exchange and radiative properties). However, a 68 survey of the current literature clearly reveals that our knowledge of sea ice-climate interactions in the 69 Southern Ocean essentially relies on the last 40 years of satellite imagery, with longer-term records far 70 71 more elusive because of the paucity of sea ice archives.

72 In this study, we demonstrate the suitability of laminated sedimentary units for the reconstruction of 73 past sea ice dynamics in Antarctica. Laminated diatom ooze deposits have been documented in 74 different regions of the Antarctic margin, including the Antarctic Peninsula (Bahk et al., 2003; Hjort et 75 al., 1997; Leventer et al., 2002; Maddison et al., 2005), the East Antarctic Margin (Alley et al., 2018; 76 Denis et al., 2006; Maddison et al., 2012; Maddison et al., 2006), and the Ross Sea (Finocchiaro et al., 77 2005; McKay et al., 2016). Such laminated records often reflect weak post-depositional reworking, 78 which makes them suitable archives for resolving seasonal and sub-seasonal processes driven by 79 short-lived climate variability. Although the formation of laminated units is dependent on local and 80 regional conditions, collectively, their deposition is generally attributed to spring and summer algal bloom events associated with seasonal sea ice retreat (Abram et al., 2014; Finocchiaro et al., 2005; 81 Leventer et al., 2002; Maddison et al., 2012). As such, analysis of such laminated sediments using 82 appropriate proxy signatures of the respective bloom events has the potential to provide insights into 83 local sea ice dynamics (at least) and any changes to these over time. 84

In this study, we analysed laminated sediments from a piston core collected from the Edisto inlet 85 (Western Ross Sea) (HLF17-1, Fig. 1; Fig. 2) to reconstruct past sea ice variability during the late 86 87 Holocene (last ca. 2.6 ka) via an expanded record characterized by continuous diatom-rich 88 laminations. Since the 1970s, sea ice has increased over the Ross Sea, followed by rapid negative anomalies in recent years (e.g. 2016) (Parkinson, 2019; Turner et al., 2017). The reasons behind the 89 90 strong variability of sea ice in the Southern Ocean is a matter of on-going discussion, although 91 compiled evidence suggests that there are likely multiple factors that collectively involve anomalies in 92 the atmospheric and, consequently, oceanic circulation (Meehl et al., 2019). Our study focuses on 93 land-fast ice dynamics, which is a major sea ice type common to Antarctic coastal regions. Sea ice 94 reconstructions have traditionally been based on ecological proxies such as diatom assemblages, whose relative abundance in Antarctic sediments reflects changes in sea surface conditions (Armand et 95 al., 2005; Gersonde and Zielinski, 2000; Leventer, 1998). Over the last decade, sea ice diatom-96 produced Highly Branched Isoprenoid (HBI) lipids have emerged as novel biomarker proxies for 97 Arctic and Antarctic sea ice (Belt, 2018, 2019; Belt and Müller, 2013; Collins et al., 2013; Massé et 98 99 al., 2011; Vorrath et al., 2019). For the Southern Ocean, the di-unsaturated HBI termed IPSO₂₅ ("Ice 100 Proxy for the Southern Ocean with 25 carbon atoms") has been suggested as a biomarker of land-fast 101 ice. In a preliminary study (Belt et al., 2016), it was suggested that this source-specific HBI might 102 trace the occurrence of the sympagic (i.e. living within sea ice) diatom Berkeleya adeliensis, which 103 flourishes predominantly in the bottom sections of fast ice and within the underlying platelet ice (Riaux-Gobin et al., 2013). Otherwise, unlike its close structural counterpart IP₂₅ in the Arctic (see 104 105 Belt, 2018 for a recent review), there have been no in-depth investigations into the relationship 106 between IPSO₂₅ distributions and Antarctic sea ice characteristics (*e.g.* sea ice concentration, ice type, 107 presence of polynyas, etc.). A relatively small number of Holocene sea ice reconstructions based on 108 IPSO₂₅ have assumed a positive relationship between sedimentary IPSO₂₅ concentration and sea ice 109 extent in a general sense, an interpretation supported in some cases by diatom assemblage data (Denis et al., 2010; Etourneau et al., 2013; Lamping et al., 2020; Massé et al., 2011). Finally, in a recent pilot 110 study of surface sediments from the West Antarctic Peninsula, Vorrath et al. (2019) concluded that a 111 number of different factors likely need to be considered when using IPSO₂₅ (and combined biomarker 112

indices based on IPSO₂₅) for paleo sea ice reconstruction, especially given the complex nature of Antarctic sea ice dynamics. As such, it seems likely that the use of IPSO₂₅ as a sea ice proxy needs to be carried out on a context-specific, rather than a generic, basis.

This study presents the first high-resolution reconstruction of late-Holocene sea ice dynamics in the 116 Western Ross Sea inferred from IPSO₂₅. Among its various attributes, the comparatively time-efficient 117 analysis of IPSO₂₅ makes it an ideal tool to be used in high-resolution studies dealing with laminated 118 119 sequences. However, since IPSO₂₅ is still a relatively new proxy (see Belt, 2018 for a review) and the 120 likely need to use it according to the specific sea ice setting (see above), our study also involves the 121 analysis of some diatom assemblages and other complementary biogeochemical and lithological 122 parameters, which include the stable isotopic composition of bulk organic carbon, inorganic elemental 123 composition, and grain size measurements.

The overarching objectives of this study were threefold: (i) to investigate the extent to which proxy data obtained from dark and light laminae found in HLF17-1 reflected annual, seasonal or subseasonal fast ice dynamics; (ii) to use this information to infer the millennial-scale variability of sea ice in the inner-shelf of the Ross Sea over the last 2.6 ka BP; (iii) to rationalise findings through consideration of other climatic parameters pertinent to the late Holocene.

129

130 2. Regional setting

Edisto Inlet is small ice-filled elongated NNE-SSW fjord situated behind Cape Hallett, along the Victoria Land Coast in the north-western Ross Sea. The fjord is approximately 15 km long and 4 km wide, with a maximum water depth of approximately 500 m and a sill 400 m deep, which divides the fjord from Moubray Bay to the north. Geophysical data (3.5 kHz sub-bottom profiler) acquired during the 2005 PNRA (National Antarctic Research Program) Italian expedition highlighted that the studied core was collected in a very expanded Holocene sequence characterized by soft biogenic laminated sediments (Fig. 1c).

138

3. Materials and methods

140 3.1. Sampling and subsampling

141 Piston core HLF17-1 (72° 18.4842' S - 170° 03.2592' E; 465 m water depth) was collected in February 142 2017 in the Edisto inlet on board of the R/V Italica (Fig. 1). The core, 14.65 m long, was split into 1-m long sections and stored at 4°C on the vessel. At the same site, a box corer (HLF17-2BC, 56 cm) was 143 retrieved and kept refrigerated under the same conditions as the piston core. In the laboratory, open 144 145 core sections were analysed using an AVAATECH core scanner for high-resolution image acquisition 146 and XRF analyses (0.5-cm sampling interval). Sediments were sub-sampled throughout every 5 cm at 147 1-2 cm thick intervals (n=295), while distinct laminae (n=34) were subsampled from section III. All 148 subsamples were frozen and freeze-dried prior to analysis.

The HLF17-1 core site was visited previously in 2016 on board the *R/V Italica*, during which time a 11.43 m long core (HLF16-1) was retrieved and split into sections as described above. Unfortunately, freezing of the core due to failure of the cooling system resulted in the expansion and loss of sediments from the liner ends. Despite this handling issue, the stratigraphy of remaining sediments was well preserved. HLF17-1 and HLF16-1 were correlated with each other using distinctive lamination patterns (Supplementary Material, Fig. S1). HLF16-1was then sub-sampled (10-cm thick intervals) and wet-sieved to isolate carbonate microfossils for the age-depth model of HLF17-1.

156

157 3.2. Bulk analyses

Freeze-dried samples were powdered and homogenized in an agate mortar. Acidified sediments (1.5 M 158 HCl) were analysed using a Thermo Fisher Elemental Analyser (FLASH 2000 CHNS/O) coupled with 159 160 a Thermo Finnigan Delta plus isotope ratio mass spectrometer (IRMS) for organic carbon (OC, wt.%), 161 total nitrogen (TN, wt.%) and stable carbon isotopes (δ^{13} C, ‰) (D'Angelo et al., 2018; Tesi et al., 2012). Opal content (wt.%) was measured according to the leaching method developed by Mortlock 162 163 and Froelich (1989). Sediments were placed in Teflon tubes and alkaline dissolution was performed with 0.5 M Na₂CO₃ solution at 80 °C for 5 h. Dissolved silica was measured according to the 164 165 molybdate-blue spectrophotometric method ($\lambda = 812$ nm). Data are reported as weight percentage 166 (wt.%) of SiO₂ x (0.4H₂O).

