Authigenic ferrimagnetic iron sulfide preservation due to non-steady state diagenesis: A perspective from Perseverance Drift, Northwestern Weddell Sea

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Abstract

We document magnetic mineral diagenesis with high resolution magnetic susceptibility, hysteresis, isothermal remanent magnetization, and other rock magnetic measurements through a shallow sulfate-methane transition (SMT) at Perseverance Drift-a high-accumulation rate Holocene biosiliceous Antarctic marine sediment deposit. The structure of the SMT is defined with porewater measurements from the same core, allowing direct comparison. Dissolution of the detrital (titano)magnetite assemblage, with preferential dissolution of stochiometric magnetite, occurs in the upper SMT. Higher coercivity magnetic minerals dissolve more slowly, continuing to dissolve through the entire SMT and could be a source of ferric iron for microbial respiration following exhaustion of porewater sulfate, as suggested by accumulation of porewater ferrous iron below the SMT. Superparamagnetic ferrimagnetic mineral enrichment/depletion occurs in three phases through the SMT and is coupled tightly to the availability of dissolved ferrous iron relative to dissolved sulfide. High concentrations of authigenic remanence-bearing iron sulfides, including greigite and hexagonal 3C pyrrhotite, which can be detected using remanence parameters but not in-field concentration dependent parameters, accumulate in a transient horizon at the base of the SMT during this early diagenesis, where sulfide is present but limited relative to dissolved ferrous iron. Formation of this remanence-bearing iron sulfide horizon is likely facilitated by continued iron reduction through the SMT. Non-steady state perturbations that shift the porewater profile, such as changes in carbon flux or sedimentation rate, can lead to preservation of these transient horizons, much like well documented preservation of manganese oxide layers in marine sediments following similar shifts to porewater profiles.

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32

33 Key Points

Magnetic mineral diagenesis is studied in Antarctic sediments using high resolution porewater
 and rock magnetic data

- Magnetic mineral dissolution and authigenesis is tightly coupled to dissolved sulfide and ferrous
- 37 iron concentrations
- Remanence-bearing iron sulfides form below the sulfate-methane transition and can be
 preserved following non-steady state diagenesis
- 40

41 1. Introduction

42 Primary sedimentary magnetic mineral assemblages reflect sediment provenance and 43 depositional processes; however, following deposition these magnetic mineral assemblages can alter by 44 dissolution of components of the primary detrital assemblage or through growth of new authigenic 45 magnetic minerals (Liu et al., 2012; Roberts, 2015). Understanding magnetic mineral diagenesis is 46 important for interpreting paleomagnetic and environmental magnetic records. Fundamentally, 47 magnetic mineral diagenesis is driven by changes in pore water geochemistry (e.g. Canfield and Berner, 48 1987; Dillon and Bleil, 2006; Garming et al., 2005; Karlin and Levi, 1983; Kars et al., 2018; Larrasoaña et 49 al., 2007; Leslie et al., 1990), which reflects microbial activity, carbon (organic or methane) flux, oxygen 50 (and other electron acceptors) availability, and/or sedimentation rate.

51 Post-depositional alteration of magnetic mineral assemblages may complicate paleomagnetic 52 data, such as by smoothing or overprinting the signal through acquisition of a chemical remanent 53 magnetization (CRM) at depth (e.g. Jiang et al., 2001; Rowan and Roberts, 2006). This can introduce 54 significant uncertainty to magnetostratigraphic studies as the timing and depth ranges the authigenic 55 minerals carrying the CRM form is poorly constrained in most systems (Roberts et al., 2018b). However, 56 for environmental magnetic studies, magnetic mineral diagenesis is tightly coupled to iron and sulfur 57 cycling and can preserve signals of the biogeochemical history of these systems (e.g., Kars and Kodama, 58 2015; Tarduno, 1994). A link to paleoenviromental conditions that could drive non-steady state changes 59 in porewater conditions can often be made using the degree of alteration tracked by magnetic 60 measurements, with diverse applications including sea-ice conditions in the Arctic Ocean (Brachfeld et 61 al., 2009), abrupt changes in Asian monsoon intensity (Chang et al., 2016a), discharge events in the 62 Santa Barbara Basin (Blanchet et al., 2009), bottom water ventilation and export productivity in the 63 eastern Mediterranean (Larrasoaña et al., 2003) and others.

64 To move toward a more quantitative understanding of uncertainties in paleomagnetic data and 65 how magnetic minerals can be used to understand diagenetic histories and their relationship with 66 paleoenvironmental changes requires further porewater and magnetic observations from a wide range 67 of depositional environments. Here we document how the magnetic properties of rapidly accumulating 68 biosiliceous hemipelagic Antarctic sediments evolve in relation to sediment porewater geochemistry. 69 The sediments studied in detail here were deposited during the latest Holocene (~800 cal yrs B.P. to 70 present, Kyrmanidou et al., 2018) which limits the recording of environmental (and related depositional) 71 variability associated with changes in Earth's orbit that often influence other records of paired 72 porewater and magnetic variations (e.g. Dillon and Bleil, 2006; Garming et al., 2005).

