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3	The origin of lithogenic sediment in the southwestern Ross Sea and implications
4	for iron fertilisation
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Austral summer iron (Fe) fertilisation in the Ross Sea has previously been observed in 27 28 association with diatom productivity, lithogenic particles and excess Fe in the water column (Collier et al., 2000). This productivity event occurred during an early break 29 30 out of sea ice via katabatic winds, suggesting that aeolian dust could be an important 31 source of lithogenic Fe required for diatom growth in the Ross Sea. Here we 32 investigate the provenance of size-selected dust deposited on sea ice in McMurdo 33 Sound, southwestern (SW) Ross Sea. The isotopic signature of McMurdo Sound dust  $(0.70533 < {}^{87}Sr/{}^{86}Sr < 0.70915$  and  $-1.1 < \epsilon_{Nd}(0) < 3.45$ ) confirms that dust is locally 34 35 sourced from the McMurdo Sound debris bands and comprises a two-component 36 mixture of McMurdo Volcanic Group and Southern Victoria Land lithologies. In 37 addition, we investigate the provenance of lithogenic sediment trapped in the water column: the isotopic signature ( $\epsilon_{Nd}(0)=3.9$ ,  ${}^{87}Sr/{}^{86}Sr=0.70434$ ) is differentiated from 38 39 long-range transport dust originating from South America and Australia. Elevated 40 lithogenic accumulation rates in deeper sediment traps in the Ross Sea, suggest that 41 sinking particles in the water column cannot simply result from dust input at the 42 surface. This discrepancy can best be explained by significant upwelling and 43 remobilisation of lithogenic Fe from the sea floor.

44

#### 45 Key words

- 46 Dust provenance, dust, iron, McMurdo Sound, Antarctica
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# 51 **1. Introduction**

52 Atmospheric dust is potentially an important source of dissolved iron (DFe) which is 53 the limiting nutrient required for primary production in vast regions of the remote 54 Southern Ocean, including Antarctica's marginal seas (Boyd et al., 2010; Sedwick et 55 al., 2000). Despite being seasonally iron (Fe) limited, the high-nutrient, high 56 chlorophyll (HNHC) regime of the Ross Sea is the most biologically productive continental shelf region in Antarctica, and supports intense phytoplankton blooms in 57 58 the austral summer (Arrigo et al., 2008). Although the flux of Fe into the Ross Sea 59 plays a critical role in determining its productivity, the origin(s) of this Fe remains 60 poorly constrained (Sedwick et al., 2011).

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### 62 1.1. Dust deposition in Antarctica

63 Global 'background' dust is characterized by fine particles having a mass modal 64 diameter  $<5 \,\mu$ m, long atmospheric residence time and modern mass deposition rates in the order of 0.001-0.02 g m<sup>-2</sup> yr<sup>-1</sup> in the Southern Ocean (Wagener et al., 2008; and 65 references therein). The isolated, snow and ice-covered central East Antarctic Plateau 66 67 (EAP) has proven to be an excellent location for investigating long-range transport of 68 dust representative of the broader Southern Hemisphere, both at present and in the 69 past (Delmonte et al., 2008; Delmonte et al., 2007). Moreover, the high East Antarctic Plateau (EAP) has much lower accumulation rates of around 0.0002-0.0006 g m<sup>-2</sup> yr<sup>-1</sup> 70 71 during the Holocene (Albani et al., 2012). Recently, it has become apparent that 72 peripheral areas of the Antarctic ice sheet, close to high-elevation ice-free mountain 73 ranges, such as the Transantarctic Mountains (TAM), can receive significant 74 additional dust inputs from exposed Antarctic sources, some of which have been ice-75 free for millions of years (Delmonte et al., 2013 and references therein).

77 The relative contribution of much smaller, patchy but proximal dust sources to the 78 atmospheric dust load over Antarctica and the Southern Ocean is not well known. The 79 largest expanse of contiguous ice-free ground in Antarctica is found in the McMurdo Dry Valleys - a series of west-to-east-oriented, glacially carved valleys located 80 81 between the high EAP and the Ross Sea in Southern Victoria Land. However, the 82 dustiest known place in Antarctica is located in the southwestern (SW) Ross Sea, 83 associated with the so-called 'debris bands' area on the McMurdo Ice Shelf (Kellogg et al., 1990) (Fig. 1). In this region, dust deposition flux (~1 g m<sup>-2</sup> yr<sup>-1</sup>) is at least two 84 85 orders of magnitude greater than fallout of long-range transport dust measured in ice 86 cores from the EAP (Atkins & Dunbar, 2009; Chewings et al., 2014; Delmonte et al., 87 2013) and is, potentially, an important source of bioavailable Fe to the Ross Sea 88 (Winton et al., 2014).