Major (%) and minor (ppm) elements were quantified using a wavelength dispersive Philips PW 1480
sequential X-ray fluorescence spectrometer (XRF). Analyses were carried out on pressed powder

pellets applying the matrix correction as presented in Dinelli *et al.* (2001). Aliquots designated for
XRF were used to make loss-on-ignition (LOI, 950°C) measurements. The reproducibility for major
and minor elements was ca. 7% and 5%, respectively, and shifted to ca. 15% for minor elements
<10ppm (Dinelli *et al.*, 2001). In this study, only the major lithogenic elements are presented (SiO₂,
Al₂O₃, TiO₂, Fe₂O₃ and K₂O; the remaining XRF data can be found in the Supplementary Material).

174 XRF analyses were performed exclusively on light and dark laminae sub-sampled from section III.

The grain-size of each bulk sediment sample was determined using a Malvern Mastersizer
Hydro2000S Diffraction Laser unit for the <2 mm size fraction. Sand, silt and clay fractions were
determined using the grain-size classification proposed by Udden-Wentworth (1922).

178 All bulk data not discussed in detail in the main text are presented in the Supplementary Material.

179

180 3.3 HBIs

181 Sediments were extracted, purified and analysed using the analytical method described previously (Belt et al., 2019). Briefly, prior to extraction, 9-octylheptadec-8-ene (9-OHD; ca. 0.1 µg) was added 182 183 to sediments as an internal standard to enable quantification of IPSO₂₅ by gas chromatography-mass 184 spectrometry (GC-MS). Sediments were subsequently saponified in methanolic KOH (H₂O/MeOH, 185 1:9; 5% m/v KOH) for 60 min at 70°C. After extraction with hexane (3 x 3 ml), the centrifuged supernatant containing HBIs was dried under a N₂ stream. The dry extract was re-dissolved in hexane 186 187 (500 μ l) and purified using open column chromatography (SiO₂, 38-63 μ m). The eluted hexane fraction (3 x 2 ml) was dried under a N₂ stream and re-dissolved in 300 µl of hexane prior to GC-MS 188 analysis. 189

HBIs were quantified via GC–MS using an Agilent 7820a chromatograph fitted with a J&W DB5-MS
column (30 m length, 0.25 mm i.d., 0.25 µm film thickness) coupled to a 5977b Mass Selective
Detector (MSD). The oven temperature ramp was programmed from 60°C to 280°C at 10°C/min.
During the ramp, the MSD operated in both Selective Ion Monitoring (SIM) and SCAN modes.

The sea ice algae-derived biomarker IPSO₂₅ (HBI diene II) and a tri-unsaturated HBI made by certain
open water diatoms (Belt *et al.*, 2017), referred to here as HBI III, were identified by comparison of

their mass spectra with those published in previous studies (Belt, 2018; Belt et al., 2012). The injection 196 197 of C8-C40 alkanes (Sigma-Aldrich) was used to further check the retention indices of IPSO25 and HBI 198 III. Quantification of IPSO₂₅ and HBI III (Fig. S2) was achieved by integrating peaks of ions m/z199 348.3 and 346.3, respectively, in SIM mode followed by normalizing to the corresponding peak area 200 of the internal standard (9-OHD) and an instrumental response factor obtained by analysis of a 201 purified standard. Data are presented as ng/g opal since both biomarkers are only produced by diatoms 202 and to circumvent the effect from dilution with lithogenic particles and other sources of OC. For 203 example, other algae common to Antarctica such as Phaeocystis Antarctica are within the non-silica (and non-HBI) producing Prymnesiophyte class, and may potentially contribute to the total OC 204 205 (Arrigo et al., 2000; Arrigo and van Dijken, 2004). Normalisation of IPSO₂₅ to the opal content thus 206 eliminates such possible types of dilution effect. In any case, we also provide our HBI biomarker data 207 in ng/g OC and ng/g dry sediment in the Supplementary Material (Fig. S3 and S4).

208

209 3.4. Diatom analyses

210 An aliquot of each dry sediment (ca. 0.2 g) was treated in a beaker containing a solution of distilled H_2O (40 ml for each sample), H_2O_2 (60 ml for each sample; concentration 40%) to dissolve organic 211 matter, and Na₄P₂O₇ (100 mg) to disaggregate sediment particles. The suspensions were heated (70° C; 212 213 45 min), followed by the addition of 10 ml of HCl (concentration 10%). Suspensions were then heated 214 $(70^{\circ} \text{ C}; 15 \text{ min})$ and rinsed repeatedly with distilled water in order to reach a pH of ca. 5–6. Rinsing 215 was performed every 8 hours to allow diatoms to settle. The resulting suspensions were then reduced 216 to a volume of 50 ml. A coverslip was placed inside a petri dish and a known volume of suspended 217 material (ca. 150–300 μ L) was pipetted into it, together with distilled water in order to achieve a 218 heterogeneous distribution of diatoms on the surface of the dish. After removing excess water, 219 coverslips were glued to microscope slides using the Norland Optical Adhesive 61 (NOA61) and dried 220 under UV light. At least 300 diatom frustules were counted for each slide following the method 221 proposed by Crosta and Koc (2007), which revised Schrader and Gersonde (1978) and Armand (1997). Corethron pennatum was counted as one when more than half a valve was present. The 222 223 relative abundance of each diatom species in a sample was calculated as the percentage of valves of a

given species in relation to the total number of valves counted in each sample. In addition, relative biovolume contribution was calculated following the equation and individual biovolumes proposed for each species by Alley *et al.* (2018).

227

228 3.5. Chronology

A detailed description of HLF17-1 chronology is presented in section 4.3. Here, we present the 229 230 analytical methods used to constrain the different dated horizons encompassed by the age-depth model (Table 1). The top of HLF17-1 core was dated using the short-lived radionuclide ²¹⁰Pb (maximum 231 penetration depth, Fig. S5). ²¹⁰Pb activity was derived from its daughter nuclide ²¹⁰Po via alpha 232 233 spectrometry, following the procedure presented elsewhere (Frignani et al., 2005). XRF core scanning 234 data (0.5 cm resolution) were used to identify possible cryptotephra layers within the core. A 235 significant anomaly in some element ratios (i.e. Zr/Sr and Nb/Sr) was found at 136.5 cm 236 corresponding to a peak in volcanic ash concentration (cryptotephra). The bulk layer was treated in 237 H₂O₂ (40% concentration) to dissolve organic matter and with HCl (10% concentration) to dissolve 238 carbonate. Resulting sediment was impregnated in epoxy resin and the textures and composition of glass particles were studied at the Istituto Nazionale di Geofisica e Vulcanologia, Sezione di Pisa 239 (INGV-Pisa) using a scanning electron microscope (SEM), Zeiss EVO MA coupled with Oxford-240 241 Aztec Energy EDS Analysis System. Standards of volcanic glass were analysed to test the accuracy of 242 data during the SEM analyses (Fig. S6).

Radiocarbon dating of bulk OC (*n*=5) and carbonate samples (*n*=6) was performed via accelerated
mass spectrometry (AMS) at The National Ocean Sciences Accelerator Mass Spectrometry (Woods
Hole Oceanographic Institution, USA) and at the Poznan Radiocarbon Laboratory (Poland). The
Bayesian age-depth model was constructed in R using the Bchron package (Haslett and Parnell, 2008).

247

248 3.6. Statistical analyses

T-tests were used to determine whether the populations of dark and light laminae were statistically different from each other at p < 0.01 (Excel, Analysis ToolPak). When the variance was different among populations, we ran the test assuming unequal standard deviation. Complete linkage

agglomerative hierarchical clustering (AHC) was carried out using IPSO₂₅, δ^{13} C (OM), and *C. pennatum* (% biovolume) as descriptive variables in order to determine their ability to separate light and dark laminae into distinct groups. Subsequently, the same explanatory variables were used to build a classification tree (CT) model via the Classification and Regression Trees (CART) algorithm of Breiman *et al.* (1984) to determine thresholds that best separated light and dark laminae. Primary and surrogate splits were considered, and the CT was pruned by minimizing classification accuracy after 10-fold cross-validation (Fig. S7).

259

260 **4. Results and discussion**

This section initially focuses on the origin of dark and light laminae, followed by the reconstruction ofcentennial-scale fast ice variability in the Edisto inlet throughout the late Holocene.

263

264 **4.1 Dark and light laminae composition**

Core HLF17-1 exhibited well-defined dark and light laminae on a mm- to cm-scale throughout the record. In order to understand the origin of laminations and their relationship with sea ice, we subsampled individual laminae from section III of core HLF17-1 (n=34, Fig. 2) since these were sufficiently thick to be collected without cross-contamination from adjacent sections. Overall, *t*-test analyses of IPSO₂₅, geochemical and taxonomic data revealed that dark and light laminae were statistically different from each other with, however, important differences depending on the parameter (Table 2).

272 For example, the mean IPSO₂₅ concentration was much higher in the dark laminae compared to the 273 light laminae (Table 2; p < 0.01) (Fig. 3b). We interpret this as an indication of the opening of the inlet 274 during the summer ice break-up following the spring sympagic bloom, resulting in relatively high 275 IPSO₂₅ concentrations in the dark laminae. In contrast, we suggest that lighter sediments with low 276 IPSO₂₅ reflect largely (later) ice-free conditions, with lower contributions from sympagic algae. Related inferences were made previously by Massé et al. (2011) following analysis of IPSO₂₅ in 277 laminated near-surficial sediments from Adélie Land (East Antarctica). Thus, alternating IPSO₂₅ 278 279 concentrations were identified in contrasting light/dark laminated sediment sections believed to

represent varying seasonal input; however, the precise seasonal characteristics of the individual
laminae were not confirmed as part of the study by Massé *et al.* (2011) and the relationship between
IPSO₂₅ concentration and lamina colour/brightness was also not reported.