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75 2. Geologic Background

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77 2.1 Perseverance Drift, Northeastern Antarctic Peninsula

78 Perseverance Drift is located north of Joinville Island on the continental shelf, Northeastern 79 Antarctic Peninsula/Northwestern Weddell Sea, in the Joinville-D' Urville Trough (Figure 1). The deposit 80 was first surveyed in 2004 by the R/V Laurence M. Gould (cruise LMG0404) to investigate if sediment 81 drifts form when tidal/geostrophic currents slow at channel openings on the Antarctic shelf (Domack 82 and the LMG0404 Scientific Party, 2004). Two Kasten cores (1.5 to 6 m, large volume, and square cross 83 section gravity corers that are designed to preserve the sediment water interface; Kuehl et al., 1985), 84 Cores KC3 and KC16, were recovered during this expedition (Darley, 2014). In 2012 as part of the Larsen 85 Ice Shelf System Antarctica (LARISSA; Wellner et al., 2019) program, the R/V Nathaniel B. Palmer (cruise 86 NBP1203) revisited the drift and conducted a more extensive (although incomplete due to weather and 87 time limitations) multi-beam and sub-bottom acoustic survey, which revealed a complex seafloor with 88 sediment thicknesses of at least 90 m (Kyrmanidou et al., 2018; Vernet and the NBP1203 Scientific Party, 89 2012). A Jumbo Kasten Core (JKC36; 529 cm; 63.089 °S, 55.395 °W; 806 m water depth) and Jumbo Piston Core (JPC36; 2399 cm; 63.089 °S, 55.399 °W; 806 m water depth) were recovered during this 90 91 expedition. The NBP1203 geophysical survey indicates that while Cores KC3, JKC36, and JPC36 were 92 taken from the drift, Core KC16 was recovered from a nearby sedimentary basin separate from the drift 93 (Vernet and the NBP1203 Scientific Party, 2012).

94 Ice flowed over the Joinville-D' Urville Trough during the Last Glacial Maximum (Lavoie et al., 95 2015) and following deglaciation sediments accumulated at high rates, averaging ~760 cm/kyr for the 96 last ~3.3 ka at the Core JKC36/JPC36 location, and provides an ultra-high resolution archive of Holocene 97 paleoenvironmental and palaeoceanographic conditions with abundant carbonate shells for establishing 98 a geochronology (Kyrmanidou et al., 2018). For the last three millennia, diatom assemblages reflect high 99 primary productivity, mostly without nutrient limitation, and benthic foraminifera assemblages are 100 consistent with high primary productivity with variable sea ice cover (Kyrmanidou et al., 2018). Benthic 101 foraminifera assemblages suggest a paleoceanographic change at ~1.8 ka to colder conditions with more 102 sea ice, which is interpreted as the onset of Neoglacial conditions for the Northeasternmost Antarctic 103 Peninsula (Kyrmanidou et al., 2018).

104 Perseverance Drift sediments also contain the authigenic mineral ikaite (CaCO₃ \bullet 6H₂O), which 105 becomes glendonite after loss of hydration and is often used as a paleoenvironmental indicator (e.g., 106 Spielhagen and Tripati, 2009). Ikaite is found in some organic-rich biosiliceous sediments around the 107 North/Northeastern Antarctic Peninsula at sites in the path of water masses exported from the Weddell 108 Sea, including: the Bransfield Basin, Firth of Tay, and Vega Drift; but notably not the Palmer Deep nor 109 western Antarctic Peninsula inner shelf deposits (Domack et al., 2007; Lu et al., 2012; Suess et al., 1982; 110 Zhou et al., 2015). Based on modern observations, ikaite formation is associated with cold (≤ 10 °C) 111 bottom waters and high porewater phosphate concentrations (Zhou et al., 2015). The latter may be 112 related to organic matter remineralization, or, as is hypothesized for other Northeastern Antarctic 113 Peninsula sites, may be released during Fe-bearing mineral dissolution (Zhou et al., 2015).