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Dust provenance in Antarctica can be determined from the <sup>87</sup>Sr/<sup>86</sup>Sr and <sup>143</sup>Nd/<sup>144</sup>Nd 90 91 radiogenic isotope composition of dust in snow and ice by comparison with potential 92 source areas (PSAs) (e.g. Delmonte et al., 2010). This geochemical method allows 93 mantle-derived (basaltic rocks, tephra and soils derived from them, weathered and 94 eroded mafic rocks) and crustal-derived sediments and soils to be identified. Both the 95 geochemical fingerprint and particle size of dust deposited on the EAP suggests it 96 originates from arid regions in southern South America during glacial periods (Delmonte et al., 2008; Gaiero et al., 2007). However, for dust deposited during 97 98 interglacial periods, that is when dust input to inner Antarctica was extremely low, the 99 source is less certain (Delmonte et al., 2007), and an Australian contribution is likely 100 (Delmonte et al., 2008; Delmonte et al., 2007; Revel-Rolland et al., 2006). In addition

101 to atmospheric circulation, dust transport efficiency is dependent on particle size; for 102 example, long-range dust deposited on the EAP has a mass-modal size of  $\sim 2-3 \ \mu m$ 103 (Delmonte et al., 2002). When investigating the provenance of dust, the fractionation 104 of Sr isotopes into different grain size fractions needs to be considered, as there is a correlation between grain size and <sup>87</sup>Rb/<sup>86</sup>Sr ratios and thus <sup>87</sup>Sr/<sup>86</sup>Sr ratios. In coarse 105 106 (fine) grained suspended particulate matter Sr is enriched (depleted) in less radiogenic 107 Sr isotopic ratios (e.g. Andersson et al., 1994). In contrast, Nd isotopic ratios are not 108 influenced to the same extent by particle size (e.g. Andersson et al., 1994).

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# 110 1.2. Iron-fertilisation in the Ross Sea

111 The Ross Sea is one of the most productive regions in the Southern Ocean and an 112 important oceanic sink for atmospheric carbon dioxide (CO<sub>2</sub>) (e.g. Arrigo et al., 2008). The environmental factors responsible for controlling the rates of 113 114 phytoplankton production and incomplete utilisation of inorganic macronutrients 115 include: grazing (Banse, 1991), temperature (Bunt & Wood, 1963), light availability 116 (e.g. Mitchell et al., 1991), micro-nutrient availability (e.g. Fe and Mn) (Sedwick & 117 DiTullio, 1997; Sedwick et al., 2000), or a combination of these (e.g. Arrigo et al., 118 2000). Collier et al. (2000) show that a Ross Sea diatom productivity event, captured 119 during the 1996-1997 deployment of moored Antarctic Environment and Southern 120 Ocean Process Study (AESOPS) sediment traps, is correlated with elevated lithogenic 121 particle accumulation rates, excess Fe (determined from high Fe/Al ratios), and with 122 an early breakout of sea ice caused by katabatic winds. They go on to suggest that 123 there may be a causal relationship between the retreat of sea ice, the supply of 124 particulate Fe and diatom production and export. The source of this lithogenic Fe to 125 the Ross Sea is unknown but could be derived from either dust released into the ocean 128

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126 from melting sea ice from local and/or from distal sources, or new particulate Fe 127 derived from ice shelves, icebergs, upwelling of resuspended continental sediments from the sea floor, circumpolar deep water or some combination thereof.

130 Multiple sources of new Fe to the Ross Sea region have been identified, which 131 include local dust sourced mainly from the McMurdo Ice Shelf (Atkins & Dunbar, 132 2009; Chewings et al., 2014; de Jong et al., 2013; Winton et al., 2014), sea ice melt 133 (de Jong et al., 2013; Sedwick & DiTullio, 1997), and lithogenic sediments 134 resuspended from the sea floor (de Jong et al., 2013; Gerringa et al., 2015; Marsay et 135 al., 2014; Sedwick et al., 2011). However, the relative importance of these sources for 136 stimulating primary production remains an open question. Winton et al. (2014) 137 estimate that the supply of soluble aeolian Fe in dust from the debris bands, southern McMurdo Sound to the adjacent ocean could support up to ~15 % of primary 138 139 production in the area. The implication being that Fe supporting the remaining 85 % 140 of productivity was derived largely from other sources, such as lithogenic sediment 141 resuspended from the sea floor (de Jong et al., 2013; Gerringa et al., 2015; Kustka et 142 al., 2015; Marsay et al., 2014; McGillicuddy et al., 2015; Sedwick et al., 2011).

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144 As Fe is critical for seasonal phytoplankton growth in the Ross Sea, this study aims to 145 further investigate the source(s) of lithogenic Fe as a driver of the vast austral summer 146 phytoplankton blooms in the SW Ross Sea. We do this by examining the provenance 147 of lithogenic material sinking in the upper 200 meters below sea level (mbsl) of the 148 water column and compare its origin to both known local and global sources. Here we 149 report the Sr-Nd isotopic composition of i) size-selected dust from snow samples on 150 sea ice from McMurdo Sound, and ii) sediment trap material from the Research on Ocean - Atmosphere Variability and Ecosystem Response in the Ross Sea (ROAVERRS) moorings program (1996-98) that represents accumulation of sediment settling out of the water column. When investigating PSAs to Antarctica, previous studies have size-selected the PSA samples prior to Sr analysis to be comparable to that of the fine size range of dust deposited in Antarctica (Delmonte et al., 2008), and a similar approach is used here.

157

158 **2. Methods** 

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160 2.1. Samples used in this study

Previous studies have focused on dust flux and particle size distribution patterns in
McMurdo Sound (Atkins & Dunbar, 2009; Chewings et al., 2014; Dunbar et al.,
2009). This study is based on the following samples:

- Samples of dust-laden snow collected from sea ice along a south-north X-Y
  transect in McMurdo Sound, collected in November 2010 and described in
  Chewings et al. (2014) and in Winton et al. (2014) (Fig. 1b).
- 167 A Ross Sea sediment trap sample from 200 mbsl, collected between 25 Dec. 1997
- and 3 Jan. 1998 from the ROAVERRS program Chinstrap site (76° 20.5'S, 165°
- 169 1.78'E) in the SW Ross Sea. This site was anchored in 830 m water depth in the
- 170 southern extension of the Drygalski Basin (Fig. 1a).
- 171
- 172 2.2. Nd and Sr isotopic ratios and concentrations
- 173