If our interpretations of variable IPSO₂₅ in laminated sections from core HLF17-1 are correct, the 283 distribution of diatoms would be expected to follow the lamina colour in line with the IPSO₂₅ data. 284 Instead, the *t*-test showed that, with the exception of *Fragilariopsis obliquecostata* (p < 0.01), 285 286 differences between dark and light laminae were not statistically significant for the species identified, 287 and the differences observed for F. obliquecostata were, in any case, small (Table 2). This contrasts 288 with a qualitative examination of the light laminae, which gave a visual impression of being 289 dominated by C. pennatum mats when compared to the dark laminae. Similar observations were made 290 by Alley et al. (2018) in laminated sediments collected in Iceberg Alley (East Antarctica). However, 291 Alley et al. (2018) also noted that C. pennatum is a relatively large diatom, so its abundance, relative 292 to the total number of organisms, can lead to underestimations with respect to biomass or biovolume.

293 Thus, once the relative concentrations of diatoms in core HLF17-1 were re-calculated on a biovolume 294 basis (calculations based on Alley et al., 2018, Table 3), C. pennatum became the most abundant species in the light laminae (Fig. 3c; Table 3), supporting the visual qualitative examination, and low 295 IPSO₂₅ concentrations, as predicted. In fact, since C. pennatum can adjust its buoyancy to uptake 296 297 nutrients beneath the pycnocline, C. pennatum-rich sediments likely accumulate towards the end of the 298 summer when surface waters are ice-free, well-stratified and nutrient depleted (Alley et al., 2018; Leventer et al., 2006; Salter et al., 2012), again, consistent with the low IPSO₂₅ values. In contrast, for 299 300 dark laminae, where IPSO₂₅ is generally high, the deposition of phyto-detritus is likely associated with 301 the progressive accumulation of sympagic diatoms, as fast ice retreats in early summer, in combination 302 with the first algal blooms, as further evidenced by increases to well-known sea ice associated species 303 such as *F. obliquecostata* and *F. curta* (Table 3).

Consistent with this interpretation, the average stable carbon isotopic composition (δ^{13} C) of the OM in the dark laminae was relatively enriched in ¹³C compared to the light laminae (Table 2; *p*<0.01) (Fig. 306 3d). Indeed, we propose that the isotopic fingerprint of laminae likely reflects the influence of variable sea ice cover on the overall photosynthetic fractionation of CO₂, especially since it is well-known that

sea ice-derived OM is normally relatively enriched in ¹³C compared to OM produced in pelagic 308 309 settings (Massé et al., 2011). Under normal CO2-replete conditions, such as those associated with cold open waters, preferential uptake of ¹²CO₂ during photosynthesis leads to isotopically light OM; 310 however, within the confined environment of semi-enclosed sea ice, CO₂ becomes a limiting substrate, 311 with consequential reduction in isotopic fractionation and OM that becomes relatively enriched in ¹³C 312 (Geilfus et al., 2014; Geilfus et al., 2015). This effect can be augmented by the formation of melt 313 ponds and percolation of meltwater into the ice matrix during ice brake-up, which can further decrease 314 315 the dissolved CO_2 concentration (Geilfus *et al.*, 2015), while the uptake of "heavy" bicarbonate as an 316 alternative carbon source further increases δ^{13} C (OM) values (Lehmann *et al.*, 2004).

317 Previous studies have suggested that other factors can also regulate the $\delta^{13}C$ of phytoplankton 318 including nutrient availability, growth rate and morphology (i.e. volume/surface cell ratio) (Crosta et al., 2005; Popp et al., 1999; Popp et al., 1998). While we cannot comment on the quantitative 319 importance of the first two aspects in the current context, we note that some previous in-situ and batch 320 culture studies indicate that the fractionation of ¹³C during photosynthesis decreases with increasing 321 322 volume/surface area ratio of the diatom cell (Crosta et al., 2005; Popp et al., 1998). However, in the 323 current study, the most depleted δ^{13} C values for OM were found in the light laminae dominated by C. *pennatum*, which is characterised by a high volume/surface area ratio. Together, these data support our 324 325 hypothesis that the progressive increase in CO_2 concentration in surface waters upon transitioning from ice-covered to ice-free conditions likely exerted first-order control on the $\delta^{13}C$ signature of 326 diatomaceous OM, with most depleted δ^{13} C values aligned with light laminae associated with summer 327 pelagic conditions. Conversely, dark laminae contain a higher contribution from ¹³C-enriched 328 329 sympagic OM following ice break-up and higher IPSO₂₅. Consistent with this, although we did not measure δ^{13} C for IPSO₂₅ in the current study, this biomarker has been previously been shown to be 330 significantly ¹³C-enriched in both sea ice and sediments ($\delta^{13}C = ca. -5$ to -18 ‰; Belt *et al.*, 2016, and 331 references cited therein). 332

We note that the diatom *Berkeleya adeliensis* was not observed in any of the laminae analysed, despite the identification of IPSO₂₅ in all cases. This might be a result of the occurrence of production of

 $IPSO_{25}$ by other, as yet unidentified, sources; however, HBIs (including $IPSO_{25}$) are only produced by 335 certain diatoms (see Belt, 2018 for a recent review) and none of the other species identified here 336 337 (Table S1) are known to biosynthesise IPSO₂₅. Alternatively, the absence of *B. adeliensis* in all laminae may simply reflect its poor preservation in the water column and in sediments (due to lightly 338 silicified valves) following its release from fast ice, as reported previously (Riaux-Gobin et al., 2011; 339 Tanimura et al., 1990). Indeed, Tanimura et al., (1990) showed that B. adeliensis can account for a 340 341 large fraction of diatom assemblages in sea ice and melt ponds, yet is frequently absent in surface sediments due to dissolution. In addition, we note that some of the C. pennatum observed in the 342 current study was present as fragmented frustules. On the other hand, IPSO₂₅ appears sufficiently 343 344 stable in sediments to permit identification in sediments spanning the Quaternary, at least (Belt, 2018, 345 2019).

346

347 The distributions of lithogenic material are also consistent with the biogenic content and associated interpretations of dark and light laminae. Specifically, dark laminae, on average, showed higher 348 349 concentrations of lithogenic elements, including TiO₂, Al₂O₃ and Fe₂O₃, and K₂O, compared to the light laminae (Table 2; p<0.01) (Fig. 3e). Following our interpretations based on IPSO₂₅ concentration, 350 diatom assemblages and δ^{13} C (OM), we suggest that the relatively higher lithogenic content in the dark 351 laminae likely reflects the opening of the inlet, when fast ice thaws and releases mineral dust deposited 352 on sea ice during the preceding months of the year (Atkins and Dunbar, 2009). In fact, according to 353 354 Atkins and Dunbar (2009), the flux of dust that accumulates over the sea ice in McMurdo Sound (Ross 355 Sea) decreases with increasing distance from the coast and accounts for a large fraction of the total 356 lithogenic particle flux to sediments.

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358 4.2 Annual variability and lamination pattern

Overall, our collective biogenic and geochemical proxy data obtained from well-defined laminae suggest that fast ice evolution during summer in the Edisto inlet resulted in the deposition of laminae possessing different colour and composition. Further, by co-plotting the IPSO₂₅, δ^{13} C (OM) and *C*. *pennatum* (% biovolume) datasets (Fig. 4), the two proposed dominant fast ice patterns can be

visualized and summarized as follows. At the beginning of summer, the progressive opening of the 363 inlet takes place, leading to the accumulation of dark laminae characterized by high IPSO₂₅, enriched 364 δ^{13} C (OM) and low C. pennatum (% biovolume) (Fig. 4). We infer that the biogenic material that 365 366 forms these darker laminae has two main sources: (i) sympagic diatoms living within the brine matrix 367 of sea ice, which are subsequently released as the ice thaws, and (ii) diatoms that flourish in surface 368 waters as the inlet opens in summer (e.g. F. curta). Further, since satellite images show that the 369 complete opening of the inlet can take a few weeks, we believe that the deposition of sea ice diatoms 370 and sea ice-associated diatoms likely coexist. This is confirmed by the co-occurrence of IPSO25 and 371 HBI III in the dark laminae (Fig. S3), the latter being derived from certain pelagic diatoms (e.g. (Belt et al., 2017) and commonly found in marginal sea ice zones (Belt et al., 2015; Smik et al., 2016; 372 Vorrath et al., 2019). 373

When protracted opening of the inlet occurs (i.e. ice free conditions), the surface nutrient limitation 374 375 likely favours diatoms that can uptake nutrients underneath the pycnocline typical of oligotrophic 376 waters which develop towards the end of the summer (Alley et al., 2018). Under these conditions, C. 377 pennatum dominates, generating light laminae (Fig. 4), possibly in the form of rapidly sinking mats 378 bound together by exopolymer mucus secretions and entangled spines (Alley et al., 2018). Our 379 interpretation would clearly benefit from contemporary in situ studies dealing with the composition 380 (diatoms, biomarkers, etc.) of seasonal sinking particulate material inside Antarctic inlets. To the best 381 of our knowledge, however, sediment traps have been deployed for this kind of setting only for short periods (Thomas et al., 2001), although longer studies in the Southern Ocean have been carried out 382 383 primarily in deep- and mid-shelf environments (Smith et al., 2000)..