2.2 Magnetic Mineral Diagenesis at the Sulfate Methane Transition 115 116 Significant magnetic mineral dissolution can occur at shallow depths in marine sediments at the 117 sulfate-methane transition (SMT) where downward diffusion of seawater sulfate (SO₄²⁻) is used as an 118 electron acceptor to fuel anaerobic oxidation of methane (AOM and CH_{4}) (Knittel and Boetius, 2009) 119 (Equation 1). This reaction produces elevated hydrogen sulfide (HS⁻) concentrations when dissolved 120 ferrous iron (Fe²⁺) is limited, and leads to the dissolution of iron minerals like (titano-) magnetite (Fe₂₋ 121 $_x$ Ti_xO₄, which is a possible reservoir for ferric iron (Fe³⁺) as long as x < 1) after exhaustion of more reactive 122 species of iron (Canfield, 1989; Canfield and Berner, 1987) (Equation 2). 123 $CH_4 + SO_4^{2-} \rightarrow HCO_3^{-} + HS^{-} + H_2O[1]$ 124 $Fe^{2+} + HS^{-} \rightarrow FeS + H^{+}[2]$ 125 126 127 In anoxic sediments with a SMT, AOM may be the principal driver of the sulfate⁻ profile, but 128 sulfate reduction also occurs following the use of oxygen, nitrate, manganese, and iron as electron 129 acceptors in the oxidation of organic carbon (Froelich et al., 1979) (Equation 3). 130 131 Organic Matter + SO_4^{2-} \rightarrow CO_2 + NH_3 + HS^- + H_3PO_4 + H_2O [3] 132 Generally, detrital magnetic mineral dissolution can be identified by a large decrease in 133 134 concentration dependent magnetic parameters (referred to here as a dissolution front) and a coarsening 135 of bulk magnetic 'grain-size' parameters, due to the preferential dissolution of finer magnetite (e.g. 136 Karlin and Levi, 1983; Rowan et al., 2009). Additionally, following complete reduction of SO_4^2 and 137 reaction of the product, HS⁻, with iron to form paramagnetic pyrite (FeS₂) (Berner, 1984, 1970), HS⁻ could 138 become limited with respect to Fe²⁺ in systems where there is continued dissolution of iron-bearing 139 minerals. This geochemical setting is hypothesized to provide conditions suitable to preserve 140 intermediaries along the pyrite formation pathway, including the ferrimagnetic mineral greigite (Fe₃S₄) 141 (Blanchet et al., 2009; Hunger and Benning, 2007; Kao et al., 2004; Roberts et al., 2011; Rowan et al., 142 2009) and/or hexagonal 3C pyrrhotite. Both minerals have distinct room temperature magnetic 143 remanence properties, despite early assumptions of the latter only having antiferromagnetic properties 144 (Horng, 2018; Horng and Roberts, 2018). The depths at which these magnetic iron sulfides form relative 145 to the SMT is still poorly understood (Roberts et al., 2018b). 146 Methods and Materials

147 148

149 3.1 Core NBP1203 JKC36

150 Core JKC36 was collected from Perseverance Drift using a 6 m Jumbo Kasten Corer. Upon 151 recovery, Core JKC36 smelled strongly of H_2S and degassed CH_4 , suggesting a shallow SMT that 152 motivated this study and careful sampling by the NBP1203 shipboard geomicrobiology group.

- 153 Environmental magnetic samples were taken on ship soon after collection of pore water samples and
- 154 before the sediments altered significantly from warmer temperatures and oxygen exposure.

155 Core JKC36 consists of diatomaceous mud containing mm-scale laminations, streaks, and

156 patches of dark organic materials from 0 – 350 cm below sea floor (cmbsf). Below 350 cmbsf the

sediment becomes increasingly mottled, with laminations observed at a few intervals, and increased

158 evidence for bioturbation. Beyond laminations and evidence for bioturbation, the lithology is fairly

consistent, excluding an interval from 193 – 203 cmbsf described as a lighter grayish olive diatomaceous
 mud with laminae and 205 cmbsf as a moderate olive brown silty mud with sharp contacts. Calcareous

161 shells and shell fragments are abundant but are only documented in the uppermost 50 cm of the core

162 and below 300 cmbsf. Macroscopic authigenic ikaite crystals were found at 303-305 cmbsf, 463-465

163 cmbsf, 479-481 cmbsf, and 505 cmbsf (Figure 2). ²¹⁰Pb dating indicates that the Kasten Corer recovered

164 an intact sediment-water interface, with little, if any, sediment lost (Kyrmanidou et al., 2018).

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166 3.2 Sediment Magnetism

167 While on ship, 1 cm horizon samples were taken from JKC36 at 1 cm intervals from 0 to 20 168 cmbsf and at 2 cm intervals from 20 to 519 cmbsf and kept cool by refrigeration or ice packs until the 169 samples could be freeze-dried at Montclair State University. 150-300 mg of the sediment were packed 170 into gelatin capsules. Low field, mass normalized magnetic susceptibility (χ_{if}) was measured three times 171 and averaged using an AGICO KLY-4 Kappabridge at an applied field of 300 A/m. Thermomagnetic curves 172 $\chi(T)$ were also measured for bulk sediment samples with the Kappabridge in an argon atmosphere, 173 heating from room temperature at a rate of 0.2 °C/s to ~700 °C and then cooling to ~40 °C. Select 174 intervals were measured again in ambient air without an argon atmosphere, to compare magnetic 175 mineral alteration at high temperatures. The argon atmosphere thermomagnetic curves were unmixed 176 using the method of Reilly et al. (2016), which adapts the method of Heslop and Roberts (2012) and uses 177 the Simplex Identification via Split Augmented Lagrangian (SISAL) method of Bioucas-Dias (2009). The 178 method was modified to unmix the measured curves rather than the derivative of that curve due to the 179 noisier nature of these weakly magnetic sediments and includes both heating and cooling curves. End 180 members were determined from 100 iterations with each iteration using 16 of the 18 curves to quantify 181 uncertainty associated with any one sample having too large an influence on the analysis.

