Observations on the relationship between the Antarctic coastal diatoms Thalassiosira antarctica Comber and Porosira glacialis (Grunow) Jorgensen and sea ice concentrations during the late Quaternary

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Abstract :

The available ecological and palaeoecological information for two sea ice-related marine diatoms (Bacillariophyceae), Thalassiosira antarctica Comber and Porosira glacialis (Grunow) Jorgensen, suggests that these two species have similar sea surface temperature (SST), sea surface salinity (SSS) and sea ice proximity preferences. From phytoplankton observations, both are described as summer or autumn bloom species, commonly found in low SST waters associated with sea ice, although rarely within the ice. Both species form resting spores (RS) as irradiance decreases, SST falls and SSS increases in response to freezing ice in autumn. Recent work analysing late Quaternary seasonally laminated diatom ooze from coastal Antarctic sites has revealed that sub-laminae dominated either by I antarctica RS, or by R glacialis RS, are nearly always deposited as the last sediment increment of the year, interpreted as representing autumn flux. In this study, we focus on sites from the East Antarctic margin and show that there is a spatial and temporal separation in whether T antarctica RS or P. glacialis RS form the autumnal sub-laminae. For instance, in deglacial sediments from the Mertz Ninnis Trough (George V Coast) P. glacialis RS form the sub-laminae whereas in similar age sediments from Iceberg Alley (Mac.Robertson Shelf) T antarctica RS dominate the autumn sub-lamina. In the Dumont d'Urville Trough (Adelie Land), mid-Holocene (Hypsithermal warm period) autumnal sub-laminae are dominated by T antarctica RS whereas late Holocene (Neoglacial cool period) sub-laminae are dominated by P. glacialis RS. These observations from late Quaternary seasonally laminated sediments would appear to indicate that P. glacialis prefers slightly cooler ocean-climate conditions than T antarctica. We test this relationship against two down-core Holocene quantitative diatom abundance records from Dumont d'Urville Trough and Svenner Channel (Princess Elizabeth Land) and compare the results with SST and sea ice concentration results of an Antarctic and Southern Ocean Holocene climate simulation that used a coupled atmosphere-sea ice-vegation model forced with orbital parameters and greenhouse gas concentrations. We find that abundance of P. glacialis RS is favoured by higher winter and spring sea ice concentrations and that a climatically-sensitive threshold exists between the abundance of P. glacialis RS and T antarctica RS in the sediments. An increase to >0.1 for the ratio of P. glacialis RS:T antarctica RS indicates a change to increased winter sea ice concentration (to >80% concentration), cooler spring seasons with increased sea ice, slightly warmer autumn seasons with less sea ice and a change from similar to 7.5 months annual sea ice cover at a site to much greater than 7.5 months. In the East Antarctic sediment record, an increase in the ratio from <0.1 to above 0.1 occurs at the transition from the warmer Hypsithermal climate into the cooler Neoglacial climate (similar to 4 cal kyr) indicating that the ratio between these two diatoms has the potential to be used as a semiquantitative climate proxy.

Keywords : East Antarctica, Quaternary, diatoms, sea ice

1. Introduction

73	Most of the ecological and palaeoecological information available for sea ice-related marine
74	diatoms Thalassiosira antarctica and Porosira glacialis suggests that these two species have
75	similar sea surface temperature (SST), sea surface salinity (SSS) and sea ice proximity preferences
76	and similar seasonal occurrences (summarised in Sections 1.1 and 1.2). For example, in Ross Sea
77	surface waters that emerge from beneath the ice shelf, T. antarctica cells are associated with a
78	diatom assemblage that also includes P. glacialis and P. pseudodenticulata (Cunningham and
79	Leventer, 1998). Along the Mac.Robertson Shelf in Iceberg Alley (Figure 1) deglacial seasonally
80	laminated sediments contain abundant P. glacialis resting spores (RS) that are found within T.
81	antarctica RS-dominated sub-laminae that were deposited during autumn (Stickley et al., 2005).
82	On the other hand, in Dumont d'Urville Trough, Adélie Land (Figure 1), sub-laminae dominated by
83	T. antarctica RS are found in mid-Holocene (relatively warm, Hypsithermal) laminated sediments
84	(Core MD03-2601, Denis et al., 2006), whereas sub-laminae dominated by P. glacialis RS are

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85 found in late Holocene (cool, Neoglacial) sediments (Core MD03-2597, Maddison, 2006). Further, deglacial/early Holocene laminated sediment from Mertz Ninnis Trough, George V Coast (Figure 86 87 1), also contain sub-laminae dominated by P. glacialis RS (Maddison et al., 2006). The T. 88 antarctica RS and P. glacialis RS sub-laminae from both sites, and from both time intervals, occur 89 at the top of late summer/autumn terrigenous-rich diatom laminae, which supports the contention 90 that these two diatoms have broadly similar ecological preferences (Stickley et al., 2005). 91 However, as pointed out by Denis et al. (2006), the more coastal position of the Mertz Ninnis 92 Trough and MD03-2597 core sites (and associated P. glacialis RS sub-laminae) compared to the core site MD03-2601 (and associated T. antarctica RS sub-laminae) suggests that P. glacialis 93 94 prefers cooler temperatures with higher sea ice concentrations than T. antarctica. In this study we 95 review the available ecological information for these two diatom species. We then consolidate the perceived relationship between the two taxa as sedimentary indicators of autumnal conditions using 96 97 laminated sediment sequences and published ecological information. Finally, we investigate down-98 core Holocene diatom assemblage records from two sites in coastal East Antarctica in order to establish whether the relationship developed from laminated sediment records and ecological 99 100 information can be used as a more quantitative proxy for changing environmental conditions during 101 the Late Quaternary.

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103 1.1. Ecology of Thalassiosira antarctica Comber 1896

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Thalassiosira antarctica is an Antarctic diatom species rarely found living within sea ice (Hasle and
Heimdal, 1968; Leventer and Dunbar, 1987; Fryxell and Kendrick, 1988; Zielinski and Gersonde,
107 1997), however, it is commonly described as a sea ice-associated diatom (i.e. living in waters with a
high concentration of sea ice) linked with low sea surface temperature (SST) and low sea surface
salinity (SSS) (Villareal and Fryxell, 1983). Blooms of *T. antarctica* are often recorded in waters
associated with newly forming sea ice, such as the formation of platelet ice next to the Ross Sea ice

111 shelf (Cunningham and Leventer, 1998) and both frazil ice and platelet ice in the Weddell Sea (Smetacek et al., 1992; Gleitz et al., 1998). In the Weddell Sea, it has also been recorded under 112 113 turbulent conditions (Gleitz et al., 1998) and from low salinity, nitrate-deplete, high pH crackpools 114 associated with summer melting ice (Gleitz et al., 1996). Although it is rare to find *T. antarctica* in sea ice samples due to its requirement for open water to bloom (Bárcena et al., 1998) and its 115 sensitivity to low light intensities (Doucette and Fryxell, 1985; Fryxell et al., 1987), it has been 116 observed in some spring sea ice samples which suggests over-wintering in sea ice or re-suspension 117 118 from the sediments (Villareal and Fryxell, 1983).