Our general description of two discrete sea ice scenarios can be refined further by reference to satellite images of the inlet, which reveal marked inter-annual variability of fast ice conditions in summer. For example, Figure 5 shows satellite images from three years (*i.e.* 2012, 2017, 2018) that clearly describe the full spectrum of proposed ice cover and its transitions. Thus, in 2017, ice break-up took place at the beginning of the summer, followed by protracted opening of the inlet. Under these conditions, we infer the deposition of both dark and light laminae (Fig. 5a). In contrast, in 2012, thawing of fast ice was delayed, and the inlet never completely opened by the end of the summer. In this scenario, it is

likely that deposits were dominated by dark laminae due to only partial and ephemeral opening of the
inlet (Fig. 5b). Finally, in 2018, fast ice persisted throughout the summer along the coastal region. We
envision that these circumstances prevented the accumulation of biogenic material or, alternatively,
resulted in only a thin deposit (*i.e.* condensed unit) (Fig. 5c).

Overall, satellite images from the Edisto inlet taken over the last decade suggest that fast ice dynamics during summer can sometimes be more complex than a simple sequential accumulation of dark and light laminae, as inferred for 2017. This is confirmed by the laminations patter of HLF17-1 that does not exhibit recurring patterns at multi-year scale (*i.e.* over cm or dm) (Fig. 2), likely reflecting the dynamic annual variability in summer sea ice conditions revealed by satellite images shown herein from 2012, 2017 and 2018.

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402 **4.3.** Local reservoir effect and Bayesian age-depth model

403 Calibrations of radiocarbon ages obtained from carbon pools that are not in equilibrium with the atmosphere require a correction due to the influence of the reservoir effect (Stuiver and Polach, 1977). 404 405 In addition to the global mean marine reservoir (R) modelled when using the Marine13 curve (Reimer 406 *et al.*, 2013), a further offset (ΔR) needs to be applied to accommodate local influences. Here, we built our age-depth model using ¹⁴C ages derived from both carbonate and organic carbon matrices, with 407 additional dates obtained from excess ²¹⁰Pb and one tephra horizon. Considering the mixed nature of 408 the radiocarbon tests (i.e. organic and inorganic carbon), we used two different local reservoir 409 410 corrections prior to calibration. Based on U/Th dating of coral samples trapped by the fringing Ross Sea ice shelf (Hall et al. (2010), the ΔR value of the carbonate matrix (ΔR_{CaCO3}) in the Ross Sea has 411 412 remained relatively stable ($\Delta R_{CaCO3} = 791 \pm 121$ yrs) over the last 6,000 years. In our records, carbonate 413 tests included benthic foraminifera, planktonic foraminifera and Ophiuroidea found in core HLF16-1 (Table 1), and were correlated with core HLF17-1 using distinct lamination patterns common to both 414 415 cores (Fig. S1, Supplementary Material). For consistency, only benthic foraminifera were used in the 416 final age model, while additional carbonate samples (planktonic foraminifera and Ophiuroidea) from 417 the same core depths were used to independently test the robustness of our chronology and

assumptions behind using ΔR_{CaCO3} (Table 1). Following calibration, all carbonate skeletal elements belonging to the same horizon displayed similar calendar ages, confirming that the ΔR_{CaCO3} , in addition to being stable over the late-Holocene (Hall *et al.*, 2010), is also appropriate for diverse calcifying organisms (Table 1).

422 In contrast, the ΔR of the acid insoluble organic fraction (ΔR_{OC}) is poorly constrained in Antarctic sediments and is usually derived empirically by radiocarbon dating of surface bulk sediments and their 423 interpolation (Domack et al., 2001; Licht et al., 1998; Mezgec et al., 2017). In the Edisto inlet, Mezgec 424 425 et al. (2017) used core-top data (box-corer BAY05-bc21, 0-1 cm) to propose a value of 1580±45 years 426 as a general R_{OC} (global + ΔR). However, using the same approach, the remarkably young ¹⁴C age 427 (200±30 yrs BP; Table 1) of the surface sediment from box core HLF17-2BC (0-0.5 cm) collected at the same site as piston core HLF17-1, suggests that our core top likely contained "bomb" ¹⁴C, which 428 429 prevents its use to assess ΔR_{OC} .

430 Given the evident limitations of using surface sediments to estimate the local reservoir effect, we derived the ΔR_{OC} by pairing benthic foraminifera (ultimately used in the age-depth model) with 431 432 organic carbon from the same horizons, relying on the well-constrained ΔR_{CaCO3} . This alternative approach assumes that the accumulation of foraminifera and the deposition of organic carbon in the 433 same horizons were coeval with negligible post-depositional transport. This allowed us to subtract the 434 ¹⁴C age of foraminifera (adjusted for ΔR_{CaCO3}) from the ¹⁴C age of the organic fraction to obtain the 435 ΔR_{OC} for each level and its relative uncertainty. The average of the three values gave an estimate of the 436 final mean ΔR_{OC} (1320 ± 135 yrs, 1 σ). Additional dates used in the age-depth model include the 437 maximum penetration of excess ²¹⁰Pb (150±20 yrs before 2017, Fig. S1 and S4; ca. seven half-lives 438 439 according to Arias-Ortiz et al. (2018)) and a tephra layer recently found in the Edisto inlet sediments (Fig. S1 and S5) (Mount Rittmann volcano eruption, (Di Roberto et al., 2019), which has been 440 identified in the Talos Dome core (696 ± 2 yrs BP; ice record) (Narcisi et al., 2012), Taylor Dome core 441 442 $(709 \pm 71 \text{ yrs BP}; \text{ ice record})$ (Hawley *et al.*, 2003) and West Antarctic Ice Sheet (WAIS) Divide cores $(687 \pm 7 \text{ yrs cal BP})$ (Di Roberto *et al.*, 2019). For our final age-depth model, we used the more precise 443 444 age assignment derived from the annually-counted portion of the WAIS Divide 2014 chronology.

Our final Bayesian age-depth model for HLF17-1 was built on seven dates (Table 1; Fig. 6). The agedepth model showed that, since 2.6 ka BP, the relatively stable and high accumulation rate in the inlet
(ca. 0.7 cm/yr) was followed by an abrupt decrease around 0.7 ka BP (*ca.* 0.2 cm/yr) (Fig. 6).

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449 4.4. Late-Holocene reconstruction of sea-ice dynamics in the Edisto inlet

450 In order to reconstruct the fast ice dynamics during the late-Holocene, we analysed 295 sediment 451 samples throughout piston core HLF17-1. Following our interpretation about the emplacement of laminated strata (section III), we assumed that the down-core distribution of IPSO₂₅ and $\delta^{13}C$ (OM) in 452 453 bulk sediments co-varied as a function of the relative contribution of light and dark laminae. Since the two types of laminae could be further classified, statistically, into two distinct groups based on AHC 454 455 and CT analyses (see Section 3), we identified suitable thresholds for IPSO₂₅ (0.42 ng/g opal) and δ^{13} C (-27.56‰) to best separate the laminae analysed in section III (Fig. 4) into light and dark groupings 456 457 (Supplementary Material, Fig. S6). By applying these thresholds to bulk sediments, we obtained two sub-groups of IPSO₂₅ and δ^{13} C (OM) which succinctly represented the two dominant types of fast ice 458 459 coverage in summer: initial summer break-up and ice-free conditions (dashed lines in Fig. 7a,b). As described previously (Section 2), we expected that the third condition - inlet covered throughout the 460 summer - would not be uniquely characterised in our laminated sequence. Rather, when this occurred, 461 462 permanent fast ice probably resulted in either a gap in accumulation or a condensed sediment unit. 463 Regardless of this limitation, comparison between the down-core sediment brightness and clustered IPSO₂₅ and δ^{13} C (OM) bulk data revealed a good agreement, especially for IPSO₂₅ (Fig. 7a,b). This 464 465 further confirmed that the correlation between sediment colour and geochemical signature observed in section III (laminae) holds throughout the record. 466