182 Magnetic hysteresis parameters were measured using a Princeton Measurements Corporation 183 MicroMag 3900 vibrating sample magnetometer (VSM) with a peak 1 T field (H). The raw hysteresis 184 loops were mass-normalized and a high-field slope correction was calculated between 0.7 and 1 T. 185 Measurement averaging times were chosen based on sample raw magnetization (M) in a 1 T field, 186 ranging from 0.25 to 1 s. The bulk coercivity (H_r) , saturation remanent magnetization (M_r) , saturation 187 magnetization (M_s), and mass normalized high field susceptibility (χ_{hf}) were determined from these 188 loops. The VSM was also used to measure DC demagnetization of remanence curves, using an initial 189 saturation field of 1 T and an increasing incremental back-field H in -10 mT steps, with the coercivity of 190 remanence (H_{cr}) determined from where M=0 on the curve. S-ratios (Stober and Thompson, 1979) and 191 HIRMs (Stoner et al., 1996) were calculated by measuring isothermal remanent magnetizations (IRM) on 192 the VSM after applying a 1 T field and then a backfield of -0.3 T, with S-ratio = -IRM_{-0.3T}/IRM_{1T} and HIRM = 193 $(IRM_{1T} + IRM_{.0.3T})/2$. First order reversal curves (FORCs; n = 140 per sample; Pike et al., 1999) were 194 measured on the VSM for selected samples using a 0.5 T saturating field, 0.25 s averaging time, H_{b} range 195 of -30 mT to 30 mT, and H_c range of 0 to 100 mT. FORCs were processed using FORCinel v. 3.03 software 196 (Harrison et al., 2018; Harrison and Feinberg, 2008) and VariFORC smoothing (Egli, 2013).

197 IRM acquisition was measured for selected gelatin capsules following demagnetization with a
 200 mT peak alternating field (AF) using an ASC Scientific D-2000 AF demagnetizer. Stepwise IRM was
 imparted using an ASC Scientific IM-10-30 Impulse Magnetizer and measured on an AGICO JR-6 spinner
 magnetometer at 50 mT intervals from 0 to 0.3 T, 100 mT intervals from 0.3-1 T, and 1 T intervals from
 201 1-5 T.

202 Low temperature measurements were made with a Quantum Design Magnetic Properties 203 Measurement System (MPMS) at the Institute for Rock Magnetism, University of Minnesota. Field 204 Cooled (FC) remanence measurements were made by cooling the sample to 20 K in a 2.5 T field and 205 measuring the magnetization at ~5 K intervals as the sample warmed to 300 K. Zero Field Cooled (ZFC) 206 remanence measurements were made with the same protocol, except the sample was cooled to 20 K in 207 zero applied field. MPMS magnetic susceptibility ($\chi_{If-MPMS}$) was measured as a function of temperature 208 and field frequency at ~10° increments during warming from 20 K to 300 K, with an applied field of 240 209 Am⁻¹ at 5 frequencies (1, 5.6, 31.6, 177.6, and 997.3 Hz). Frequency dependent magnetic susceptibility 210 (χ_{FD}) , reported as a percent, was calculated as the difference between $\chi_{If-MPMS}$ at 31.6 and 1 Hz,

211 normalized by the $\chi_{If-MPMS}$ at 1 Hz.

To complement the magnetic analyses, heavy mineral extracts were prepared from bulk freezedried sediment at 0, 148, 206, and 450 cmbsf through centrifuging in a sodium polytungstate solution with a density of 2.88 g/cm³ and freezing with dry ice (Skipp and Brownfield, 1993). Heavy mineral extracts were mounted on carbon tape and were carbon coated for imaging using a Hitachi S-3400N scanning electron microscope (SEM) and for qualitative chemical analysis on a Bruker X-flash energy dispersive spectrometer (EDS), using a 15 keV accelerating voltage.

To compare sediment magnetic properties with those of potential source rocks, 23 bedrock samples were chosen from the Polar Rock Repository (PRR) at the Byrd Polar Research Center, Ohio State University, that were proximal to our study area (Joinville Island and nearby smaller islands) and which attempt to represent regional lithologies, including sedimentary, volcanic, and intrusive igneous samples. Rock chips (~50-150 mg) were immobilized with quartz Fiberfrax[®] inside gelatin capsules and were measured for χ_{if} and hysteresis parameters in the same fashion as for the Core JKC36 sediments. All PRR sample high temperature χ_{if} measurements were made in an argon atmosphere.

225

226 3.3 Porewater Geochemistry

227 Subcores for porewater geochemical analysis were collected from target depths immediately 228 after removing each Jumbo Kasten Core panel. Duplicate sediment samples for CH₄ analysis (1 mL each) 229 were collected with sterile 3 mL pre-cut open-end syringes and transferred to N₂ purged 10 mL gas 230 chromatography (GC) vials pre-filled with 2 mL of 1 M NaOH then capped and sealed with teflon-coated 231 butyl rubber septa. Vials were equilibrated at 30 °C for 30 minutes then analyzed by manual headspace 232 sampling and injection on a shipboard Agilent 6890N gas chromatograph using a Plot-Q column and 233 flame ionization detection (FID). Final concentrations were corrected for porewater content (mL $H_2O/$ 234 mL sediment).