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120 Thalassiosira antarctica appears to be a summer and autumn bloom species. It has been observed as a common component of the early summer phytoplankton around the Antarctic Peninsula 121 (Sommer, 1991) and was recorded in summer sediment traps in the Ross Sea (Leventer pers, comm. 122 123 in Taylor and McMinn, 2002). It has been described as a major component of phytoplankton blooms in non-stratified or weakly stratified Antarctic surface waters (Cremer et al., 2003) (strong 124 water column stratification is usually associated with spring sea ice melt). T. antarctica has also 125 been associated with autumn bloom conditions in the Ross Sea with production of resting spores 126 related to the seasonally-late development of solid ice cover (Cunningham and Leventer, 1998). 127

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129 However, contrary to these observations of natural populations, laboratory culture experiments revealed that T. antarctica can grow well in reduced light conditions in temperatures as low as -4°C 130 131 (Bartsch, 1989; Aletsee and Jahnke, 1992), and that vegetative T. antarctica cells can survive in sea ice (Aletsee and Jahnke, 1992) and for prolonged periods of darkness (up to 214 days) without 132 133 forming resting spores (Peters and Thomas, 1996a). It is believed that the formation of 134 physiologically resting cells (identical in appearance to the vegetative cells) is induced by nutrient stress (Peters and Thomas, 1996b), or by low light intensities beneath the summer diatom bloom 135 136 and/or associated nutrient depletion (Taylor and McMinn, 2001). However, culture experiments

carried out at -1.5°C failed to induce resting spore formation by nitrogen depletion (Villareal and
Fryxell, 1983). Significant lipid accumulation occurs in the resting spores as they form (Doucette
and Fryxell, 1985) and this lipid synthesis has been attributed to a synergistic interaction between
reduced light levels, reduced SST and increased SSS in other Antarctic sea-ice related
phytoplankton species (Smith and Morris, 1980; Palmisano and Sullivan, 1982). In conclusion, the
exact trigger for resting spore formation in *T. antarctica* is not well understood.

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144 Observations of T. antarctica from the sediment record are almost exclusively the resting spore stage (Fryxell et al., 1981) (Figure 2). T. antarctica resting spores are heavily silicified and this has 145 146 been suggested as a mechanism to facilitate rapid sinking out of freezing surface waters (Doucette and Fryxell, 1985). The heavily silicified resting spores are more readily transported, hence, often 147 become concentrated in Antarctic near-coastal sediments (Gersonde and Wefer, 1987; Hemer and 148 149 Harris, 2003). From spatially extensive surface sediment records, *T. antarctica* resting spores are most abundant beneath regions where February SSTs of 0 to 0.5°C and where sea ice is present for 150 at least 6 months per year with winter sea ice concentrations >70% and unconsolidated summer sea 151 152 ice concentrations between 15-40% (Armand et al., 2005). Early Holocene sediments from Prydz 153 Bay are characterised by abundant *T. antarctica* resting spores (average of 31% of the assemblage) 154 and this has been related to seasonally warmer, more open marine conditions (Taylor and McMinn, 155 2002; Taylor and Leventer, 2003). In the Bransfield Strait, western Antarctic Peninsula, maximum 156 abundance of T. antarctica resting spores in Holocene sediments is related to cold climate episodes (Bárcena et al., 1998), persistent influence of cold Weddell Sea water (Gersonde and Wefer, 1987; 157 Leventer, 1991; Zielinski and Gersonde, 1997) and proximity to ice edge (Heroy et al., 2008). In 158 159 deglacial and Holocene seasonally laminated Antarctic coastal sediments, T. antarctica resting 160 spores commonly dominate the last recorded flux of the season, indicating their prior prevalence in mid- to late-summer blooms (as vegetative cells) and a relationship with falling temperatures (ice 161 162 formation) and falling light levels (Maddison et al., 2005; Stickley et al., 2005).

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164 1.2. Ecology of Porosira glacialis (Grunow) Jörgensen 1905

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166 Generally, less is known about the ecological preferences of *Porosira glacialis* than for T. antarctica. Porosira glacialis is a bipolar diatom species associated with cold coastal waters 167 adjacent to sea ice (Hasle, 1973; Taylor et al., 1997; Zielinski and Gersonde, 1997). Of the two 168 169 species of Antarctic Porosira, P. pseudodenticulata is commonly observed living in pack ice and 170 fast ice samples whereas P. glacialis is rarely recorded living in sea ice (Watanabe, 1988; Scott et al., 1994; Armand, 1997). It has been suggested that *P. glacialis* predominantly grows in the open 171 172 ocean beyond the sea ice edge (Zielinski and Gersonde, 1997), however, P. glacialis has been observed in waters with high concentrations of slush and wave-exposed shore ice although, again, 173 not living within the ice (Krebs et al., 1987). Similar to T. antarctica, in culture experiments, P. 174 175 pseudodenticulata survived prolonged periods of darkness (up to 272 days) by forming physiologically resting cells, not by forming resting spores (Peters and Thomas, 1996a). Resting 176 spore formation was not induced at -1.5°C (Villareal and Fryxell, 1983), and the exact mechanism 177 178 that promotes resting spore formation is not known. 179

P. glacialis is recorded in sediments predominantly as resting spores (Figure 2) and located 180 181 shoreward of the maximum winter sea ice extent (Armand, 1997). From the sediment record, P. glacialis appears to grow in summer waters with SSTs ranging from -2 to 1.5°C (slightly cooler 182 183 maximum temperature than T. antarctica) (Zielinski and Gersonde, 1997), and reaches maximum abundances with February SSTs of 0 to 0.5°C (Armand et al., 2005). From a spatially extensive 184 185 sediment surface data set, P. glacialis is most abundant beneath regions that experience at least 7.5 186 months per year sea ice cover (slightly longer than T. antarctica), with <30% summer sea ice concentration and highly compacted winter sea ice (65-85% concentration) (Armand et al., 2005). 187 188 In deglacial, seasonally laminated sediments from Iceberg Alley, Mac.Roberston Shelf, P. glacialis

RS are abundant in *Thalassiosira antarctica* RS sub-laminae that record the final flux event of the year, thus suggesting a similarity in their growth requirements (Stickley et al., 2005). In Holocene laminated sediments from the Mertz-Ninnis Trough, George V Coast, *P. glacialis* RS sub-laminae are interpreted as representing late summer/autumn deposition as sea ice concentration in the Mertz Glacier polynya increased (Maddison et al., 2006).

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- 1962.Materials and Methods
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198 2.1. Materials

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This study uses a suite of Late Quaternary sediment cores recovered from the East Antarctic 200 201 margin. Cores from Mertz Ninnis Trough (NBP0101 JPC10/KC10A), Iceberg Alley (NBP0101 JPC43B) and Svenner Channel (NBP0101 JPC24) were recovered during RVIB Nathaniel B. 202 Palmer cruise NBP0101 during 2001 (Table 1, Figure 1). Cores from the Dumont d'Urville Trough 203 204 (MD03-2597/MD03-2601) were recovered during the MD130-Images X-CADO cruise of RV Marion Dufresne II in 2003 (Table 1, Figure 1). All cores are, in part, seasonally laminated with 205 206 the remaining stratigraphy being either intermittently laminated or homogenous siliceous-mud ooze 207 (Crosta et al., 2005; Leventer et al., 2006; Maddison, 2006; Maddison et al., 2006). Intervals 208 selected for lamina-scale analyses and down-core diatom assemblage analysis in this study are 209 illustrated in Figure 1 and listed in Table 1.