467 Thus, we propose use of the same IPSO₂₅ and δ^{13} C thresholds (dashed lines Fig. 7a,b) to identify 468 periods throughout the late Holocene when fast ice break-up followed by an ice-free inlet was more 469 frequent (grey shaded areas) relative to periods characterized by partial opening of the inlet (see Fig 5a 470 and 5b, respectively). Using this approach, we identify a significant change in the relative frequency of 471 inlet opening and ice-free waters during the late-Holocene. Specifically, our results suggest that

between 2.6 ka BP and 0.7 ka BP the inlet opened regularly during summer, while after 0.7 ka BP, 472 there was an abrupt shift characterized by relatively less frequent ice-free conditions (e.g. Fig. 7a,b). It 473 474 is worth noting that following the shift in ice conditions at 0.7 ka BP, the sediment accumulation rate 475 of the record decreased abruptly from 0.7 to 0.2 cm/yr. This likely occurred due to the absence (or lower frequency) of light laminae deposition, resulting in relatively thinner sediment strata over 476 477 summers. Finally, the bulk grain-size exhibited a similar temporal shift, becoming progressively 478 coarser after ca. 1.3 ka BP. This overall trend was expected based on the analysis of dark and light 479 laminae, which also displayed small, but statistically significant differences (Table 2; p<0.01), 480 although the reason why dark laminae were coarser is, as yet, unknown. We speculate that this might 481 be attributable to the relatively higher concentration of lithogenic material in the dark laminae (mode of dust ranges between 76 and 129 µm, (Atkins and Dunbar, 2009). Nevertheless, the difference 482 between light and dark laminae is somehow counterintuitive considering that the former are dominated 483 484 by C. pennatum, which, as described before, is considerably larger than other diatoms more abundant in dark laminae. However, since C. pennatum was mostly present as fragments, despite its high 485 abundance, it is possible that lithic particles played a major role in the overall grain-size distribution. 486

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488 4.5. Fast ice dynamics over the late-Holocene and climate forcing

489 In order to understand the origin of the shift in sea ice conditions observed around 0.7 ka BP, we 490 compared our results with the extensive dataset compiled by Stenni et al. (2017) within the umbrella 491 of the PAGES Antarctica2k programme. The entire database consists of water stable isotope data $(\delta^{18}O \text{ and } \delta D)$ from 112 records, which permitted a high-resolution reconstruction of past temperature 492 and relative anomalies (relative to 1900-1990 CE) over the last 2 ka BP for seven climatically distinct 493 494 regions of Antarctica. The composite temperature anomalies reconstruction (10-yr-binned averages; 495 Fig. 7d) for the Victoria Land Coast region based on ice cores near to HLF17-1 (Fig. 1a) shows a clear 496 and abrupt cooling at 0.7 ka BP, which follows the general long-term cooling over Antarctica that 497 started at ca 1.2 ka BP. Previously, Stenni et al. (2017) argued that this hemispheric-scale cooling 498 could have been driven by major volcanic eruptions, in agreement with previous studies (McGregor et 499 al., 2015). Although the cause of this cooling over the last two millennia is beyond the scope of the

500 current study, it is worth noting that the land fast ice reconstruction based on our marine record agrees 501 well with the temperature anomalies recorded in the Victoria Land Coast. Thus, we attribute less 502 frequent ice-free conditions since 0.7 ka BP to a colder climate inferred from ice core data (Stenni *et* 503 *al.*, 2017), although we also note that our marine record resolves summer dynamics only, while ice 504 cores provide a year-round signature.

We also compared our results with those of Mezgec *et al.* (2017) who analysed Holocene diatom assemblages in cores taken from Cape Hallett and Wood Bay (Ross Sea). Interestingly, their data show a sharp increase in the relative concentration of *F. curta* during the late Holocene, which is consistent with the occurrence of more frequent dark laminae in the current study (Table 2 and 3), thus supporting our interpretation of less protracted opening of the inlet during summers along the coast.

Another important factor that can have large-scale implications for fast-ice dynamics over Antarctica 510 511 is the non-annular response (i.e. dipole) of the Southern Ocean to the Southern Annular Mode (SAM) 512 (Bertler et al., 2018; Lefebvre and Goosse, 2005; Lefebvre et al., 2004). This dipole consists of a different regional response in sea ice, with decreases in the Weddell Sea and around the Antarctic 513 514 Peninsula, and an increase in the Ross and Amundsen Seas during years with a positive SAM index. 515 Due to a low-pressure anomaly in the Amundsen-Bellingshausen sector during positive SAM years, 516 the Weddell and Bellingshausen Seas are subject to more northerly winds, while the Ross Sea tends to 517 have more southerly winds (Lefebvre et al., 2004), inducing a significant cooling at the surface and an 518 increase in the Ross Sea ice cover (Lefebvre and Goosse, 2005; Lefebvre et al., 2004). However, as far 519 as our data are concerned, comparison between our reconstructed fast ice dynamics and the SAM 520 derived from proxy records over the last 1000 yrs (Fig. 7e) (Abram et al., 2014) does not suggest a 521 direct influence of the latter. In particular, the abrupt shift that characterizes our record around 0.7 ka 522 BP even opposes that expected from a negative SAM (Lefebvre et al., 2004). Likewise, the negative 523 SAM should also have resulted in warmer conditions over the Ross Sea, which does not concur with 524 the sharp cooling reconstructed in the Victoria Land Coast region based on ice core records (Stenni et 525 al., 2017) (Fig. 7d).

However, stronger westerlies in the region of the Antarctic Circumpolar Current during positive SAMinduce an intensified eastward surface ocean current and, as a result of Ekman drift, a stronger

northward surface current south of 45°S. This promotes upwelling over the margin of the Modified 528 529 Circumpolar Deep Water, whose characteristics - warm and salty - can promote sea ice thawing in 530 the Ross Sea despite the general wind-driven surface cooling (Lefebvre and Goosse, 2005; Lefebvre et al., 2004). Thus, the negative SAM during the late-Holocene might have reduced the upwelling with 531 direct effects on fast ice thawing. However, considering the small age uncertainties around the tephra 532 layer (687±7 yrs BP), the shift in our record almost certainly occurred before the change in the SAM 533 534 index. Therefore, the role of upwelling on fast ice dynamics remains somewhat elusive, although a 535 negative SAM after 0.7 ka presumably contributed to some preservation of coastal sea ice throughout the summer. Overall, we believe that the abrupt atmospheric cooling recorded over the Victoria Land 536 537 Coast region (Stenni et al., 2017) likely exerted first-order control on the rapid shift of fast ice coverage in the inner Ross Sea during the late-Holocene, but other factors are more challenging to 538 identify at this stage, in part, owing to the complex behaviour of sea ice cover, more generally (Meehl 539 540 et al., 2019).

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542 **5.** Conclusions

This study provides the first high-resolution late-Holocene reconstruction of fast ice dynamics in the 543 Ross Sea (Edisto inlet) based on multiple proxies (IPSO₂₅, diatom census counts and bulk properties) 544 545 analysed in a 14.6 m long laminated diatom ooze record (HLF17-1). Our results indicate that the 546 emplacement of laminated strata in the Edisto inlet reflect different fast ice cover during the summer months. In early summer, fast ice break-up releases sympagic diatoms living within the ice and 547 548 platelet ice beneath. As the inlet opens, algal blooms in melt ponds promote sinking of phytodetritus, 549 which accumulates with the sympagic diatoms. Overall, the opening of the inlet generates dark 550 laminae characterized by relatively high concentrations of the sympagic biomarker IPSO₂₅, high lithogenic material and enriched δ^{13} C (OM) with a relatively heterogeneous distribution of diatoms. In 551 552 late summer, when ice-free conditions persist, the diatom Corethron pennatum can adapt to stratified and oligotrophic waters, becoming the dominant species and generating thick mats at the seabed. The 553 resulting composition of light laminae is characterized by low IPSO₂₅ concentrations, low lithogenic 554 material, and depleted $\delta^{13}C$ (OM). 555

556 Building on the knowledge gained at lamina level, our down-core results show an abrupt change in 557 summer fast ice dynamics over the late-Holocene. In particular, while the inlet appears to have opened 558 regularly during summers since 2.6 ka BP, light laminae became suddenly less frequent around 0.7 ka 559 BP, indicating an abrupt shift towards less recurrent ice-free conditions in the inlet. Comparison with ice core data from the region revealed that the abrupt shift in fast ice dynamics was likely the 560 561 expression of colder climate conditions. Our results, both at seasonal (laminae) and late-Holocene 562 scales, provide new insights into the application of the newly established IPSO₂₅ sea ice proxy, and in particular its usefulness for paleo reconstructions of summer fast ice dynamics. 563

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Captions Fig.1. Study area in the Ross Sea. (a) Map showing the 40-y average of sea ice concentration (%) in February based on satellite images. The red line displays the 10% contour. The location of HLF17-1 core and ice cores of the Victoria Land Coast is shown with filled red and green circles, respectively; (b) Red filled circle shows the location of HLF17-1 in the Edisto inlet; (c) Chirp profile showing the sediment strata geometry. Parallel reflectors show continuous sedimentation throughout the Holocene Fig 2. HLF17-1 piston core (14.65 m) from top (section XV) to bottom (section I). Fig. 3. Dark and light lamina analysis in section III. (a) Brightness (green line) and sub-sampled horizons (n=34); relative frequency of (b) IPSO₂₅, (c) Corethron pennatum (d) δ^{13} C and (e) TiO₂ in the light (red open bars) and dark (gray filled bars) laminae, respectively Fig. 4. Composition of laminae sub-sampled in section III. Blue and red dashed lines display the two dominant clusters which reflect contrasting sea ice coverages, namely fast ice thawing and ice-free conditions, respectively. Fig. 5. Satellite images (a, 2017; b, 2012; c, 2018) of Edisto inlet (red ellipse) which provide the spectrum of sea ice coverage which, in turn, exerts first-order control on the formation of laminated sediments. 2017 was characterized by an early opening followed by protracted ice free conditions of the inlet. In 2012, thawing was incomplete and occurred toward the end of the summer. Finally, 2018 represented the end of the spectrum as sea ice persisted throughout the summer in the inlet

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Fig. 6. Median calendar age (years before present) and 2σ error from the Bayesian age-depth model of core HLF17-1. Gray colors show the dates used in the model. Additional carbonate dates (ophiurae and planktonic foraminifera) were used to test, independently, the age-model and the assumption behind the local marine reservoir age of the carbonate fraction.