All other samples for porewater analysis were collected in sterile 60 mL pre-cut open-end syringes. Immediately after recovery, syringes were wrapped in parafilm and vinyl tape and were transferred to a N₂ purged glove box maintained at 1 °C for subsampling. All remaining manipulations

- 238 were conducted in a glove box. Sediment was transferred to gas-tight acid-washed polycarbonate Oak
- 239 Ridge tubes and centrifuged at 10,000 RCF for 20 min at 4 °C. The supernatant was collected and filtered
- $240 \qquad through a 0.22 \ \mu m \ polyether sulfone \ syring e \ filter \ and \ aliquoted \ for \ sulfide, \ sulfate \ and \ ferrous \ iron$
- 241 assays. Sulfide samples were preserved by equal volume addition to anoxic 20% Zn acetate solution and
- 242 later quantified colorimetrically by the Cline method (Cline, 1969). Samples for sulfate and other anions
- 243 were stored at 4 °C in the dark until analyzed by ion chromatography on a Metrohm 861 Advanced
- 244 Compact IC unit using a Metrosep A SUPP 5-250/4.0 anion column. Samples for Fe²⁺ (aq) were preserved
- 245 by equal volume dilution with 0.1 M HCl and later quantified colorimetrically by the Ferrozine assay
- 246 (Stookey, 1970).
- 247

248 4. Results

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250 4.1 Porewater Geochemistry

251 Core JKC36 has a shallow SMT centered around 70 cmbsf, with $[SO_4^{2-}]$ declining in concentration 252 from ~23-27 mM in the upper 20 cm to background levels at 80 cmbsf. Sulfide, [S²], is present above 253 background levels (> .5 mM) between 10 and 225 cmbsf with peak concentration of 6 mM. Methane, 254 CH₄, is present above background levels below 60 cmbsf (Figure 2). As Core JKC36 degassed methane 255 upon recovery, CH₄ levels above the solubility limit at 1 atmosphere and 0 °C (~ 2.3 mM) are highly 256 variable and reflect this degassing (Yamamoto et al., 1976). [Fe²⁺] is present in high concentrations (~6 257 μ M) at the core top but decreases in the upper 10 cm to background levels between 10 and 80 cmbsf, 258 where sulfate is actively reduced and sulfide accumulates in the porewater. Between 80 and 180 cmbsf, $[Fe^{2+}]$ is elevated, despite the presence of $[S^{2-}]$ in the porewater, with peak concentrations (~3 μ M) at 259 260 140 cmbsf. Below 180 cmbsf, [Fe²⁺] returns to background levels.

AOM coupled to microbial sulfate reduction is likely occurring between 60 and 80 cmbsf, where [SO₄²⁻] and CH₄ are both present in the porewater (**Figure 2**). Accumulation of [Fe²⁺] directly below the exhaustion of sulfate, suggests that iron reduction is also occurring within the SMT. This iron reduction must be occurring at a rate where [Fe²⁺] is being produced faster than it can react with [S²⁻]. Iron reduction at and below the SMT could be explained by iron dependent AOM, which uses ferric iron instead of sulfate as an electron acceptor (Beal et al., 2009; Riedinger et al., 2014).

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268 4.2 Sediment Magnetism

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270 4.2.1 Concentration Dependent Magnetic Parameters

The χ_{if} profile for Core JKC36 is marked by a steady decline in values (~10 to 7 x 10⁻⁷ m³/kg) in the upper 30 cm and a sharp drop-off immediately below that, consistent with dissolution fronts described in sulfate reducing sediments (e.g. Karlin and Levi, 1983; Rowan et al., 2009) (**Figure 3**). This dissolution front is also observed in the concentration dependent parameter M_s, which tracks the total in-field ferrimagnetic saturation magnetic moment, and M_r, which tracks the total remanent ferrimagnetic

- 276 moment. The dissolution front observed in these three parameters occurs in the upper portion of the
- 277 SMT when [S²⁻] is only about one-third of the highest measured concentration (~2.3 mM). Below the

278 dissolution front, χ_{if} , M_s , and M_r have high frequency variability (on the order of 10-20 cm), with a shift 279 from relatively lower mean values (χ_{if} = 4.1 x 10⁻⁷ m³/kg) to higher mean values (χ_{if} = 5.0 x 10⁻⁷ m³/kg) 280 around 330 cmbsf.