174 2.2.1. Sample processing

176 McMurdo Sound surface snow on sea ice samples

177 The samples analysed in this work were size-selected in order to be comparable to 178 provenance measurements made on dust from ice core PSAs and with similar studies 179 on dust in Antarctica (e.g. Delmonte et al., 2008; Delmonte et al., 2010; Delmonte et 180 al., 2004). We analysed both the bulk (all particle sizes) and fine ( $<10 \mu m$ ) fraction of 181 McMurdo Sound dust to check for particle size induced bias in the isotopic 182 fractionation of samples. The coarse fraction was removed from bulk samples by 183 using a pre-washed 10 µm SEFAR Nitex® open mesh while the fraction 0.4µm<Ø<10µm was collected on 0.4 µm Isopore<sup>TM</sup> polycarbonate membranes. After 184 185 filtration, the membranes were put into pre-cleaned Corning® tubes filled with ~10 186 ml of ultra-pure water, and micro-particles were removed from the filter by 187 sonication. Samples were transported to the Department of Geosciences, Swedish 188 Museum of Natural History, Sweden where the liquid was evaporated in acid-cleaned 189 15 ml Savillex® beakers. Dry dust samples, ranging between 0.1 and 1.2 mg, were 190 weighed a minimum of five times to obtain a mean weight, which was used for 191 subsequent calculations.

192

193 Sediment trap samples

Isotopic analysis of bulk sediment revealed that the biogenic fraction of the sediment (~up to 70 % total mass estimated from AESOPS sediment trap data reported in Collier et al. (2000)) incorporated marine Sr and thus the isotopic signature could not be distinguished from that of seawater (Table 1). To remove the biogenic silica and calcium carbonate fraction of the sediment we leached the sediment with 6 M HCl in Savillex® beakers and centrifuged following the method of Freydier et al. (2001). The lithogenic residue was then rinsed three times with ultra-pure water and dried. 201

## 202 2.2.2. Sample digestion

203 The chemical treatment of the dust samples and leached sediment, including digestion 204 and elemental separation (Rb-Sr and Sm-Nd) using ion exchange chromatography, 205 was performed at the Swedish Museum of Natural History following the established method Delmonte et al. (2008). The samples were spiked with a mixed <sup>147</sup>Sm/<sup>150</sup>Nd 206 207 spike and <sup>84</sup>Sr-enriched spike for the isotope dilution determination of the 208 concentrations. Samples were digested in an acid mixture of 1.5 ml, of HNO<sub>3</sub>, HF and 209 HClO<sub>4</sub> heated to 90 °C in closed Savillex® beakers for 24 h. The solution was 210 evaporated to complete dryness on a hot plate, and the residue re-dissolved in 4 ml 6 211 M HCl.

212

### 213 *2.2.3. Ion exchange*

To achieve separation of potential interfering elements (Fe, Ba, Rb, Sm, Ce, and Pr), and obtain high column yield and low blanks, the residue was subjected to chemical procedures described in Delmonte et al. (2008). The total blank, including dissolution, chemical separation and mass spectrometry, was frequently monitored in each ion exchange batch and blank concentrations were <5 pg for Nd and <130 pg for Sr.

219

220 2.2.4. Mass spectrometry

Isotopic analysis of Nd and Sr was performed with a Thermo Scientific TRITON Thermal Ionisation Mass Spectrometer (TIMS). Neodymium was loaded mixed with colloidal graphite, Alfa Aesar, on double rhenium filaments and analysed as metal ions in static mode using rotating gain compensation. Concentrations and ratios were calculated assuming exponential fractionation. The calculated ratios were normalised to  $^{146}$ Nd/ $^{144}$ Nd = 0.7219. Epsilon units are calculated as follows:

228 
$$\varepsilon_{Nd}(0) = [({}^{143}Nd/{}^{144}Nd)_{sample}/({}^{143}Nd/{}^{144}Nd)_{CHUR} - 1] \times 10^4;$$

229 CHUR, chondritic uniform reservoir with 
$$(^{143}Nd/^{144}Nd)_{CHUR} = 0.512638$$

230

The external precision for <sup>143</sup>Nd/<sup>144</sup>Nd is estimated from analysis of the nNdß 231 232 standard (Wasserburg et al., 1981) by analysing a range, 4-12 ng loads, of nNdß 233 standard. The external precision becomes larger for smaller loads, with an estimated 234 precision of about 40 ppm for small loads, for intermediate size about 30 ppm and 235 about 20 ppm for larger loads. These values have been used to estimate the errors for the samples in Table 1. The mean  $^{143}$ Nd/ $^{144}$ Nd ratio for the nNdß was 0.511895±22 236 237 (n=20). Literature values for repeated analysis of standard nNdß (Andreasen & Sharma, 2006) yielded  ${}^{143}$ Nd/ ${}^{144}$ Nd = 0.511892±3 (2 $\sigma$ , n = 23) and thus no accuracy 238 239 correction was applied.