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Fig. 7. Fast ice dynamics in the Edisto inlet and general climate conditions over the late-Holocene. 618 619 Top filled blue circles show the dated horizons used in the age-depth model (a) IPSO₂₅ and brightness 620 (11-point weighted average to match the IPSO₂₅ resolution), red and blue filled circles were defined 621 according to the cluster analysis (see text for further details); (b) δ^{13} C and brightness (11-point 622 weighted average to match the IPSO₂₅ resolution), purple and pink filled squares were defined 623 according to the cluster analysis (see text for further details); (c) sediment accumulation rate and grainsize; (d) 10-yr-binned averages of temperature anomalies in the ice cores (relative to 1900–1990 CE) 624 (brown symbols) from the Victoria Land Coast ice cores (see Fig.1a) and 5-point weight average 625 626 (yellow line) (Stenni et al., 2017); (e) Reconstruction of annual Southern Annular Mode index (7-yrbinned average, blue line and 70-yr loess filter, red line) (Abdram et al 2014). Grey boxes define 627 628 period characterized by regular opening of the inlet over summer (e.g. Fig. 5a).

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850 Fig 1.



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853 Fig. 2



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856 Fig. 3





859 Fig. 4



860

862 Fig. 5



863 dark & light laminae

only dark laminae

lack of accumulation

865 Fig. 6



868 Fig. 7



871 Highlights

- 872 Dark and light sediment laminae exhibit different chemical and ecological features
- 873 Dark and light sediment laminae reflect different fast ice coverage during summer
- Enriched δ^{13} C and high IPSO₂₅ (dark laminae) track early thawing of fast ice
- 875 High concentration of *C. pennatum* (light laminae) identifies protracted ice-free conditions
- 876 Summer ice-free conditions became less frequent since 0.7 ka BP along the north-western Ross Sea

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core	horizon	depth (cm)*	depth range (cm) type of sample		ID lab	dating method	age (y)	error (y)	modelled median age	minimum 95% CI	maximum 95% CI
HLF16-1	VII 0-10	402	±5	benthic forams**	Poz-110839	^{14}C	2330 yBP	30	1083	927	1253.65
HLF16-1	VII 0-10	402	±5	ophiuroideae	Poz-110430	¹⁴ C	2220 yBP	30			
HLF16-1	VII 0-10	402	±5	bulk OC	Poz-110836	^{14}C	2890 yBP	30			
HLF16-1	IV 60-70	775	±5	benthic forams**	OS-147514	^{14}C	2800 yBP	15	1589	1387.35	1778.65
HLF16-1	IV 60-70	775	±5	bulk OC	Poz-111163	^{14}C	3210 yBP	35			
HLF16-1	I 70-80	1091	±5	benthic forams**	Poz-110838	^{14}C	3200 yBP	30	2055.5	1853.45	2260
HLF16-1	I 75-75.5	1091.25	±0.25	ophiuroideae	Poz-110431	¹⁴ C	3250 yBP	30			
HLF16-1	I 70-80	1091	±5	planktic forams	OS-147503	^{14}C	3300 yBP	35			
HLF16-1	I 70-80	1091	±5	bulk OC	Poz-110837	^{14}C	3875 yBP	30			
HLF17-2BC	0-1	0.5	±0.25	bulk OC	Poz-108920	^{14}C	200 yBP	30			
HI F17-1	XV 8-10	9	+0.5	bulk OC**	OS-141035	¹⁴ C	1940 vBP	15	58	5	134
HI F17-1	XV 18 20	10	±0.5 +1	sadimant**	05-141055	210 Db	150 v bafora 2017	20	154	110	100
HLF17-1	AV 10-20	19	±1	Sediment		FU	150 y belore 2017	20	134	117	190
HLF17-1	XIV 72-73	136.5	±1	sediment**		tephra***	687	7	687	6/5	698
HLF17-1	I 90-91	1456	±0.5	bulk OC**	Poz-92969	¹⁴ C	4220 yBP	50	2623.5	2353.8	2868.1

Table 1. Dated horizons using AMS, $^{\rm 210}{\rm Pb}$ and tephra layer

* depth refers to HLF17-1

** used in the age depth model

*** Di Roberto et al., 2019

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Table 2. Composition of laminae from section III of HLF17-1. Data were grouped according to the lamina colour: light vs dark. Statistically significant differences between groups for each variable were assessed based on T-test (p<0.01). Fragilariopsis obliquecostata Fragilariopsis separanda Fragilariopsis cylindrus Corethron pennatum Fragilariopsis curta Chaetoceros RS O pa l por osit y cl a y Si O 2 0 C δ¹³ C IPS O₂₅ sil t Ti O2 Al₂ O₃ Fe₂ O₃ dep th Nt ot sa nd laminae