281 The χ_{bf} profile tracks the non-ferrimagnetic contributions to magnetic susceptibility and, under 282 ideal circumstances, reflects the relative proportion of paramagnetic minerals (often iron-bearing) to 283 diamagnetic minerals. In reality, $\chi_{\rm bf}$ must be interpreted with caution because if high-coercivity minerals 284 are present, like hematite and goethite, which saturate in fields above 700 mT, their induced 285 magnetization can influence the slope of the magnetization as a function of field strength and therefore the calculation of χ_{hf} (Brachfeld, 2006). With that in mind, χ_{hf} decreases in the upper 30 cm along with χ_{hf} 286 287 M_{s} , and M_{r} , which is consistent with steady dissolution of iron-bearing paramagnetic minerals along with 288 ferrimagnetic minerals above the dissolution front (Figure 3). However, there is no sharp dissolution 289 front in the χ_{hf} profile like that observed in the χ_{hf} , M_s , and M_r profiles. Also, unlike χ_{hf} , M_s , and M_r , the χ_{hf} 290 values increase after a minimum around 48 cmbsf and reach a local maximum around 98 cmbsf, where 291 [S²⁻] also reaches its highest measured value. This is consistent with the formation of new authigenic 292 paramagnetic minerals—likely pyrite. Below this horizon, χ_{hf} undergoes high frequency variation without 293 the increase around 330 cmbsf, as was observed in χ_{lf} , M_s , and M_r . There is a peak in χ_{hf} at 206 cm; 294 however it is difficult to know if this is related to a higher concentration of paramagnetic minerals or an 295 artifact of the higher concentration of high-coercivity minerals at the same horizon, discussed below.

296 HIRM values, which track the concentration of minerals that contribute to magnetic remanence 297 with coercivities greater than 300 mT, are about two orders of magnitude lower than M_r values, which 298 track the total magnetic remanence of the sediments (median values are 4.2 x 10⁻⁵ versus 2.5 x 10⁻³ Am²/ 299 kg, respectively; Figure 3). This suggests that to first order, X_{if} , M_s , M_r , largely reflect lower coercivity 300 ferrimagnetic mineral concentrations. HIRM values do not decrease significantly in the upper 30 cm like 301 the other concentration dependent parameters. However, HIRM values decline steadily below the χ_{if} , 302 M_s, and M_r dissolution front to a local minimum around 174 cmbsf, near the base of the SMT, which 303 suggests continued steady dissolution of minerals with coercivities greater than 300 mT through the entire interval of elevated [S²⁻], including following the exhaustion of $[SO_4^{2-}]$ around 80 cmbsf and 304 305 accumulation of $[Fe^{2+}]$ below that (Figure 2). This is consistent with other diagenetic environments, 306 where high coercivity minerals like hematite that may be a minor contributor to the magnetization of 307 the primary detrital assemblage dissolve more slowly than (titano)magnetite (Dillon and Bleil, 2006; Garming et al., 2005; Liu et al., 2004; Yamazaki et al., 2003) and implies that these high-coercivity 308 309 minerals are likely a source of ferric iron within the SMT for iron reduction.

- Starting at the base of the SMT, HIRM values (> $\sim 1 \times 10^{-4} \text{ Am}^2/\text{kg}$) indicate horizons with enrichments of magnetic minerals with coercivity greater than 300 mT, particularly at 206 cmbsf and
- 312 442 cmbsf (up to 5.5 and 1.2 x 10^{-4} Am²/kg, respectively; Figure 3). These horizons have sharp bases and
- 313 grade upward to lower HIRM values. The 206 cmbsf horizon reaches HIRM values an order of magnitude
- greater than underlying or overlying sediments and is accompanied by a spike in M_r values (up to 9.6 x
- 315 10^{-3} Am²/kg), but it is not detected in χ_{lf} or M_s. This suggests that the enrichment of this high coercivity
- 316 mineral contributes significantly to magnetic remanence but does not reach high enough concentrations
- 317 to be detected using in-field concentration dependent measurements in the presence of the background
- 318 ferrimagnetic mineral assemblage.

320 4.2.2 High Temperature Magnetic Susceptibility

321 $\chi_{\rm lf}$ as a function of heating to ~700 °C and cooling back to room temperature has variable 322 behavior with the most pronounced difference being that cooling curves are always weaker than or 323 about equal to heating curves from 0-30 cmbsf (above the dissolution front). Cooling curves are 324 generally stronger than heating curves below the dissolution front (Figure 4a). Principal component 325 analysis of the $\chi_{\rm f}$ temperature curves indicates that the first and second components explain about 86% 326 and 7% of the variance, respectively, with significantly less variance explained by subsequent 327 components. Accordingly, we adopt a 3 end-member solution using a regularization parameter (τ) of 328 0.7848, which is the value that uses that smallest simplex that still has a good fit to the data (Figure 4e) 329 (see Reilly et al. (2016) for discussion).