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211 The early Holocene and Neoglacial (Figure 1) laminated sediment sequences from Mertz Ninnis

212 Trough were deposited approximately 30 km from the coast, beneath the site of a persistent winter

- 213 polynya (Maddison et al., 2006). The two cores from Dumont d'Urville Trough were recovered
- from biogenic drift deposits that accumulated in small depressions, approximately 30 km and 60 km

215 offshore, within the trough (Crosta et al., 2005). The Dumont d'Urville Trough region is an open coastline with a typical regime of advance and retreat of seasonal sea ice, with the episodic presence 216 217 of a polynya (Arrigo and Van Dijken, 2003). Today, the sites are covered by sea ice for 8-9 months 218 of the year (Schweitzer, 1995). During the deposition of the deglacial laminated sediments in 219 Iceberg Alley the long, across-shelf trough that comprises the basin was surrounded by a calving 220 bay re-entrant (Leventer et al., 2006). This meant that even though the site is approximately 70 km 221 offshore today, it would have been subject to a seasonal sea ice retreat and advance regime typical 222 of more coastal sites. Svenner Channel is a coast-parallel trough on the eastern margin of Prydz Bay. The core site is approximately 60 km from the coast and is subject to a typical seasonal sea 223 224 ice advance and retreat regime that results in a yearly sea ice presence of 10 months.

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226 2.2. Lamina analyses: BSEI and LM sediment fabric analysis and quantitative diatom
227 assemblage analysis

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Laminated sediments (Figure 1) were embedded in resin and polished thin sections were produced 229 230 for microscope analysis following the methods outlined in Maddison et al. (2006, NBP0101 JPC10/KC10A, MD03-2597), Denis et al. (2006, MD03-2601) and Stickley et al. (2005, NBP0101 231 232 JPC43B). Laminated sequences (polished thin sections) were logged for sediment fabric and 233 diatom assemblages at the 100 micron-scale using either scanning electron microscopy 234 backscattered electron imagery (BSEI, Pike and Kemp, 1996; Pearce et al., 1998) or optical light 235 microscopy (LM, Denis et al., 2006) (Table 2). For core sites NBP0101 JPC10/KC10A and MD03-2597 additional quantitative diatom assemblage analyses were carried on specific lamination types 236 237 following the method outlined in Maddison et al. (2006). These additional analyses facilitated 238 quantitative comparison of the relative abundance of T. antarctica RS and P. glacialis RS in the 239 laminations, as opposed to the more qualitative techniques involved in thin section analysis using 240 BSEI and LM. Data are presented as absolute abundance (valves per gramme), relative abundance

241 and Hyalochaete Chaetoceros resting spore-free (CRS-free) relative abundance (Table 3). CRS-

242 free abundance is commonly used in Antarctic diatom assemblage analysis when CRS

243 overwhelmingly dominate every assemblage (Leventer et al., 1996; Allen et al., 2005). The CRS-

244 free abundance allows the ecological information within the minor species assemblage to be

critically examined.

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- 247 2.3. Down-core quantitative diatom assemblage analysis
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Down-core quantitative diatom assemblage analysis was carried out on MD03-2601 (Dumont 249 250 d'Urville Trough) and NBP0101 JPC24 (Svenner Channel) following the methods outlined in 251 Crosta et al. (2007). These records allow the relationship between T. antarctica RS and P. glacialis RS, deduced from lamination-scale analyses of mainly deglacial sediments, to be investigated over 252 253 the duration of the Holocene. Porosira glacialis RS and P. pseudodenticulata RS are grouped together, however, P. glacialis RS overwhelmingly dominates the group. P. pseudodenticulata has 254 no distinct central annulus, a slightly larger diameter and fewer areolae in 10 µm (~60-80 µm; 10-255 256 12) than *P. glacilalis* (30-40 µm; 25-26) which often has a distinct central annulus. The two forms of *T. antarctica* RS are grouped (T1 and T2, Buffen et al., 2007), however, the cool *T. antarctica* 257 258 RS form (T1) overwhelmingly dominates in both Dumont d'Urville Trough and Svenner Channel 259 sediments.

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261 2.4. Modelled Holocene surface temperatures and sea ice concentrations

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The diatom abundance and sediment fabric data and interpretations will be compared with the results of a 9000-yr Holocene climate simulation using the ECBilt-CLIO-VECODE coupled atmosphere-sea ice-ocean-vegetation model forced with annually varying orbital parameters (monthly insolation at 60°S) and atmospheric greenhouse gas (carbon dioxide and methane)

267 concentrations derived from ice cores (Opsteegh et al., 1998; Goosse and Fichefet, 1999; Brovkin et 268 al., 2002; Renssen et al., 2005). In the model, the vernal equinox is fixed at day 81 and the year is 269 divided into 12 months of 30 days. All other forcings (e.g. solar constant, other greenhouse gases, 270 ice sheet configuration) are set at AD 1750 values, hence, changes in ice sheet volume are not taken 271 into account in the simulation used here. As well as comparing with the published East Antarctic modelled summer temperature record of Renssen et al. (2005), we will also compare our results to a 272 273 sub-set of the whole model output for the Adélie Land sector that encompasses the Dumont 274 d'Urville Trough core sites (south of 60°S, 140°E-150°E). Two 1000-year time periods from the Adélie Land mean monthly surface temperature and sea ice concentrations were selected to provide 275 276 a measure of typical mean monthly conditions; the interval 4999-4000 yr BP was selected as typical 277 of the Hypsithermal and 1999-1000 yr BP selected as representative of the Neoglacial. These two time periods were compared to investigate statistically significant differences in modelled monthly 278 279 parameters between the Hypsithermal and Neoglacial. Significant differences in the means were found using a standard z-test (with 2-tailed critical values) for assessing differences in the means 280 281 between large sample sets (Table 4).

- 282
- 283 **3. Results**
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- 285 3.1 Laminated sediments BSEI, LM and quantitative diatom assemblage analysis
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Data for the lamina-scale diatom assemblage analyses, using BSEI, LM and quantitative abundance,
come from both published sources and from new analyses. BSEI lamina characterisations for
sediments from the Mertz-Ninnis Trough (NBP0101 JPC10/KC10A) are summarised from
Maddison (2006) and Maddison et al. (2006). Similar data for the Dumont d'Urville Trough
(MD03-2597) are summarised from Maddison (2006) and LM lamina data (MD03-2601) are

summarised from Denis et al. (2006). Lamina-scale diatom assemblage data for sediments from

293 Iceberg Alley (NBP0101 JPC43B) are a combination of both published (Stickley et al., 2005,

interval 21.84-23.96 m) and previously unpublished data (Stickley and Pike: interval 19.30-21.84

m). Tables 2 and 3 provide a summary of these data containing only the lamina or sub-lamina datarelevant to this study.