K

 $\overset{2}{\mathbf{0}}$

	cm	%	%	‰	%		ng/g opal	%	%	%	%	%	%	%	%	%	%	%	%	%	%
light		0	0						7	1		1	5								
HLF17 1 III 14-15	117 9.5	1 2	4 4	28 .6	43 .8	0.91	0.16	11 .2	5. 7	3. 2	3. 6	7. 3	6. 7	2. 9	9. 2	0. 8	59 .0	0. 4	2.7	1.8	1. 1
HLF17 1 III 17-18	118 2.5	0. 1 2	0. 5 2	27 .7	45 .2	0.92	0.31	13 .2	7. 1	9. 7	1 9. 0	2 6. 1	1 9. 0	1 2. 5	7. 7	1. 9	65 .3	0. 6	3.7	3.1	1. 0
HLF 17 1 III 20-21	118 5.5	0. 1 0	0. 4 8	- 29 .0	43 .7	0.93	0.23	13 .3	7 7. 3	9. 4	8. 2	5. 8	5 1. 2	3. 2	1 4. 3	3. 2	62 .7	0. 8	4.9	4.2	1. 1
HLF17 1 III 33-34	119 8.5	0. 1 3	0. 5 8	- 28	40	0.90	0.22	11	7 8. 6	1 0. 1	5. 3	1 1. 3	4 5. 2	1 0. 7	1 2. 9	0. 0	64 .4	0. 7	4.3	3.6	1.
HLF17 1 III 34 5-35	120	0. 1	0. 6	27	46	0.91	0.32	17	7 4.	8.	7. 0	8. 2	3 3. 4	5.	1 4. 3	1.	67	0.	4.6	4.6	1.
HLF 17 1 III	121	0. 1	0. 4	- 30	47	0.04	0.32	18	7 2.	9.	4.	3 6.	2	7.	1 2.	2.	64	0.	1.0	4.0	0.
40-47 HLF 17 1 III	1.5	0. 1	0. 5	.4 - 29	.2 50	0.94	0.55	.5	5 7 5.	2 9.	3 7.	1 0.	5 6.	4.	6.	0.	.4 n.	n.	л.	2.4 n.	9 n.
51-52 HLF17 1 III	6.5 122	0 0. 1	8 0. 4	.3 - 29	.4 37	0.93	0.42	.3	2 7 8.	6 9.	4 9.	8 2 0.	8 3 6.	0 6.	1 1.	9 3.	m. 63	m. 0.	m.	m.	m. 0.
57-58 HLF 17 1 III	2.5 122	1 0. 1	5 0. 6	.5 - 28	.9 47	0.93	0.18	.7 12	9 7 8.	4 9.	5 8.	3 5.	1 4 0.	3 6.	0 1 2.	8 8.	.2 68	7 0.	3.5	3.6	8
63-64 HLF17 1 III	8.5 123	4 0. 1	3 0. 4	.6 - 29	.6 45	0.92	0.31	.5 13	5 7 7.	1 8.	3 9.	5 1 6.	1 4 5.	0 2.	1 9.	6 2.	.4 66	7 0.	3.6	3.4	0 0.
71-72 HI F17 1 III	6.5 123	2 0.	9 0. 6	.9 - 30	.1	0.94	0.21	.4	9 7 5	7	4 1 7	8	3 2 5	8	3 1 8	2	.4	4	2.3	1.9	7
73.5-74.5	9	3 0.	0	.0 -	.6	0.93	0.18	.5	4 7	0	4	0 4	0 2	3	2	3	.6	5	2.8	2.3	8
80.5-81.5	6	0	4 0 0.	.3	.2	0.92	0.20	.2	0 7	7	1. 3 1	4. 6 1	4. 2 3	1	5 1	8	.6	8	4.5	3.9	1.
HLF17 1 III 84-5-85.5	0	1 1 0.	5 6 0.	.8 -	44 .6	0.93	0.24	12 .9	7. 3 7	9. 9 1	2. 5 1	4. 0	1. 6 2	6. 4 1	1. 4	7. 8 1	68 .4	0. 6	3.6	3.1	1. 0
HLF 17 1 III 86.5-87.5	125 2	1 0 0.	5 1 0.	28 .4	37 .1	0.92	0.22	12 .9	6. 2 7	0. 8 1	4. 7 1	8. 3 1	4. 9 3	3. 4	8. 6	0. 7	68 .4	0. 6	3.8	3.0	0. 9
HLF17 1 III 91-92	125 6.5	1	4 8	28 .2	34 .1	0.91	0.42	12 .9	4. 7	2. 3	0. 5	8. 6	5. 8	1. 9	9. 6	1. 2	63 .2	0. 9	5.0	4.2	1. 2
	mea n	0. 1 2	0. 5 2	28 .8	42 .9	0.92	0.26	13 .6	7 6. 3	1 0. 0	9. 2	1 6. 9	3 6. 5	5. 9	1 1. 2	3. 8	64 .9	0. 6	3.7	3.2	1. 0
	s.d.	0. 0 2	0. 0 8	0. 9	5. 8	0.01	0.08	2. 1	1. 8	1. 4	5. 0	1 1. 4	1 2. 4	3. 7	3. 0	3. 3	3. 0	0. 2	0.8	0.9	0. 1
dark																					
HLF17 1 III 1-2	116 6.5	0. 1 4	0. 7 3	26 .4	36 .2	0.87	0.76	13 .4	7 9. 0	7. 6	3. 0	0. 8	4 8. 4	3. 4	1 9. 9	3. 4	65 .6	1. 1	6.0	5.3	1. 3
HLF17 1 III 3-4	116 8.5	0. 1 3	0. 7 4	26 .8	34 .7	0.89	0.42	14 .6	7 7. 5	8. 0	8. 0	1. 0	4 1. 6	3. 9	1 8. 4	4. 6	64 .8	0. 9	5.2	4.7	1. 3
HLF17 1 III 6-7	117 1.5	0. 1 3	0. 6 9	25 .3	28 .0	0.85	2.13	15 .0	7 7. 6	7. 4	2. 4	0. 0	7 2. 2	2. 5	9. 1	4. 5	65 .5	1. 2	6.8	5.7	1. 5
HLF17 1 III 11-12	117 6.5	0. 1 2	0. 6 6	- 25 .9	30 .4	0.89	0.57	11 .2	7 9. 4	9. 4	4. 9	6. 4	4 6. 0	4. 5	2 3. 0	1. 5	65 .6	0. 8	4.5	3.9	1. 2
HLF 17 1 III 16-17	118 1.5	0. 1 4	0. 8 4	- 27 .0	35 .3	0.90	0.80	13 .8	7 7. 8	8. 4	3. 9	3. 6	5 1. 3	2. 2	2 2. 4	1. 4	67 .3	0. 8	4.7	4.5	1. 2
HLF 17 1 III 18-19	118 3.5	0. 1 3	0. 6 9	27	37	0.89	0.31	17	7 5. 3	7.	5. 2	1.	3 9. 0	2 7. 2	9. 0	5. 4	66 .1	0. 8	4.8	4.3	- 1. 2
HLF17 1 III 27.28	119	0. 1	0. 7 8	26	30	0.88	0.81	13	7 8.	8.	- 9. 7	3.	5 2. 3	- 4. 1	1 7.	0.	66 1	1.	57	48	- 1. 1
HLF17 1 III 39-40	120 4.5	4 0. 1	0. 4	.9 - 28	.1 27 .0	0.88	0.43	.0 12 .2	7 6.	1 1.	4. 7	4 9. 3	1 8.	4 0.	3. 7	1. 9	.1 64 .2	1. 0	5.7	4.9	4 1. 3

Image: bold of the second se	T-test	s.d.	1	0	0	7	0.02	0.66	8	7	0	1	5	9	7	2	3	8	1	0.9	0.7	2
I 6 0. - - 7 1 4 1 -			0. 0	0. 1	1.	4.			1.	1.	1.	4.	9.	1 4.	9.	5.	1.	1.	0.			0.
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$		mea n	0. 1 2	0. 6 4	- 27 .0	34 .2	0.89	0.83	14 .1	7 7. 0	8. 9	6. 3	8. 3	4 0. 8	7. 4	1 6. 4	2. 4	65 .4	0. 9	5.2	4.6	1. 2
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	HLF17 1 III 95.5-96.5	126 1	0. 1 3	6 5	25 .9	34 .0	0.87	2.50	13 .2	7. 8	9. 0	4. 1	3. 8	4 9. 3	3. 5	8. 1	3. 5	66 .3	1. 0	6.2	5.3	1. 3
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	HLF17 1 III 88.5-89.5	125 4	1 2 0	6 4 0	27 .2	30 .5	0.89	1.37	12 .9	7. 8 7	9. 3	4. 4	3. 3	6. 5	4. 8	1. 8	2. 3	65 .7	1. 0	5.8	5.0	1. 3
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	HLF17 1 III 83.5-84	124 8.75	1 1 0.	5 6 0.	28 .6	45 .3	0.93	0.30	17 .7	3. 0 7	9. 3	7. 6 1	2. 2	6. 0 4	4. 4	8. 7 1	2. 0	63 .9	0. 7	3.8	3.6	0. 9
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	HLF 17 1 III 82.5-83	124 7.75	1 1 0.	5 7 0.	26 .9 -	29 .6	0.92	1.78	14 .6	6. 7 7	8. 7	5. 2 1	3. 9 2	0. 4 1	5. 3	2. 8 1	2. 7	64 .5	0. 8	4.2	3.9	1. 1
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	72.5-73.5	8	2 0.	6 0.	.4	.7	0.90	0.42	.6	8. 0 7	9. 3	4. 8	0. 7 1	6 2	8	2. 8 2	2. 5	.9	9	5.0	4.4	2
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	HLF17 1 III 60-61	122 5.5	1 2 0.	6 4 0.	26 .4 - 27	35 .8	0.88	0.83	14 .6	6. 8 7	8. 7	3. 8	1. 9 1	3. 7 4	5. 0	5. 1 1 2	2. 6	63 .5	1. 1	6.0	5.6	1. 4
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	HLF17 1 III 55-56	122 0.5	1 3 0.	6 8 0.	25 .3	39 .6	0.88	1.12	11 .3	9. 3 7	9. 4	5. 9	0. 4	7. 6 5	2. 3	8. 7 1	1. 8	67 .0	0. 9	5.2	4.9	1. 1
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	HLF17 1 III 48-49	121 3.5	1 0 0	4 6 0	28 .1	30 .2	0.92	0.36	13 .1	, 7. 8 7	9. 1	4. 2	5. 7	6. 6	9. 9	9. 3	1. 5	64 .5	0. 7	4.1	4.0	0. 9
1 6 .0 2 6 9 4 0. 0. - 7 1 4 1 HLF17 1 III 120 1 6 27 34 17 4. 8. 1. 4. 8. 1. 68 0. 1. 42.5-43.5 8 2 0 .1 .2 0.91 0.42 .3 2 5 1 9 6 7 5 3 .2 7 3.9 3.4 0 0. 0. - 7 1 4 9 7 1. 7 3.9 3.4 0 HLF17 1 III 120 1 6 27 36 .4 6 9. 4. 9. 7 1. 65 0. 1. 43.5-44 8.75 3 4 .3 .3 0.92 0.29 .1 3 6 3 2 9 0 8 3 .3 8 4.1 3.7 1	HLF17 1 III 44.5-45	120 9.75	0. 1 0	0. 5 1	28 .8	41 .8	0.93	0.23	14 .9	/ 5. 6 7	9. 5	2. 6	3 6. 3	2 5. 0 2	5. 9	1 4. 1	1. 1	60 .6	1. 0	6.2	5.0	1. 4
1 6 .0 2 6 9 4 0. 0. - 7 1 4 1 HLF17 1 III 120 1 6 27 34 17 4. 8. 1. 48 0. 1. 42.5-43.5 8 2 0 .1 .2 0.91 0.42 .3 2 5 1 9 6 7 5 3 .2 7 3.9 3.4 0	HLF17 1 III 43.5-44	120 8.75	0. 1 3	0. 6 4	27 .3	36 .3	0.92	0.29	14 .1	6. 3	9. 6	4. 3	9. 2	2 7. 9	1. 0	7. 8	1. 3	65 .3	0. 8	4.1	3.7	1. 1
	HLF17 1 III 42.5-43.5	120 8	1 0. 1 2	6 0. 6 0	.0 - 27 .1	34 .2	0.91	0.42	17 .3	2 7 4. 2	6 8. 5	1 1. 1	4. 9	9 4 1. 6	4 4. 7	1 8. 5	1. 3	68 .2	0. 7	3.9	3.4	1. 0