330 The composition of the surficial sediments (0 cmbsf) is almost entirely composed of end-331 member 1 (EM1) and EM1 is likely representative of the primary detrital magnetic mineral assemblage. 332 It is the only end-member with a cooling curve weaker than the heating curve, which suggests the 333 presence of maghemite, which inverts to hematite at high temperatures (Gehring et al., 2009; Özdemir 334 and Banerjee, 1984) (Figure 4d). χ_{if} increases with initial heating starting with a relatively slow increase 335 at around 150 °C and increasing to a relatively faster rate around 230 °C. Peak $\chi_{\rm ff}$ is achieved at around 336 300 °C, after which it steadily decreases until about 557 °C, where it begins a sharp drop centered at 337 around 580 °C. This is the only end-member with a pronounced decline at about 580 °C, which suggests 338 that EM1 is the only end-member with a significant proportion of stochiometric magnetite. The cooling 339 curve has about the same slope as the heating curve while cooling from 550 and 350 °C, which suggests 340 that the gradual χ_{μ} decline in this temperature range is related to the presence of titanomagnetite of 341 variable titanium concentrations (Lattard et al., 2006).

342 End-members 2 (EM2) and 3 (EM3) have similar heating curves, with only a slight χ_{f} increase 343 between 200 and 300 °C and a steady decline between about 350 and 600 °C with no sharp drop-offs 344 (Figure 4d). Cooling curves generally retrace the heating curve during cooling to about 500 °C and then 345 have stronger χ_{if} than the heating curve while cooling to room temperature. The major difference 346 between EM2 and EM3 is that EM3 has a sharp χ_{f} increase on cooling centered around 265 °C with a 347 maximum $\chi_{\rm f}$ at about 233 °C and decrease in $\chi_{\rm f}$ as it continues to cool, suggesting alteration in EM3 348 samples that does not occur in EM2 samples. This could be indicative of the creation of an 349 antiferromagnetic hexagonal pyrrhotite polytype during heating, one that is only ferrimagnetic between 350 its λ transition ~200-220 °C and its Curie temperature of ~275-295 °C. We note this polytype is different 351 from the hexagonal 3C pyrrhotite with room temperature ferrimagnetic properties (Horng, 2018) 352 discussed elsewhere in this study. This interpretation is consistent with previously published reports of 353 pyrrhotite polytypes produced during the alteration of greigite and other iron sulfides at high 354 temperatures or under oxic conditions (Chang et al., 2008; Roberts et al., 2011; Schwarz and Vaughan, 355 1972; Sweeney and Kaplan, 1973).

356 Downcore variability in the end-member abundances suggest that the EM1 primary 357 ferrimagnetic mineral assemblage (or as close to the primary assemblage as the 0 cmbsf sample 358 captures), contains stochiometric magnetite, maghemite, and titanomagnetite with a range of 359 compositions. EM1 steadily decreases in its relative abundance from 0 to 30 cmbsf, above the 360 dissolution front, along with the steady decrease in room temperature χ_{lf} , which suggests that 361 dissolution through this interval is preferentially removing stochiometric magnetite and maghemite 362 relative to titanomagnetite (Figure 4b-c). A preference for dissolution of stochiometric magnetite over

- 363 titanomagnetite has been observed elsewhere in diagenetic sediments, likely because Ti⁴⁺ substitution
- reduces the ratio of Fe^{3+} to Fe^{2+} in the mineral (Dillon and Bleil, 2006). There is no clear trend in end-
- 365 member abundance below the dissolution front nor in end-member abundances where there is elevated
- 366 pore water [S²⁻], but the ferrimagnetic assemblages are dominated by titanomagnetite with or without
- 367 mineral phases that alter to an antiferromagnetic hexagonal pyrrhotite polytype. Enrichment of the EM3
- 368 assemblage that contains the mineral phases that alter to an antiferromagnetic hexagonal pyrrhotite
- 369 polytype often coincides with HIRM peaks (Figure 3).
- 370

371 4.2.3 Concentration Independent Magnetic Parameters

- 372 Concentration independent magnetic parameters can be influenced by relative changes in 373 magnetic coercivity and magnetic mineralogy. Hysteresis loop ratios M_r/M_s and H_{rr}/H_c are often 374 interpreted as reflecting changes in magnetic grain-size or domain state (e.g. Day et al., 1977); however, 375 they can also be influenced by variations in relative proportions of magnetic mineralogy, surface 376 oxidation, and numerous other factors (e.g. Roberts et al., 2018a). As we have already documented 377 differences in relative magnetic mineralogical abundances during diagenesis in the SMT (Figures 3 and 378 4), we keep this in mind in our discussion of concentration independent magnetic parameter. For 379 discussion, we divide the core into five zones based on magnetic mineral parameters (particularly M,
- 380 and H_{cr}/H_c) and porewater geochemistry variations.

381 Zone 1 (0-33 cmbsf) represents the interval from the top of the core to the base of the 382 dissolution front, during the initial downcore $[S^2]$ rise in the upper portion of the SMT (Figure 5a). The 383 change in magnetic mineral concentration in this interval, tracked by M_s, has a strong correlation with M_r/M_s (r = 0.83), H_{rr}/H_c (r = -0.94), and S-ratio (r = 0.95) but a weak correlation with H_{rr} (r = 0.34) (Figures 384 385 5). This reflects 'coarsening' of the magnetic mineral assemblage due to preferential dissolution of fine magnetic minerals. The absence of a trend in the H_{cr} values and significant correlation with S-ratio, also 386 reflects mineralogic changes. The lack of H_{cr} trend above the dissolution front could reflect the 387 388 competing influence of dissolution of stable single domain magnetite, lowering the coercivity of 389 remanence-bearing minerals and preferential preservation of titanomagnetite (Figure 4c), and slower 390 dissolution of high coercivity minerals (Figure 3), which increases the bulk coercivity of remanence.