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298 3.1.1. Thalassiosira antarctica RS sub-laminae

Analysis of lamina occurrence data from the spectacularly laminated deglacial sediments of the East 299 300 Antarctic margin reveals that when sub-laminae characterised by T. antarctica RS or P. glacialis RS occur, usually: (1) either one or the other species is dominant; and (2) they are found in a similar 301 302 position within the seasonal succession of diatom assemblages and lamina types, i.e. at the end of the season of diatom production and subsequent flux to the sediment, late summer or autumn 303 (Stickley et al., 2005; Denis et al., 2006; Maddison, 2006). In deglacial Iceberg Alley sediment, 43 304 305 of 68 T. antarctica RS sub-laminae occur in this position above a summer lamina (Table 2) and, consequently, directly below the following spring diatom ooze lamina from which they are 306 separated by a sharp contact delineating the winter hiatus. Further, when a summer lamination is 307 308 not present, 25 of 68 occurrences of *T. antarctica* RS sub-laminae follow a transitional lamination 309 (Table 2), defined as having some characteristics of both spring diatom ooze laminae and summer terrigenous-rich laminae (Stickley et al., 2005). These transitional laminae nearly always grade 310 311 upwards from a spring diatom ooze lamina and usually grade into a summer terrigenous-rich 312 lamina, however, sometimes these transitional laminae are abruptly overlain by the following spring 313 diatom ooze laminae. Usually, it is at the top of these latter transitional laminae that T. antarctica 314 RS sub-laminae occur. When T. antarctica RS sub-laminae occur within the annual sediment 315 increment, the mean annual sediment thickness is 2% greater than if sub-laminae do not occur. 316 Further, if a summer lamination is followed by a T. antarctica RS sub-lamina, the summer lamina is, on average, 7% thicker than one that is not associated with a sub-lamina (Table 2). Qualitative 317

- analysis of *T. antarctica* RS sub-laminae from Iceberg Alley also reveals that *P. glacialis* RS are
- 319 more abundant in the sub-laminae than in the summer, terrigenous-rich laminae.
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LM analysis of lamina sequences from the Holocene sediments of the less coastal MD03-2601 core 321 322 from Dumont d'Urville Trough shows the presence of sub-laminae of T. antarctica RS associated with late summer/autumn in Hypsithermal-age sediments and the absence of the sub-laminae in 323 324 Neoglacial-age sediments (Table 2, Denis et al., 2006). During the Hypsithermal period, annual 325 sediment thickness was 46 mm compared with 21 mm during the Neoglacial, and mean thickness of T. antarctica RS sub-laminae during the Hypsithermal was 0.371 mm (Table 2, Denis et al., 2006). 326 327 In contrast to the other, more coastal core site from Dumont d'Urville Trough (MD03-2597) these sections from MD03-2601 do not contain any *P. glacialis* RS sub-laminae during the Neoglacial. 328

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- 330 3.1.2 Porosira glacialis RS sub-laminae

Nineteen out of twenty two occurrences of *P. glacialis* RS sub-laminae in deglacial Mertz-Ninnis 331 Trough sediment appear in a late summer/autumn position in the seasonal lamina succession (Table 332 333 2). Quantitative diatom abundance analysis of Mertz-Ninnis Trough laminations reveals that although, in absolute terms, *P. glacialis* RS have a similar abundance in summer mixed diatom 334 laminae as they do in P. glacialis RS sub-laminae (Table 3, Maddison et al., 2006), P. glacialis RS 335 comprise, on average, 11.35% of the CRS-free assemblage in P. glacialis RS sub-laminae and only 336 337 2.29% in the summer mixed assemblage laminae (Table 3, Maddison et al., 2006). At this site, T. 338 antarctica RS are absent from both the summer mixed diatom laminae and the P. glacialis RS sub-339 laminae (Table 3, Maddison et al., 2006). When a P. glacialis sub-lamina occurs within the annual 340 sediment increment, the mean annual sediment thickness is 18% greater than if a sub-lamina does 341 not occur (Table 2). BSEI analysis of a 0.28 m-long (17 year) sequence of early Neoglacial-age laminations from this site (NBP0101 KC10A; Table 1 and Figure 1) does not reveal any T. 342 343 antarctica RS or P. glacialis RS sub-laminae (Maddison, 2006).

345	In Neoglacial laminated sediments from the more coastal Dumont d'Urville Trough core (MD03-
346	2597), in 23 out of 26 years when P. glacialis RS sub-laminae occur, the sub-laminae appear in a
347	late summer/autumn position (Table 2, Maddison, 2006), separated with a sharp contact from the
348	overlying spring lamination. Quantitative diatom abundance analysis reveals that, in absolute
349	terms, P. glacialis RS are more than 2.5 times as abundant in P. glacialis RS sub-laminae than they
350	are in terrigenous-rich, summer mixed assemblage laminae. Further, T. antarctica RS are more
351	than 2.5 times as abundant in <i>P. glacialis</i> RS sub-laminae than in terrigenous-rich, summer mixed
352	assemblage laminae (Table 3, Maddison, 2006). When P. glacialis RS sub-laminae occur within
353	the annual sediment increment, the mean annual sediment thickness is 12% greater than if sub-
354	laminae do not occur (Table 2).
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356	3.2. Holocene records from Dumont d'Urville Trough and Svenner Channel
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358	Holocene relative abundance of <i>T. antarctica</i> RS and <i>P. glacialis</i> RS from Dumont d'Urville
359	Trough (MD03-2601) and Svenner Channel (NBP0101 JPC24) (Figure 1) are shown in Figure 3. In
360	general, at both sites, relative abundances of <i>T. antarctica</i> RS are higher than <i>P. glacialis</i> RS. In
361	the Dumont d'Urville Trough, the relative abundance of <i>T. antarctica</i> RS steadily rises to a peak at
362	~8.9 cal kyr BP (Figure 3B). Abundance then generally stays above 10% until ~3.4 cal kyr BP
363	when it falls below 10% until ~2 cal kyr BP. Abundance then becomes very low (<5%) between 2
364	cal kyr BP until ~1.1 cal kyr BP, after which it recovers to about 10% at the top of the core.
365	Relative abundance of <i>P. glacialis</i> RS displays an approximately inverse pattern. Abundance
366	remains above 2% from ~10.8 cal kyr BP until ~8.7 cal kyr BP (Figure 3C), with a slight dip in
367	abundance around 9.8 cal kyr BP. At 8.7 cal kyr BP, abundance falls abruptly below 2% until ~3.4
368	cal kyr BP, when it abruptly rises to above 2% until ~2.8 cal kyr BP when it falls below, and stays

below, 2% until ~2 cal kyr BP. After 2.0 cal kyr BP, abundance rises to above 2% and then steadily
falls towards the top of the core (~1 cal kyr BP).

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In Svenner Channel, T. antarctica RS relative abundance rises from 11.2 cal kyr BP to a peak of 372 373 >20% around 9.8 cal kyr BP (Figure 3E). Abundance dips below 20% around 9 cal kyr BP, rises to above 20% at ~8.7 cal kyr BP and then increases steadily to a peak at ~4.4 cal kyr BP. Relative 374 abundance begins to decline and falls below 20% at ~3.4 cal kyr BP; it remains below 20% until the 375 376 top of the core (~0.67 cal kyr BP). The relative abundance of *P. glacialis* RS remains below ~2% from 11.2 cal kyr BP until ~8.7 cal kyr BP when it begins to steadily rise (Figure 3F). At ~4.3 cal 377 378 kyr BP, abundance rises above 2% and stays high until ~1 cal kyr BP when it falls below 2% at the 379 top of the core.

380

The relationship between T. antarctica RS and P. glacialis RS can be further investigated by 381 looking at the ratio between the two diatoms, and also the statistical correlation between the two 382 species curves. The ratio between P. glacialis RS and T. antarctica RS (i.e. relative abundance of 383 384 P. glacialis RS/relative abundance of T. antarctica RS) in Dumont d'Urville Trough is approaching 0.5 from the base of the core (~10.8 cal kyr BP) until ~10 cal kyr BP where it abruptly falls to very 385 386 low values (Figure 3D). The ratio remains very low until ~3.4 cal kyr BP when it abruptly rises to 387 >0.1 up to the top of the core (~1 cal kyr BP). Between ~2.8 cal kyr BP and ~2 cal kyr BP, the ratio falls, however, generally remains above 0.1. In Svenner Channel, the ratio between P. glacialis RS 388 389 and *T. antarctica* RS remains generally <0.1 from ~11.2 cal kyr BP until ~4.3 cal kyr BP when it 390 begins to rise (Figure 3G). The ratio reaches peak values (~ 0.5) at ~ 2.8 cal kyr BP and generally 391 remains above 0.2 up to the top of the core (~0.7 cal kyr BP). The Pearson correlation coefficient 392 (r) between the relative abundance of T. antarctica RS and P. glacialis RS is -0.19 for the Dumont d'Urville record (n=528). This correlation coefficient appears weak, but a test of significance of r 393 394 demonstrates that threshold for significant values of r for sample sizes of n=528, at the 99%

significance level, is 0.14 (Lowry, 2008). There is no significant correlation between the relative
abundances of *T. antarctica* RS and *P. glacialis* RS in the Svenner Channel record. The correlation
coefficient is negative, but not significant (r=-0.0004; n=348). However, the significant negative
correlation between *T. antarctica* RS and *P. glacialis* RS for Dumont d'Urville Trough supports the
qualitative observations of an inverse relationship between the two taxa.