n.m= not measured

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			haetoceros RS	haetoceros dichaeta	orethron pennatum	ucampia antarctica	ragilariopsis curta	ragilariopsis cylindrus	ragilariopsis obliquecostat	hizosolenia spp.	tal		haetoceros RS	haetoceros dichaeta	orethron pennatum	ucampia antarctica	ragilariopsis curta	ragilariopsis cylindrus	ragilariopsis obliquecostat	
laminae	depth		0	0	U relative	ليا rebude	بة ce (cour	ts) %	F	R	to		0	0	U relative :	ية abudance (بة hiovolu	ne) %	<u>با</u>	
lammac	(cm)				Telative	abuuai	ice (cour	113), 70							Telative a	abuuance (biovoiui	nc), 70		
												average biovolume (µm ³) ^b	277	19227	70686	40457	642	231	4936	169
light					17.0															
HLF17 I III 14-15	1179.5	Light	3.6	0.0	17.3	0.0	56.7	2.9	9.2	0.3	90.0		0.1	0.0	89.8	0.0	2.7	0.0	3.3	4
HLF17 1 III 17-18	1182.5	Light	19.0	0.0	26.1	0.0	19.0	12.5	7.7	0.3	84.5		0.3	0.0	94.7	0.0	0.6	0.1	2.0	2
HLF 17 1 III 20-21	1185.5	Light	8.2	0.9	5.8	0.0	51.2	3.2	14.3	0.0	83.5		0.4	3.1	77.0	0.0	6.1	0.1	13.2	0
HLF17 1 III 33-34	1198.5	Light	5.3	0.0	11.3	0.0	45.2	10.7	12.9	0.3	85.6		0.2	0.0	84.4	0.0	3.1	0.3	6.7	5
HLF17 1 III 34.5-35	1200	Light	7.0	1.0	8.2	0.3	33.4	5.1	14.3	0.5	69.9		0.2	2.5	73.1	1.3	2.7	0.1	8.9	11
HLF 17 1 III 46-47	1211.5	Light	4.5	0.0	36.7	0.0	22.3	7.3	12.3	0.6	83.8		0.0	0.0	93.5	0.0	0.5	0.1	2.2	3
HLF 17 1 III 51-52	1216.5	Light	7.4	0.8	10.8	0.0	56.8	4.0	6.7	0.0	86.6		0.2	1.8	89.6	0.0	4.3	0.1	3.9	0
HLF17 1 III 57-58	1222.5	Light	9.5	0.0	20.3	0.0	36.1	6.3	11.0	0.3	83.6		0.2	0.0	91.6	0.0	1.5	0.1	3.5	3
HLF 17 1 III 63-64	1228.5	Light	8.3	4.3	5.5	0.2	40.1	6.0	12.1	0.0	76.3		0.4	14.5	68.7	1.1	4.5	0.2	10.5	0
HLF17 1 III 71-72	1236.5	Light	9.4	0.3	16.8	0.1	45.3	2.8	9.3	0.0	84.0		0.2	0.4	93.0	0.5	2.3	0.1	3.6	0
HLF17 1 III 73.5-74.5	1239	Light	17.4	0.0	9.0	0.2	25.0	3.3	18.2	0.0	73.1		0.6	0.0	84.2	0.9	2.1	0.1	12.0	0
HLF17 1 III 80.5-81.5	1246	Light	1.3	0.0	44.6	0.0	24.2	2.1	10.5	0.0	82.7		0.0	0.0	97.9	0.0	0.5	0.0	1.6	0
HLF17 1 III 84-5-85.5	1250	Light	12.5	2.5	14.0	0.0	31.6	6.4	11.4	0.0	78.3		0.3	4.3	88.4	0.0	1.8	0.1	5.0	0
HLF 17 1 III 86.5-	1252	Light	14.7	0.3	8.3	0.0	24.9	13.4	8.6	0.7	70.8		0.5	0.8	76.1	0.0	2.1	0.4	5.5	14
87.5 HLF17 1 III 91-92	1256.5	Light	10.5	1.9	18.6	0.0	35.8	1.9	9.6	0.3	78.6		0.2	2.4	89.0	0.0	1.6	0.0	3.2	3
		average	9.2	0.8	16.9	0.0	36.5	5.9	11.2	0.2			0.3	2.0	86.1	0.3	2.4	0.1	5.7	3
		s.d.	5.0	1.2	11.4	0.1	12.4	3.7	3.0	0.2			0.2	3.7	8.7	0.5	1.6	0.1	3.8	4
dark																				
HLF17 1 III 1-2	1166.5	Dark	3.0	7.7	0.8	0.0	48.4	3.4	19.9	0.0	83.2		0.2	43.8	17.6	0.0	9.2	0.2	29.0	0
HLF17 1 III 3-4	1168.5	Dark	8.0	2.6	1.0	0.0	41.6	3.9	18.4	0.4	75.9		0.7	15.9	21.9	0.0	8.6	0.3	29.4	23
HLF17 1 III 6-7	1171.5	Dark	2.4	1.8	0.0	0.0	72.2	2.5	9.1	0.0	87.9		0.5	27.1	0.0	0.0	36.6	0.5	35.4	0
HLF17 1 III 11-12	1176.5	Dark	4.9	0.0	6.4	0.0	46.0	4.5	23.0	0.0	84.7		0.2	0.0	75.5	0.0	5.0	0.2	19.1	0
HLF 17 1 III 16-17	1181.5	Dark	3.9	0.9	3.6	0.0	51.3	2.2	22.4	0.5	84.8		0.2	3.3	50.7	0.0	6.6	0.1	22.2	16
HLF 17 1 III 18-19	1183.5	Dark	5.2	1.6	1.3	0.0	39.0	27.2	9.0	0.2	83.6		0.6	13.2	38.0	0.0	10.4	2.6	18.6	16
HLF17 1 III 27-28	1192.5	Dark	9.7	2.1	3.4	0.0	52.3	4.1	17.6	0.0	89.2		0.7	10.0	59.5	0.0	8.3	0.2	21.4	0
HLF17 1 III 39-40	1204.5	Dark	4.7	0.5	9.3	0.1	18.9	40.4	3.7	0.3	77.9		0.2	1.4	86.4	0.7	1.6	1.2	2.4	6
HLF17 1 III 42.5-43.5	1208	Dark	11.1	0.2	4.9	0.3	41.6	4.7	18.5	0.2	81.5		0.6	0.7	67.5	2.1	5.2	0.2	17.8	5
HLF17 1 III 43.5-44	1208.8	Dark	4.3	0.3	19.2	0.1	27.9	1.0	17.8	0.3	70.9		0.1	0.3	89.5	0.3	1.2	0.0	5.8	2
HLF17 1 III 44.5-45	1209.8	Dark	2.6	0.0	36.3	0.6	25.0	5.9	14.1	0.0	84.5		0.0	0.0	95.8	1.0	0.6	0.1	2.6	0
HLF17 1 III 48-49	1213.5	Dark	4.2	1.7	15.7	1.7	26.6	9.9	19.3	0.3	79.3		0.1	2.3	80.9	4.9	1.2	0.2	6.9	3
HLF17 1 III 55-56	1220.5	Dark	5.9	1.4	0.4	0.0	57.6	2.3	18.7	0.5	86.9		0.6	9.8	11.2	0.0	13.2	0.2	32.8	32
HLF17 1 III 60-61	1225.5	Dark	3.8	5.0	1.9	0.0	53.7	5.0	15.1	0.0	84.6		0.3	28.1	39.4	0.0	10.1	0.3	21.8	0
HLF17 1 III 72.5-73.5	1238	Dark	4.8	2.2	10.7	1.1	41.6	5.8	12.8	0.3	79.2		0.1	4.4	76.7	4.6	2.7	0.1	6.4	4
HLF 17 1 III 82.5-83	1247.8	Dark	5.2	0.5	13.9	0.8	20.4	5.3	22.8	0.5	69.4		0.1	0.8	79.2	2.5	1.1	0.1	9.1	7
HLF17 1 III 83.5-84	1248.8	Dark	17.6	0.4	22.2	0.0	16.0	4.4	18.7	0.0	79.3		0.3	0.5	93.1	0.0	0.6	0.1	5.5	0
HLF17 1 III 88.5-89.5	1254	Dark	14.4	1.0	3.3	0.0	46.5	4.8	11.8	0.5	82.3		0.9	4.5	54.0	0.0	6.9	0.3	13.5	19
HLF17 1 III 95.5-96.5	1261	Dark	4.1	3.5	3.8	0.3	49.3	3.5	18.1	0.6	83.3		0.2	11.6	46.5	2.2	5.5	0.1	15.4	18
		average	6.3	1.8	8.3	0.3	40.8	7.4	16.4	0.2			0.4	9.3	57.0	1.0	7.1	0.4	16.6	8

^a Percent biovolume =100 * (% relative abundance * V avg) / Total biovolume (from Alley et al., 2018)

^b average biovolume from Alley et al., 2018