240

Purified Sr samples were mixed with tantalum activator and loaded on a single 241 242 rhenium filament. Two hundred 8 s integrations were recorded in multi-collector static mode, applying a rotating gain compensation. Measured <sup>87</sup>Sr intensities were 243 244 corrected for Rb interference assuming <sup>87</sup>Rb/<sup>85</sup>Rb=0.38600 and ratios were calculated 245 using the exponential fractionation law and <sup>88</sup>Sr/<sup>86</sup>Sr=8.375209. External precision for <sup>87</sup>Sr/<sup>86</sup>Sr, estimated from analysing NBS SRM987 standard, was calculated as 246 247  $\pm 0.000016$  (n=12) while repeated measurements of prepared CIT #39 sea water gave 248 a reproducibility of  $\pm 0.0000082$  or 12 ppm (n=21) which was taken to be the best estimate of the external precision. Accuracy correction was applied to the <sup>87</sup>Sr/<sup>86</sup>Sr 249 ratios corresponding to a <sup>87</sup>Sr/<sup>86</sup>Sr ratio of 0.710245 for NBS SRM 987 standard 250

251 (NBS 987: literature value 0.710245, Department of Geosciences value 0.710217 ±16
252 (n=12); difference: 0.000028).

253

The accuracy of the Nd and Sr isotopic composition of small dust samples was determined using the Basalt Columbia River rock standard (BCR-2), a certified reference material. Preparation and analysis of 150 to 600  $\mu$ g aliquots of BCR-2 in each batch of ion exchange resulted in a recovery of >79 % (n=6) for concentration and >99 % (n=6) for isotopic composition. Due to the small dust samples and the difficulty of weighing such small masses we estimate a Sr and Nd concentration error of ±10 % estimated by repeated weighing of BCR-2 standards (~0.3 mg).

261

#### 262 **3. Results**

263 The Sr and Nd isotopic composition of dust are primarily related to lithology and 264 geologic age of parent materials, although the Sr isotopic composition, for particles 265 between 2-50 µm can also be influenced by their size. The Sr and Nd isotopic 266 composition of the fine (<10 µm) and bulk (all sizes included) dust samples collected 267 in this study are well characterised and reported in Table 1 and Fig. 2 along with additional isotopic data from McMurdo Sound measured in an earlier study (Winton 268 et al., 2014). Samples in Fig. 2 are grouped by geographic location. The samples 269 270 collected from the snow on sea ice in McMurdo Sound display a narrow isotopic composition (0.70533<  ${}^{87}$ Sr/ ${}^{86}$ Sr <0.70915 and -1.1<  $\epsilon_{Nd}(0)$  <3.45). Our fine dust 271 (<10  $\mu$ m) samples have relatively higher <sup>87</sup>Sr/<sup>86</sup>Sr ratios compared to the bulk samples 272 273 (Fig. 2), consistent with previous studies of size-dependent fractionation (Andersson 274 et al., 1994). The Nd isotopes do not fractionate with the particle size (with the 275 exception of MPN10-5 which could be related to the different size fractions

276 originating from different sources; Fig. 2), also consistent with previous studies. The  $\Delta^{87}$ Sr/<sup>86</sup>Sr is about 0.00115, that is slightly smaller than the  ${}^{87}$ Sr/<sup>86</sup>Sr increase of 277 278 ~0.0028 units observed between 63  $\mu$ m and 2  $\mu$ m dust particles by Gaiero et al. 279 (2007). The isotopic ratios of leached (lithogenic sediment fraction) and unleached 280 (lithogenic and biogenic sediment fraction) Ross Sea sediment obtained from the 281 upper Chinstrap sediment trap (200 mbsl) are reported in Table 1. Leaching had a 282 significant effect on the Sr isotopic ratio of this sample and removed a Sr seawater 283 overprint from the sediment. After leaching, the remaining lithogenic sediment has an isotopic signature between the McMurdo Volcanic Group and Southern Victoria Land 284 285 PSAs (Fig. 2).

286

287 **4. Discussion** 

288

289 *4.1. Dust provenance* 

290

291 4.1.1. McMurdo Sound

292 The overwhelming majority of dust deposited in snow on sea ice in McMurdo Sound 293 is locally sourced. It is not possible to detect any contribution from South American 294 or Australian sources with our approach. Sedimentological, meteorological and 295 geochemical evidence consistently points to the debris bands on the McMurdo Ice 296 Shelf (Kellogg et al., 1990) as the dominant local dust source in the McMurdo Sound 297 region (Fig. 1) (Atkins & Dunbar, 2009; Chewings et al., 2014; Winton et al., 2014). 298 Studies of spatial variability of dust and particle size for the greater McMurdo Sound 299 region show a distinct decrease in particle size and dust flux along transect X-Y (Fig. 300 1b) as part of a dust plume extending northwards from the debris bands (Atkins &

301 Dunbar, 2009; Chewings et al., 2014). As the plume extends northwards away from 302 the debris bands the particle size and dust accumulation rate decrease, although 303 secondary elevated patches of both occur near coastal headlands. Overall dust accumulation declines exponentially from 55 g m<sup>-2</sup> yr<sup>-1</sup> near the debris bands to  $\sim 0.2$ 304 g m<sup>-2</sup> yr<sup>-1</sup> 120 km north of the debris bands (Atkins & Dunbar, 2009; Chewings et al., 305 306 2014). This northward dust dispersal is consistent with the local meteorology whereby 307 the highest wind speeds, i.e. those most competent with respect to entraining silt and 308 fine sand, are predominately from the south (Fig. 1c), dispersing dust from the debris 309 bands north along the Southern Victoria Land coastline.