400

The diatom relative abundance curves are also compared with the modelled mean October to April 401 402 (i.e. austral summer) surface temperature record for the Holocene of East Antarctica (Renssen et al., 2005) (Figure 3A). The modelled surface temperature record reveals temperatures warming from 403 404 ~0.4°C warmer than pre-industrial levels at 9.0 kyr BP, to temperatures ~1.0°C warmer than preindustrial levels at ~4.3 kyr BP. Modelled temperatures then decline steadily until the end of the 405 record, approximately 300 yr BP. The Pearson correlation between the relative abundance of T. 406 407 antarctica RS and modelled mean summer temperature in Dumont d'Urville Trough and Svenner Channel is 0.44 and 0.64, respectively. The Pearson correlation between the relative abundance of 408 P. glacialis RS and modelled mean summer temperature in Dumont d'Urville Trough and Svenner 409 410 Channel is -0.48 and -0.11, respectively. The Pearson correlation between the modelled mean summer temperature and the ratio P. glacialis RS:T. antarctica RS is -0.55 for Dumont d'Urville 411 412 Trough and -0.40 for Svenner Channel. All but one of these correlation coefficient values are above 413 the threshold values of the Pearson's test of significance (r=-0.11 falls below the threshold). This highlights a positive correlation between the relative abundance of T. antarctica RS and modelled 414 415 mean summer temperature; a negative correlation between relative abundance of P. glacialis RS and modelled mean summer temperature; and a more robust negative correlation between the ratio 416 417 P. glacialis RS:T. antarctica RS and modelled mean summer temperature.

418

419 **4.** Interpretation and Discussion

421 4.1. Mechanism for formation of T. antarctica and P. glacialis resting spore sub-laminae

422

423 When sub-laminae characterised by T. antarctica RS or P. glacialis RS occur in the laminated 424 sediments from the East Antarctic margin, three features are consistent: (1) one or the other species is dominant; (2) the sub-laminae predominantly occur in the same position within the seasonal 425 426 sedimentary sequence – late summer to autumn; and (3) overall annual sediment thickness is greater 427 when sub-laminae are present (in deglacial Iceberg Alley sediments, the sub-laminae are also 428 associated with summer laminations that are thicker than average indicating increased summer production and flux to the sediments). Hence (2) and (3) can be interpreted together as indicating 429 430 that the sub-laminae are associated with years of higher diatom flux to the sediment. The occurrence of *T. antarctica* RS and *P. glacialis* RS sub-laminae (and thicker summer laminations) 431 432 could indicate either a longer duration summer season of diatom productivity, promoting greater 433 overall flux to the sediment, or enhanced summer conditions that favour higher T. antarctica and P. glacialis vegetative production, hence production and flux of the resting spores to the sediment in 434 435 autumn.

436

Thalassiosira antarctica RS and P. glacialis RS sub-lamina production is likely to be related to a 437 438 slow break up of sea ice over a sustained period in spring, i.e. a slow increase of temperatures over 439 a few weeks so that when light levels increase sufficiently to promote vegetative cell growth there is 440 a relatively high sea ice concentration that significantly effects surface water properties. Porosira glacialis and T. antarctica have both been found associated with spring sea ice-rich regions 441 442 (Villareal and Fryxell, 1983; Krebs et al., 1987). Relatively high spring sea ice concentrations 443 would lead to summer production favouring sea ice-associated diatoms such as T. antarctica and P. glacialis (standing stocks enhanced by 'icy' spring) and both species are known to be abundant in 444 Antarctic summer waters (Sommer, 1991; Zielinski and Gersonde, 1997; Cremer et al., 2003). 445 446 Finally, sea ice formation in the autumn, accompanied by lowering SSTs, increased salinity and

447 lowering light levels (Doucette and Fryxell, 1985; Fryxell et al., 1987; Cunningham and Leventer, 1998), would induce resting spore formation from a vegetative population that was sufficiently 448 449 large to produce a sub-lamina in the sediments (Stickley et al., 2005; Denis et al., 2006; Maddison 450 et al., 2006). Years without sub-laminae suggest more rapid melting and break-up of sea ice in 451 spring favouring more marginal ice zone/open water species and reduced growth of T. antarctica 452 and P. glacialis in the subsequent summer. Lower summer vegetative populations of T. antarctica 453 and P. glacialis would not lead to the production of sufficient numbers of resting spores to produce 454 a sub-lamina in autumn.

455

456 Key to confirming this sea ice-based model of sub-lamina formation are the two cores from Dumont d'Urville Trough which are geographically close to each other and would have experienced similar 457 458 changes in external forcing (i.e. by insolation and greenhouse gases). At MD03-2601, during the 459 Hypsithermal, T. antarctica RS sub-laminae were preserved (core MD03-2597 did not recover Hypsithermal-age sediments). In the Neoglacial, no sub-laminae of either species were preserved at 460 site MD03-2601, although higher relative abundances of *P. glacialis* RS are observed (Figure 3C), 461 whereas P. glacialis RS sub-laminae were preserved at MD03-2597 (Table 2). In the modern day, 462 site MD03-2597 is covered by less than 65% sea ice concentration for, on average, 111 days of the 463 464 year, whereas site MD03-2601 is subject to less than 65% sea ice concentration for only 94 days of the year (difference between means is significant at >95%; mean values extracted from satellite-465 466 derived daily sea ice concentrations for 26 years between 1979 and 2006, National Snow and Ice Data Centre, http://nsidc.org/data/nsidc-0079.html). Site MD03-2597 becomes 'ice-free' 467 468 (consistently less than 65% sea ice concentration), on average, 19 days earlier in the spring than site 469 MD03-2061 because the ice-edge not only retreats parallel with the coastline, but also retreats in a 470 westward direction along the coast towards the Dumont d'Urville Trough. This is due to the areal 471 expansion of the coastal Mertz Glacier Polynya, which merges with the open ocean in summer. So, 472 not only are there less ice-free days at MD03-2601, the sea ice retreats later in the spring. The

473 combined evidence of higher relative abundance of P. glacialis RS (MD03-2601) and the presence of P. glacialis RS sub-laminae (MD03-2597) in Neoglacial sediments, along with the cessation of T. 474 475 antarctica RS sub-laminae formation (MD03-2601), suggests that increasingly higher winter and 476 spring sea ice concentrations occurred in the Dumont d'Urville Trough region in the Neoglacial, 477 relative to the Hypsithermal period. Perhaps the environment became either too icy (given modern 478 seasonal sea ice differences detailed above) or, more likely, had too short an ice-free summer season at site MD03-2601 during the Neoglacial for even P. glacialis to be able to form large 479 enough populations to promote significant flux of resting spores to the sea floor and subsequent 480 481 sub-laminae formation.