310

311 In addition to sedimentological considerations, geochemical evidence also points to 312 dust being locally sourced. The Sr isotopic of modern seawater is homogenous (<sup>87</sup>Sr/<sup>86</sup>Sr=0.70924; Elderfield (1986)), and has a similar Sr isotopic composition to 313 314 the geology in McMurdo Sound (Fig. 2). However, the Nd isotopic composition of 315 the local geology and Ross Sea seawater (-10 $\leq E_{Nd}(0) \leq -6$ ; Basak et al. (2015)) is 316 distinguished in Fig. 2, and combined with other provenance indicators (coarse 317 particle size, high dust flux and Fe/Al elemental ratios (Atkins & Dunbar, 2009; de 318 Jong et al., 2013); see below, allows tracing of dust to local PSAs. Winton et al. 319 (2014) report two Sr and Nd isotopic ratios of the bulk sediment from snow on sea ice 320 in southern McMurdo Sound and one from Granite Harbour (Fig. 1). The values are 321 consistent with dust originating from McMurdo Volcanic Group (MVG), although 322 within the Granite Harbour embayment there is also evidence for dust-derived from 323 TAM lithologies. We rule out the possibility that the volcanic signature of McMurdo Sound dust on sea ice is derived from volcanic rocks in Marie Byrd Land (0.7026< 324  $^{87}Sr/^{86}Sr$  <0.7032 and 1.99<  $\epsilon_{Nd}(0)$  <6.87) (Futa & Le Masurier, 1983; Hole & 325

326 LeMasurier, 1994) due to the northerly direction of the prevailing winds (Chewings et327 al., 2014).

328

329 Consequently, only local potential source areas (PSAs) are considered for comparison to the new isotopic dataset. Overall, the Sr isotopic ratios for McMurdo Sound 330 331 samples analysed in this study and in Winton et al. (2014) are tightly grouped and 332 range between  $0.705 < {}^{87}\text{Sr} < 0.709$  while  $\varepsilon_{Nd}(0)$  ranges between  $3.45 < \varepsilon_{Nd}(0) < -1.1$ . 333 These new isotopic data form a linear array in Fig. 2. McMurdo Sound dust can be 334 considered the result of a two-component mixture derived from isotopically distinct 335 end-members: i) the MVG volcanic rocks and ii) Southern Victoria Land lithologies 336 found in the TAM such as Ferrar Dolerites and Beacon sandstone (Fig. 2). The 337 narrow range of isotopic ratios of McMurdo Sound dust along the south-north transect 338 X-Y represents northwards dust dispersal downwind from the debris bands, that is a 339 mixture of TAM and MVG sources, with minor localised additions of TAM dust 340 input from coastal outcrops from New Harbour and Marble Point that contribute to 341 the dominant south to north dust plume (Fig. 1). This is consistent with field 342 observations showing sediment on the McMurdo Ice Shelf debris bands is itself a 343 mixture of MVG and TAM lithologies (Kellogg et al., 1990).

344

Previous studies have shown dust deposited within embayments or adjacent to headlands along the Southern Victoria Land coastline is not widely dispersed (Barrett et al., 1983; Chewings et al., 2014; de Jong et al., 2013). Within the narrow range of isotopic ratios of McMurdo Sound dust, GH9 is isotopically distinct and displays a dominant TAM signature (Fig. 2). This sample is not situated under the main northward-directed dust plume and hence represents localised dust accumulation within the Granite Harbour embayment (Fig. 1.). In contrast, the isotopic composition of GH2 lies within the tight cluster of McMurdo Sound dust and thus highlights that the mass of dust on the sea ice immediately seawards of Granite Harbour originates from the south. A single-source from the debris bands is also consistent with Fe concentrations within dust samples that were uniform along the transport pathway (Winton et al., 2014).

357

### 358 4.1.2. Southwestern Ross Sea

359 The isotopic signature of the lithogenic fraction of sediment from the upper 200 mbsl Chinstrap sediment trap, located ~170 km north of the debris bands ( ${}^{87}$ Sr/ ${}^{86}$ Sr=0.704, 360 361  $E_{Nd}(0)=3.9$ ), falls outside of the isotopic range of dust originating in Australia  $(0.709 < {}^{87}\text{Sr} / {}^{86}\text{Sr} < 0.763, -2.9 < \mathcal{E}_{Nd}(0) < -15.4$ ; Delmonte et al. (2004); Revel-Rolland et 362 al. (2006)) and South America ( $0.704 < {}^{87}Sr/{}^{86}Sr < 0.713$ ,  $-8.9 < \epsilon_{Nd}(0) < -8.3$ ; Delmonte et 363 364 al. (2004)). These two potential Southern Hemisphere sources supply dust to the high 365 elevation EAP at very low deposition rates (e.g. Delmonte et al., 2008). The signature of the lithogenic fraction of sediment from SW Ross Sea (Chinstrap) matches that of 366 the local geology and dust on sea ice in McMurdo Sound. Thus, the lithogenic 367 particles, and their associated Fe, collected here is 'locally' sourced from the Ross Sea 368 369 region (Fig. 2).

370

# 371 4.1.3. Dust transport and deposition in the southwestern Ross Sea

372 Deposition of local dust into the SW Ross Sea can occur by direct atmospheric fallout 373 into ice-free surface waters, and released into surface waters by sea ice melt 374 associated with subsequent northwards advection (Atkins & Dunbar, 2009; Chewings 375 et al., 2014; de Jong et al., 2013). The geographical area over which local dust is 376 transported into the Ross Sea and hence contributes to Fe-fertilisation is potentially 377 large. Although dust accumulation measurements only exist up to 120 km from the 378 debris bands and decrease exponentially from the source, local dust deposition likely 379 extends far beyond this point. Extrapolating the dust flux trend observed by Chewings 380 et al. (2014) suggests that the Chinstrap site may represent a northern extension of the 381 dust and DFe dispersal pattern reported by Winton et al. (2014). We estimate an annual accumulation rate of ~ $0.01 \text{ g m}^{-2} \text{ yr}^{-1}$  at the Chinstrap site from aeolian dust, 382 383 although we do not have accumulation rate data from the Chinstrap sediment trap 384 with which to compare this estimate. Whether or not locally sourced aeolian sediment 385 this is the main source of the lithogenic sediment in the Chinstrap site remains 386 somewhat of an open question. However, when the relationship between DFe and 387 phytoplankton productivity in McMurdo Sound is considered we suggest this is 388 unlikely to be the case (Fig. 3; Section 4.2.).