482

483 4.2. Environmental controls on the P. glacialis RS:T. antarctica RS ratio in Holocene sediments

484

485 From the lamination data, T. antarctica RS and P. glacialis RS sub-laminae are both interpreted as representing years with relatively high spring sea ice concentrations, however, what threshold under 486 these conditions would govern changes in the relative abundance of the two species seen throughout 487 488 the Holocene (Figure 3), given that they appear to have similar ecologies? Here we combine our 489 sea ice-based mechanism of sub-lamina formation with published ecological preferences and 490 modelled Holocene surface temperature and sea ice concentrations from the Adélie Land sector 491 (Table 4, Figure 4) to present a model of the environmental conditions that alter the relative abundance of T. antarctica RS and P. glacialis RS in the sediments. 492

493

P. glacialis appears to prefer somewhat higher winter and spring sea ice concentrations than *T. antarctica.* Maximum abundances of *P. glacialis* RS in the sediment are found beneath regions
subject to >7.5 months per year sea ice cover (*T. antarctica* prefers >6 months), <30% summer sea
ice concentration (*T. antarctica* similar) and highly compacted winter sea ice (up to 85%)

498 concentration; *T. antarctica* prefers slightly less, but still >70%) (Armand et al., 2005). These

499 ecological observations are supported by our Holocene records and comparisons with the modelled
500 summer surface temperatures from East Antarctica (Renssen et al., 2005). Increased abundance of
501 *T. antarctica* RS in the sediments from Dumont d'Urville Trough and Svenner Channel is positively
502 correlated with modelled summer surface temperatures whereas increased abundance of *P. glacialis*503 RS is negatively correlated with modelled summer surface temperatures (Figure 3).

504

The modelled sea ice concentration and surface temperature records for the Adélie Land sector 505 suggest that the cool Neoglacial period had higher winter and early spring sea ice concentrations 506 than the mid Holocene Hypsithermal (Table 4, Figure 4), with significantly lower late spring and 507 508 early summer temperatures. In autumn, the Neoglacial had lower sea ice concentrations than the 509 Hypsithermal with slightly higher temperatures. This model output supports the interpretation from ecological and seasonal sedimentological data that P. glacialis prefers slightly cooler and 'icier' 510 winter and spring conditions than T. antarctica, which is why its abundance was enhanced during 511 the Neoglacial off both Adélie Land and Princess Elizabeth Land. A subsequent long 512 summer/autumn season, as suggested by the model data, would favour the build-up of larger 513 514 populations of *P. glacialis* and concomitantly reduced populations of *T. antarctica*. Our interpretations are in agreement with those of Crosta et al. (2008) who state that seasonal sea ice 515 516 distribution during the Holocene of East Antarctica is more complicated than a simple pattern of 517 less sea ice during the warmer Hypsithermal, and more during the cooler Neoglacial.

518

519 4.3. P. glacialis RS:T. antarctica RS – a new sediment proxy for winter and spring sea ice
520 concentrations

521

522 The results from Dumont d'Urville Trough (MD03-2601) and Svenner Channel sediments

523 presented here indicate that 0.1 appears to be a significant threshold value for the ratio *P. glacialis*

524 RS:*T. antarctica* RS. An increase above 0.1 accompanies an important environmental change to

525 increased winter sea ice concentration, cooler spring seasons with increased sea ice, and slightly 526 warmer autumn seasons with less sea ice. In the Holocene sediments of coastal East Antarctica, the 527 ratio changes from <0.1 to >0.1 between 4.3-3.4 cal kyr BP (i.e the transition from warmer 528 Hypsithermal conditions to cooler, Neoglacial conditions, Masson et al., 2000; Crosta et al., 2007). However, it is possible to use the ecological information in an attempt to be more quantitative than 529 530 this. Using the ecological preferences of P. glacialis and T. antarctica and it's distribution in 531 modern core top samples (Armand et al., 2005), an increase in the sediment ratio above 0.1 (i.e. 532 enhanced P. glacialis RS and/or relatively reduced T. antarctica RS abundance) appears to indicate a change from ~7.5 months of annual sea ice cover (favouring *T. antarctica* production), to much 533 534 greater than 7.5 months of annual sea ice cover (enhancing P. glacialis production), and an increase from ~70% winter sea ice concentration (favouring subsequent T. antarctica production) to highly 535 536 compacted winter sea ice above 80% concentration (enhancing P. glacialis production). Although 537 these may not seem like large changes, they appear to be key manifestations of the changes between warmer and cooler Holocene climate states as reflected in the diatom fossil record, and are in line 538 with the changes suggested by previous diatom evidence for the Hypsithermal-Neoglacial transition 539 540 (Crosta et al., 2008).

541

Attempts to apply the *P. glacialis* RS:*T. antarctica* RS proxy to west Antarctic Peninsula (WAP) 542 543 diatom records from Palmer Deep (Taylor and Sjunneskog, 2002) and Bransfield Strait (Heroy et 544 al., 2008) have met with limited success because the relative warmth of the WAP compared to the 545 East Antarctica margin limits the abundance of *P. glacialis* RS in the sediments to be almost always less than 10% of T. antarctica RS. However, a preliminary analysis of Holocene diatom abundance 546 547 records from the cooler NW Weddell Sea has proved promising in that stratigraphic intervals when 548 P. glacialis RS:T. antarctica RS was >0.1 (interpreted as representing cool, sea ice-rich winter and 549 spring conditions) are also intervals of high abundance of other sea ice-related diatoms such as 550 Fragilariopsis curta, F. cylindrus and Thalassiosira tumida (Anna Hey, personal communication,

551 2008). This demonstrates the potential wider applicability of the *P. glacialis* RS:*T. antarctica* RS
552 ratio around the Antarctic coast.

553

554 **5.** Conclusions

555

This synthesis of information on the abundance of Thalassiosira antarctica and Porosira glacialis 556 557 has shown that these two diatoms have similar ecological preferences. The combination of modern 558 phytoplankton analyses, laboratory experiments and late Quaternary laminated sediment records reveals that high fluxes of T. antarctica RS and P. glacialis RS to the sediment are associated with 559 560 prior high winter and spring sea ice concentrations that promote the build-up of large vegetative cell populations. In late autumn, abrupt sea ice advance, accompanied by decreasing temperatures and 561 562 irradiance and increased salinities promotes resting spore formation from high summer/autumn 563 standing stocks. The investigation of two core records from the Dumont d'Urville Trough and one from Svenner Channel showed that despite their similar ecological preferences, there were 564 interesting differences in the Holocene occurrence of the two taxa. Relative abundance of T. 565 antarctica RS peaked during the warm mid-Holocene Hypsithermal period and declined into the 566 cooler Neoglacial, whereas relative abundance of P. glacialis RS peaked during the late Holocene, 567 568 cool Neoglacial period. Increased P. glacialis RS abundance appeared to be linked with higher winter and spring sea ice concentrations than T. antarctica RS. These conclusions are corroborated 569 by a comparison with modelled Holocene sea ice concentrations and surface temperatures for the 570 571 Adélie Land sector. The model output reveals higher winter and early spring sea ice concentrations, 572 and lower spring and early summer temperatures during the Neoglacial relative to the Hypsithermal 573 (which would promote increased populations of *P. glacialis*). Investigation of the ratio of relative 574 abundance of the two diatoms, P. glacialis RS:T. antarctica RS, revealed that a threshold of 0.1 was important. The ratio abruptly increased from <0.1 to >0.1 at the Hypsithermal to Neoglacial 575 576 transition. An increase in the ratio P. glacialis RS:T. antarctica RS to greater than 0.1 appears to

indicate a change from ~7.5 months of annual sea ice cover to much greater than 7.5 months of
cover, and an increase of winter sea ice concentrations from 70% to >80%. An attempt to apply the *P. glacialis* RS:*T. antarctica* RS ratio to other core sites from West Antarctica, in particular the NW
Weddell Sea, shows that the relationship between these two diatom species may be able to reveal
information about past winter and spring sea ice concentrations around the Antarctic margin.