389

# 390 4.2. Implications for iron-fertilisation

391

## 392 4.2.1. Contribution of local dust to lithogenic iron

393 By extrapolating the dust flux trend observed by Chewings et al. (2014) and its 394 associated contribution to DFe (Winton et al. (2014) into the SW Ross Sea, we can 395 estimate the upper bound of the DFe at the Chinstrap site from the debris bands (Fig. 3). Assuming a lithogenic dust flux of  $\sim 0.01$  g m<sup>-2</sup> yr<sup>-1</sup> and an associated total Fe 396 397 content of 4 % and 11 % of this Fe is soluble (Winton et al., 2014), we estimate a maximum DFe flux of  $\sim 2$  nmol m<sup>-2</sup> d<sup>-1</sup> to the Chinstrap site (Fig. 3). However, when 398 399 this is considered relative to the spatial distribution of primary production in the SW Ross Sea, using averaged annual Sea-Viewing Wide Field-of-View Sensor 400

401 (SeaWiFS) satellite chlorophyll-a data, we find that the gradient in increasing 402 chlorophyll-a concentration with distance from the debris bands within the McMurdo 403 Sound polynya does not match the pattern of decreasing dust accumulation (Fig. 3). 404 This pattern suggests that DFe from dust not regulate growth in the SW Ross Sea. A 405 seasonal phytoplankton bloom occurs in the McMurdo Sound polynya, SW Ross Sea 406 each summer and is dominated by diatoms. The rate of primary production is greatest 407 in the centre of the McMurdo Sound Polynya. As the dust flux decreases and primary 408 production increases with distance from McMurdo Sound, it is difficult to reconcile 409 these patterns at the Chinstrap site assuming only a local dust source (Fig. 3).

410

# 411 4.2.2. Supporting evidence from the wider Ross Sea

412 Some further insight into the origin of sediment in the water column in the SW Ross 413 see may be inferred from the data published by Collier et al. (2000). They show a 414 significantly elevated lithogenic accumulation rate in deep AESOPS sediment traps 415 compared to accumulation rates measured in the upper AESOPS sediment traps at 416 other sites in the Ross Sea (e.g. MS-7; 76°30'S, 178°1'W). In addition, lithogenic Fe fluxes between 1-90  $\mu$ g m<sup>-2</sup> d<sup>-1</sup> have been measured for the upper 200 mbsl AESOPS 417 trap (MIS-7b) and 40-850  $\mu$ g m<sup>-2</sup> d<sup>-1</sup> for the deep trap (MS-7a). The greater mass of 418 419 sediment and lithogenic Fe flux in the deep traps in the Ross Sea highlight that 420 concentrations of suspended sediment in the water column at these sites cannot 421 simply result from sediment input at the surface (whereby the accumulation in each 422 trap would be the same regardless of depth). Instead, the increase in accumulation 423 with depth likely reflects resuspension and horizontal near-bottom transport 424 processes. Whilst we do not have the data to constraint these processes at Chinstrap 425 we infer, based on the provenance of the lithogenic sediment and distance from 426 known local sources, that the lithogenic sediment accumulating there is most likely 427 dominated by resuspended bottom sediments, potentially from sills either side of the 428 Drygalski Basin but also locally sourced material falling through the water column 429 sourced from atmospheric deposition or ice rafting. Although, no Fe flux data for the 430 Chinstrap site are available, future work could examine the relationship between Fe 431 fluxes in the Chinstrap sediment trap and those DFe fluxes reported for locally 432 derived dust at McMurdo Sound.

433

434 Despite McMurdo Sound representing the upper bound of locally derived dust and 435 associated DFe to Antarctic waters, previous studies have ruled out local dust as the 436 major source of DFe supply for phytoplankton blooms in the Ross Sea. Dissolved Fe 437 in McMurdo Sound dust can only support up to 15 % of primary production in the 438 region (Winton et al., 2014). Furthermore, based on regional scale estimates of dust deposition to the Southern Ocean, primary production triggered by long-range 439 440 transport dust is likely to be less significant than local dust (e.g. Edwards & Sedwick, 441 2001). Evidence from the extrapolation of the mass accumulation rate to the upper 442 200 mbsl Chinstrap trap, the sedimentological study of Chewings et al. (2014), 443 sedimentation in the water column, and the low contribution of local aeolian DFe to 444 phytoplankton blooms, suggest that it is unlikely that aeolian dust deposition is the 445 dominant process by which lithogenic Fe is supplied to the water column in the SW Ross Sea. Considered together, these lines of evidence point to a combination of 446 447 resuspended bottom sediments with smaller additions of local dust sourced from 448 atmospheric deposition or ice rafting as the sources of Fe-bearing sediment to the 449 water column in the SW Ross Sea.

### 451 4.2.3. Implications for iron-fertilisation

452 More broadly, Sedwick et al. (2011) noted that the phytoplankton-Fe limitation must 453 be overcome by continuous replenishment from new sources to sustain the significant 454 biomass observed over summer. They considered the following as potential sources of new DFe: vertical mixing, lateral advection, aerosol input, and dissolution of 455 456 particulate Fe from any or all of these sources. Consistent with upwelling of DFe as a 457 major source of DFe in the Ross Sea, Marsay et al. (2014) reported the highest DFe 458 concentrations are found within 50 m of the seafloor in the austral summer 2012. 459 Most recently, Gerringa et al. (2015) measured seawater DFe concentrations in the 460 2013-2014 austral summer, and concluded that DFe from the seafloor and land mass 461 sediments are the main sources of DFe which support phytoplankton in the upper 462 mixed layer of the Ross Sea Polynya in the early summer. Similarly, phytoplankton 463 blooms in the Pennell Bank region of the Ross Sea are supported by upwelling of DFe 464 (Kustka et al., 2015). However, Kustka et al. (2015) also highlight the spatial 465 variability of processes supplying DFe in the Ross Sea. For example, circulation patterns around bathymetric features can alter the input of DFe from increased 466 467 upwelling rates and higher concentrations of DFe. Numerical modelling of DFe supply by McGillicuddy et al. (2015) suggests that the largest sources to the euphotic 468 469 zone are wintertime mixing and melting sea ice (e.g. de Jong et al., 2013; Sedwick & 470 DiTullio, 1997) with smaller inputs from Circumpolar Deep Water and from melting 471 glacial ice.

472

# 473 Conclusions

474 Dust extracted from surface snow on McMurdo Sound sea ice enables us to document 475 the present day provenance of dust reaching the SW Ross Sea. Based on our 476 measurements of Sr and Nd isotopic ratios of dust deposited in surface snow on sea
477 ice at McMurdo Sound and in the Chinstrap sediment trap in the SW Ross Sea we
478 conclude the following:

479

480 1. The Sr and Nd isotopic signature of lithogenic sediment from the upper Chinstrap 481 sediment trap in the SW Ross Sea ( $\varepsilon_{Nd}(0)=3.9$ ,  ${}^{87}Sr/{}^{86}Sr=0.70434$ ) matches local dust 482 sources.

483

2. McMurdo Sound has been well characterised in terms of the Sr and Nd isotopic composition of locally derived dust deposited on sea ice. Dust found there displays a narrow isotopic field between  $0.70533 < {}^{87}Sr/{}^{86}Sr < 0.70819$  and  $-1.1 < \varepsilon_{Nd}(0) < 3.45$  for the bulk fraction and  $0.70807 < {}^{87}Sr/{}^{86}Sr < 0.70915$  and  $-0.94 < \varepsilon_{Nd}(0) < 0.86$  for the fine fraction. Due to Sr isotopic fractionation with particle size, the signature of the fine fraction reference adds to the PSA database for comparison to Antarctic ice core dust provenance studies.

491

492 3. Locally derived dust from McMurdo Sound is unlikely to be the major source of 493 DFe for seasonal phytoplankton blooms in the SW Ross Sea. Although, Sr and Nd 494 isotopic ratios of local dust on sea ice show similarities to lithogenic marine sediment, 495 we acknowledge the limited transport distance of coarse-sized dust in this region. As 496 dust transport varies from year to year, we cannot completely exclude the possibility 497 that local dust can contribute to DFe to the greater Ross Sea region although this is 498 not the dominant source of lithogenic Fe.

500 4. We surmise that there is significant remobilisation and upwelling of Fe from the 501 sea floor that contributes to Fe-fertilisation of phytoplankton during the austral 502 summer in the SW Ross Sea.

504 5. Source information of dust inputs to regions, such as the Ross Sea, improves the 505 ability to predict how such supply will change as the climate changes. As local 506 sources are important to the SW Ross Sea, this data could be included in models that 507 predict changes in snow and ice cover in the region.

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547

#### 548 Author Contributions

- 550 collected the samples; V.H.L.W, B.D and P.S.A prepared the samples and analysed
- 551 the data; V.H.L.W, G.B.D, C.B.A, B.D and P.S.A evaluated the data; all authors
- 552 contributed to the interpretation of the data and the writing of the manuscript.

24

553 Table 1: Nd and Sr concentrations and isotopic composition of McMurdo Sound and Chinstrap sediment trap samples analysed in this study.

	Size		Date	142	$a^{a)}\pm 2\sigma_{mean}$	b) (a)	a) a	C <sub>Nd</sub>	<sup>87</sup> Sr/ <sup>86</sup> Sr	<sup>d)</sup> ±2 <sub>smean</sub>	<sup>e)87</sup> Sr/ <sup>86</sup> Sr	<sup>f)</sup> ±2σ	Csr	
Sample	(µm)	Location	sampled	<sup>143</sup> Nd/ <sup>144</sup> Nd	*10 <sup>6</sup>	<sup>b)</sup> ε <sub>Nd</sub> (0)	<sup>e)</sup> ±2σ	(ppm)		*10 <sup>6</sup>	corrected	*106	(ppm)	Reference
McMurdo Sound														
MPR11-5	Bulk	77° 35.44 S	Nov 2011	0.512580	12	-1.1	0.2	110	0.708158	6.0	0.708186	8	140	This study
WII KIT 5		164° 31.36 E					0.2							
MPR11-5	<10	77° 35.44 S	Nov 2011	0.512590	7	-0.94	0.2	220	0.709119	5.3	0.709147	8	320	This study
		164° 31.36 E												This study
CRG7-5	Bulk	77° 05.44 S	Nov 2011	0.512633	11	-0.10	0.3	38	0.707607	4.2	0.707635	8	510	This study
		163° 41.86 E		0.012000			0.0							·
CRG7-5	<10	77° 05.44 S	Nov 2011	0.512632	512632 8	-0.12	0.3	68	0.708943	8.0	0.708971	8	570	This study
		163° 41.86 E	1107 2011	0.012002										
MPN10-5	Bulk	77° 24.52 S	Nov 2011	0.512735	6	1.9	0.4	40	0.706705	13	0.706733	13	6600	This study
		164° 18.60 E		0.512755	0	1.9	0.1					10		
MPN10-5	<10	77° 24.52 S	Nov 2011	0.512682 8	8	0.86	0.4	20	0.708188	6.4	0.708216	8	910	This study
		164° 18.60 E	1007 2011		č	0.00								This study
GH2	Bulk	76° 55.33 S	Nov 2009	0.512650	11	0.23	0.3	76	0.707299	12	0.707327	12	890	This study