582

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584

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762 **Table and Figure Captions**

763

- 764 Table 1: East Antarctica margin sediment core information, including details of location, length,
- 765 distance from coast, core intervals analysed and age of intervals analysed.

766

- 767 Table 2: Summary of lamina and sub-lamina thickness measurements from BSEI and LM of East
- 768 Antarctic laminated sediment sequences.

769

770 Table 3: Quantitative abundance of *Porosira glacialis* RS and *Thalassiosira antarctica* RS from

771 selected East Antarctic sediment laminae and sub-laminae.

772

Table 4: Summary of Adélie Land sector modelled Holocene surface temperature and sea ice

concentrations for typical Hypsithermal (4000-4999 yr BP) and typical Neoglacial (1000-1999 yr

BP) periods (extracted from Renssen et al., 2005). Surface temperatures and sea ice concentrations

are given as anomalies from the pre-industrial mean (1000-250 yr BP).

777

Figure 1: Locations of the East Antarctic margin sediment cores used in this investigation. AL =
Adélie Land and PEL = Princess Elizabeth Land. The intervals sampled from each of the cores are
shown by the black (BSEI and LM sediment fabric analyses) and grey (Holocene down-core

quantitative diatom abundance analyses). The presence of *Thalassiosira antarctica* RS sub-laminae

is indicated by white and *Porosira glacialis* RS sub-laminae by a vertical line.

783

Figure 2: Light microscope (LM) and scanning electron microscope (SEM) images of

785 Thalassiosira antarctica and Porosira glacialis resting spores. SEM micrographs taken of Au-Pd

- coated, lamina parallel sediment fracture surfaces. A-B. LM micrographs of *T. antarctica* resting
- spores (NBP01-01 JPC10, 19.536 cm). C-D. Secondary electron SEM micrographs of *T*.

788	antarctica resting spores (MD03-2597, 5307.5 cm, scale bars $G = 5$ microns, $H = 10$ microns). E-
789	F. LM micrographs of <i>P. glacialis</i> resting spores (NBP01-01 JPC10, 19.536 cm). G-H. Secondary
790	electron SEM micrographs of P. glacialis resting spores (NBP01-01 JPC10, 19.463 cm; scale bars
791	C = 10 microns, $D = 50$ microns).
792	Q I
793	Figure 3: Holocene Thalassiosira antarctica and Porosira glacialis resting spore abundance from
794	Dumont d'Urville Trough (MD03-2601) and Svenner Channel (NBP01-01 JPC24). A. Modelled
795	mean summer (October to April) surface temperature (relative to the pre-industrial mean (1000-250
796	yr BP)) from Renssen et al. (2005). B-D. Dumont d'Urville Trough records. B. Relative
797	abundance of T. antarctica RS. C. Relative abundance of P. glacialis RS. D. Ratio of P. glacialis
798	RS: <i>T. antarctica</i> RS. E-G. Svenner Channel records. E. Relative abundance of <i>T. antarctica</i> RS.
799	F. Relative abundance of P. glacialis RS. G. Ratio of P. glacialis RS:T. antarctica RS.
800	Neoglacial and Hypsithermal intervals are defined following Crosta et al. (2005).
801	
802	Figure 4: Schematic diagram illustrating differences modelled sea ice concentrations and surface
803	temperatures between Hypsithermal and Neoglacial. Differences significant at the 99% confidence

804 interval are shown with horizontal lines and differences at the 95% confidence interval are shown

805 with horizontal curves. Where the Hypsithermal and Neoglacial lines overlap there is no

statistically significant difference between the two records.

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				Water denth	Distance from	Core length/	Interval(s) analysed	Age
Location	Core	Latitude	Longitude	/ m	coast / km	m	/ m	/cal yr BP ¹
Mertz-	JPC10	66°34.334′S	143°05.168′E	850	~30	21.35	Thin sections:	6756-11384
Ninnis	KC10A			\mathbf{C}		2.50	17.36-20.60	3820-3892
Trough				6			2.05-2.38	
Dumont	MD03-2597	66°24.74′S	140°25.26′E	1025	~30	57.34	Thin sections:	925-2814
d'Urville							13 x 15-cm-intervals	
Trough							between 18.75-56.83	
Dumont	MD03-2601	66°03.07'S	138°33.43′E	746	~60	40.24	Diatom abundance:	
d'Urville			2.				0-40.25	1000-10843
Trough							Thin sections:	
							6.19-6.485	~2550
							& 18.808-19.107	~5635
Iceberg	JPC43B	66°55.943′S	64°07.376′E	465	~75	23.96	Thin sections:	10500-11500
Alley			2				19.30-23.96	
Svenner	JPC24	68°41.660′S	76°42.557′E	848	~60	15.35	Diatom abundance:	655-11082
Channel		6					0-17.00	
	Location Mertz- Ninnis Trough Dumont d'Urville Trough Dumont d'Urville Trough Iceberg Alley Svenner Channel	LocationCoreMertz-JPC10NinnisKC10ATroughMD03-2597DumontMD03-2597d'UrvilleMD03-2601DumontMD03-2601d'UrvilleJPC43BAlleyJPC24ChannelIceberg	LocationCoreLatitudeMertz- NinnisJPC1066°34.334'SNinnisKC10A66°34.334'STrough DumontMD03-259766°24.74'Sd'Urville TroughMD03-260166°03.07'SDumont d'Urville TroughMD03-260166°03.07'Sd'Urville Alley SvennerJPC43B66°55.943'SAlley ChannelJPC2468°41.660'S	LocationCoreLatitudeLongitudeMertz- NinnisJPC1066°34.334'S143°05.168'ENinnisKC10A143°05.168'ETroughMD03-259766°24.74'S140°25.26'EDumont d'Urville TroughMD03-260166°03.07'S138°33.43'EDumont d'Urville TroughMD03-260166°55.943'S64°07.376'EIceberg Alley SvennerJPC2468°41.660'S76°42.557'E	Location Core Latitude Longitude Water depth /m Mertz- Ninnis JPC10 66°34.334'S 143°05.168'E 850 Ninnis KC10A 140°25.26'E 1025 Trough MD03-2597 66°24.74'S 140°25.26'E 1025 d'Urville MD03-2601 66°03.07'S 138°33.43'E 746 Dumont MD03-2601 66°03.07'S 138°33.43'E 746 d'Urville MD03-2601 66°03.07'S 138°33.43'E 746 lceberg JPC43B 66°55.943'S 64°07.376'E 465 Alley JPC24 68°41.660'S 76°42.557'E 848	LocationCoreLatitudeLongitudeWater depthDistance depthMertz-JPC10 $66^{\circ}34.334'S$ $143^{\circ}05.168'E$ 850 ~ 30 NinnisKC10A143^{\circ}05.168'E 850 ~ 30 TroughMD03-2597 $66^{\circ}24.74'S$ $140^{\circ}25.26'E$ 1025 ~ 30 DumontMD03-2597 $66^{\circ}03.07'S$ $138^{\circ}33.43'E$ 746 ~ 60 d'UrvilleMD03-2601 $66^{\circ}03.07'S$ $138^{\circ}33.43'E$ 746 ~ 60 d'UrvilleFrough $56^{\circ}5.943'S$ $64^{\circ}07.376'E$ 465 ~ 75 AlleyJPC24 $68^{\circ}41.660'S$ $76^{\circ}42.557'E$ 848 ~ 60	LocationCoreLatitudeLongitudeWater depthDistance from length/ coastCore length/ length/ mMertz- NinnisJPC10 $66^{\circ}34.334'S$ $143^{\circ}05.168'E$ 850 ~ 30 21.35 Ninnis TroughKC10A140°25.26'E 1025 ~ 30 21.35 Dumont d'Urville TroughMD03-2597 $66^{\circ}24.74'S$ $140^{\circ}25.26'E$ 1025 ~ 30 57.34 Dumont d'Urville 	$\begin{array}{c c c c c c c c c c c c c c c c c c c $