		163° 6.17 E												
GH2	<10	76° 55. 33 S 163° 6.17 E	Nov 2009	0.512654	6	0.31	0.3	72	0.708045	5.8	0.708073	8	620	This study
GH9	Bulk	76° 58.36 S 162° 52.80 E	Nov 2009	0.512321	5	-6.18	0.3	37	0.712260	5.0	0.712288	8	290	(Winton et al., 2014)
MIS4	Bulk	77° 40.03 S 166° 35.97 E	Nov 2009	0.512815	4	3.45	0.3	56	0.705303	5.0	0.705331	8	800	(Winton et al., 2014)
MIS23	Bulk	77° 40.03 S 164° 35.79 E	Nov 2009	0.51276	6	2.38	0.3	51	0.705608	5.0	0.705636	8	520	(Winton et al., 2014)
Ross Sea Sediment														
CT1	Bulk	76° 20.5 S 165° 1.78 E	1997	0.512715	7	1.5	0.3	0.4	0.709008	5.1	0.709036	8	110	This study
CT1-leach	Bulk	76° 20.5 S 165° 1.78 E	1997	0.512836	8	3.9	0.3	0.5	0.704314	4.4	0.704342	8	20	This study

<sup>a)</sup>Internal precision, 2 standard errors of the mean. 

- 555 <sup>b)</sup>Nd isotopic ratios expressed as epsilon units  $\varepsilon_{Nd}(0) = [(^{143}Nd/^{144}Nd)_{sample}/(^{143}Nd/^{144}Nd)_{CHUR}-1]x10^4$ ; CHUR, chondritic uniform reservoir.
- <sup>556</sup> <sup>c)</sup>Uncertainty estimates based upon external precision for standard runs. Internal precision is used if it exceeds the external.
- <sup>d</sup>)Internal precision, 2 standard errors of the mean.
- <sup>e)</sup>Corrected to a NBS987 <sup>87</sup>Sr/<sup>86</sup>Sr ratio of 0.710245.
- <sup>f)</sup> Uncertainty estimates based upon external precision for standard runs. Internal precision is used if it exceeds the external.

#### 560 Figure captions

561

562 Fig. 1: a) Map of the SW Ross Sea showing the location of SW Ross Sea Chinstrap 563 sediment trap (CT1). b) Insert of McMurdo Sound within the SW Ross Sea showing 564 location of McMurdo Sound snow on sea ice samples (solid: this study, cross: Winton 565 et al. (2014) and shaded: exposed areas of unconsolidated sediment. Samples are named based on their location i.e. MP: Marble Point; CR: Cape Robert; GH: Granite 566 567 Harbour; MIS: McMurdo Sea Ice. EAIS: East Antarctic Ice Sheet, MDV: McMurdo 568 Dry Valleys, TAM: Transantarctic Mountains. Transect X-Y shown in red. c) Wind 569 roses illustrating the direction of storm events at Pegasus North and Marble Point 570 automatic weather stations (AWS). Locations of AWS shown in Fig. 1c). Modified 571 from Winton et al. (2014).

572

573 Fig. 2: Nd and Sr isotope signature of fine (black triangles) and bulk (white triangles) 574 McMurdo Sound dust, including bulk McMurdo Sound data (GH9, MIS4 and MIS23; Winton et al. (2014)) and leached and unleached Chinstrap sediment trap material. 575 576 Also plotted are data from Victoria Land potential dust sources that include different parent lithologies located in Fig. 1. (Delmonte et al., 2013; Delmonte et al., 2010; 577 578 Delmonte et al., 2004) and the isotopic composition of Ross Sea seawater (Basak et 579 al., 2015; Elderfield, 1986). Insert top right: McMurdo Sound dust highlighting the 580 fractionation between fine and coarse particle sizes and a hypothetical mixing line 581 between the two end members MVG and Southern Victoria Land, TAM.

582

Fig. 3: a) Extrapolation of the annual DFe flux from McMurdo Sound into the SW
Ross Sea. DFe data sourced from: Winton et al. (2014). The predicted dust flux at the

Chinstrap site is estimated at ~0.01 g m  $^{-2}$  yr  $^{-1}$  with a corresponding DFe flux of ~2 585 nmol  $m^{-2} d^{-1}$ . b) Also shown is the rate of primary production with distance from 586 McMurdo Sound. As the dust flux exponentially decreases, the rate of primary 587 588 production increases. Primary production inferred from the annual mean chlorophylla concentration (2000-2009) in the McMurdo Sound polynya (72.0 °S - 78.083 °S, 589 590 160.916 °E 179.040 °W) from SeaWiFS satellite \_ data 591 (http://giovanni.gsfc.nasa.gov).

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