Age models from Denis et al. (2006), Maddison (2006), Maddison et al. (2006), and Leventer et al. (2006)

	P. glacialis RS		Mean Thickness	No. of	Minimum	Maximum
Location	or T. antarctica RS	Sediment Increment Type	(standard deviation)	ments (n)	/mm	/mm
Mertz-Ninnis Trough	P. glacialis RS	Annual thickness	17.5 (16.9)	116	1.8	103.2
(NBP0101 JPC10)	0	Annual thickness if <i>P. glacialis</i> RS sub-	20.6 (14.6)	22	3.0	53.3
Early Holocene		laminae are present				
-		<i>P. glacialis</i> RS sub-lamina thickness	1.4 (1.1)	22	0.5	5.5
Dumont d'Urville Trough	P. glacialis RS	Annual thickness	18.1 (9.6)	119	1.1	46.7
(MD03-2597)	0	Annual thickness if <i>P. glacialis</i> RS sub-	20.25 (9.1)	26	7.9	40.6
Neoglacial		laminae are present				
-		<i>P. glacialis</i> RS sub-lamina thickness ¹	2.2 (1.4)	35	0.5	5.9
Dumont d'Urville Trough						
(MD03-2601) Neoglacial	—	Annual thickness	21 (14)	10		
Hypsithermal	T. antarctica RS	Annual thickness	46 (16)	6		
		T. antarctica RS sub-lamina thickness	0.371 (0.289)	8		
Iceberg Alley	T. antarctica RS	Annual thickness	21.59 (17.69)	223	1.04	99.05
(NBP0101 JPC43B)		Annual thickness if T. antarctica RS sub-	21.98 (16.16)	67	1.86	81.32
Deglacial		laminae are present				
		<i>T. antarctica</i> RS sub-lamina thickness ²	1.16 (1.70)	68	0.03	10.47
		Summer thickness	5.35 (5.45)	159	0.28	33.49
		Summer thickness if T. antarctica RS sub-	5.74 (6.86)	46	0.33	33.49
		laminae are present				
		T. antarctica RS sub-lamina thickness if	1.33 (2.04)	43	0.03	10.47
		associated with top of summer lamina ³				
		Transitional lamina thickness	9.52 (9.28)	110	0.73	49.76
		Transitional lamina thickness if followed by <i>T</i> .	8.29 (4.49)	25	1.26	18.42
		antarctica RS sub-lamina				
		T. antarctica RS sub-lamina thickness if	0.88 (0.81)	25	0.25	3.28
		associated with top of transitional lamina ⁴				

 ¹ 26 years have *P. glacialis* RS sub-laminae; 5 years have multiple sub-laminae
 ² 67 years have *T. antarctica* RS sub-laminae; 1 year has 2 sub-laminae
 ³ 46 years have *T. antarctica* RS sub-laminae; in 43/46 years the sub-lamina is at the top of the summer lamina; in 3/46 years, the sub-lamina is within the summer laminae

⁴ Four transitional laminae, overlain by *T. antarctica* RS sub-laminae, are followed by summer laminae associated with *T. antarctica* RS sub-laminae

			Mean Abundance (standard	No. of			
		P. glacialis RS or T. antarctica RS	deviation)	Measure-	Minimum	Maximum	Relative
T	Sub-lamina	abundance in lamina or sub-lamina	/ x 10°	ments	$/ \times 10^{\circ}$	/ x 10°	Abundance (%)
Location	type	types	valves/g	(n)	valves/g	valves/g	All/CK5-Iree
Mertz-Ninnis	P. glacialis RS	P. glacialis RS abundance in summer	6.9 (2.7)	5	3.9	9.5	0.38 / 2.29
Trough		mixed diatom lamina	5				
(NBP0101 JPC10)		T. antarctica RS abundance in	0 (0)	5	0	0	0/0.42
		summer mixed diatom lamina					
		<i>P. glacialis</i> RS abundance in <i>P.</i>	7.2 (3.9)	4	3.2	10.6	0.24 / 11.35
		glacialis RS sub-lamina					
		<i>T. antarctica</i> RS abundance in <i>P</i> .	0 (0)	4	0	0	0 / 0
		glacialis RS sub-lamina					
Dumont d'Urville	P. glacialis RS	P. glacialis RS abundance in summer	20.3 (9.9)	5	10.9	34.8	2.91 / 5.1
Trough	C	mixed diatom lamina					
(MD03-2597)		T. antarctica RS abundance in	2.4 (3.6)	5	0	8.6	0.39 / 0.81
· /		summer mixed diatom lamina	. ,				
		P. glacialis RS abundance in P.	53.9 (27.2)	3	33.5	84.8	7.08 / 15.48
		glacialis RS sub-lamina	~ /				
		<i>T. antarctica</i> RS abundance in <i>P</i> .	6.7 (6.7)	3	0	13.4	0.89/3.16
		glacialis RS sub-lamina	~ /				
		Y					

	Jan	Feb	Mar	Apr	May	Jun	Jul	Aug	Sep	Oct	Nov	Dec
Mean Hypsithermal surface temperature (°C)	0.79	1.58	0.56	-0.94	-1.55	-1.62	-1.61	-1.61	-1.63	-1.65	-1.53	-0.63
Mean Neoglacial surface temperature (°C)	0.68	1.59	0.63	-0.90	-1.54	-1.61	-1.61	-1.61	-1.63	-1.65	-1.58	-0.78
z value	2.79	-0.16	-2.11	-1.97	-2.05	-1.84	-0.18	0.55	0.22	-2.16	7.82	7.07
Significance level of difference between the					X							
mean surface temperatures	99%	none	95%	95%	95%	90%	none	none	none	95%	>99%	>99%
					5							
Mean Hypsithermal sea ice concentration (%)	0.29	0.25	0.33	0.53	0.80	0.88	0.88	0.87	0.83	0.76	0.58	0.41
Mean Neoglacial sea ice concentration (%)	0.28	0.24	0.30	0.50	0.79	0.88	0.89	0.88	0.85	0.77	0.59	0.41
z value	2.15	3.20	4.61	4.86	2.52	-0.53	-3.42	-3.72	-3.80	-2.76	-1.38	0.87
Significance of difference between the mean				N.								
sea ice concentrations	95%	>99%	>99%	>99%	>98.5%	none	>99%	>99%	>99%	99%	none	none

<u>% >99% >99% ></u>



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10 µm

Figure 3



Figure 4