391 Zones 2a (33-87 cmbsf) and 2b (87-131 cmbsf) represent the interval of peak [S²⁻]. Zone 2a has 392 increasing sulfide downcore, decreasing sulfate, and low [Fe2+], which indicate that sulfate reduction is 393 producing more [S²⁻] than can react with [Fe²⁺]. Zone 2b has decreasing [S²⁻] downcore, depleted [SO₄²⁻], 394 and elevated [Fe²⁺], which indicate that there is enough [Fe²⁺] to react with the [S²⁻] (Figure 2). While M_{r} / 395 M_s and H_{cr} values are comparable for both Zones 2a and 2b, H_{cr}/H_c values are distinct—driven by H_c 396 variability (means and ranges for these parameters are shown in Figure 5b). H_{cr} is a remanence 397 measurement and H_c is an in-field measurement, so the variability in H_c and lack of variability in H_c is 398 likely driven by the concentration of superparamagnetic (SP) minerals that lower the bulk coercivity in 399 the presence of a magnetic field with evidence for SP enrichment in Zone 2a but not 2b. This 400 observation suggests that the concentration of SP minerals in the SMT is strongly related to the 401 availability of [S²⁻] relative to [Fe²⁺].

402 The relationship between magnetic mineral concentration, tracked by M_s , and concentration 403 independent parameters changes in Zone 2a and 2b relative to Zone 1, with M_r/M_s (r = -0.57 and -0.78, 404 respectively) and H_{cr} (r = -0.75 and -0.78) having strong negative correlations, and S-ratio and H_{cr}/H_{c} 405 having much weaker correlations than Zone 1. Thus, rather than a coercivity 'coarsening' with decreases 406 in ferrimagnetic concentration as is observed above the dissolution front in Zone 1, below the 407 dissolution front in Zones 2a and 2b there is a coercivity 'fining' of the ferrimagnetic particles with

408 decreases in ferrimagnetic concentration.

409 Zones 3a (131-329 cmbsf) and 3b (329-519 cmbsf) represent the interval at the base of the SMT 410 and the underlying methanic zone. In general, M_r/M_s , H_{cr}/H_c , and H_{cr} values are like those found in Zone 411 2a, which suggests an altered magnetic mineral assemblage, like that found beneath the dissolution 412 front, and variable SP concentrations (Figure 5). Horizons previously discussed at 206 and 442 cmbsf 413 with high HIRM values (Figure 3) are characterized by high M_r/M_s , H_r , and H_r values and low S-ratio 414 (Figure 5). When plotting M_r/M_s versus H_{cr}/H_c on a Day Plot (Day et al., 1977), these anomalous 415 mineralogy intervals are characterized by the looping pattern often documented in sulfate reducing 416 sediments (Rowan et al., 2009; Yamazaki et al., 2003). This has been interpreted as due to the 417 enrichment of SP grains pushing values to higher H_{cr}/H_c values, and then increased abundance of single 418 domain magnetic minerals increasing M_r/M_s while decreasing H_{rr}/H_c (Figure 5b). When M_r/M_s is plotted 419 against H_{cr}, the looping pattern disappears, and the values plot along a single mixing relationship 420 between the high M_r/M_s and H_{cr} and low M_r/M_s and H_{cr} values, which supports the interpretation that 421 the looping pattern caused by H_{rr}/H_c variation is driven by the abundance of SP grains influencing the in-422 field parameter (H_c) and not the remanence parameter (H_{cr}). Unlike prior interpretations, H_{cr} values are 423 higher and S-ratio values are lower for the high coercivity horizons at 206 and 442 cmbsf than might be 424 expected from the addition of authigenic greigite alone (Horng, 2018). In other environments, similar 425 horizons with low S-ratios have been attributed to pyrrhotite (Kars and Kodama, 2015) and 426 measurements of a variety of magnetic iron sulfides suggest that these coercivities could be explained as 427 a mixture of the altered detrital assemblage observed above and below this horizon and authigenic 428 hexagonal 3C pyrrhotite (Horng, 2018).

429 Zone 3b is different than Zone 3a in that it has higher concentrations of ferrimagnetic minerals, 430 as indicated by χ_{lf} , M_s , and M_r , lower coercivity of remanence bearing minerals, as indicated by M_r/M_s 431 and H_{cr} , and lower proportions of high-coercivity (>300 mT) minerals, as indicated by higher S-ratios 432 (**Figures 5**). [S²⁻] and [Fe²⁺] are negligible at these depths, so this less-altered magnetic mineral 433 assemblage potentially reflects a preserved signal of past diagenetic conditions.

434

435 4.2.4 Detailed Rock Magnetic Investigations

436

437 4.2.4.1 First order reversal curves (FORCs)

FORCs were measured to investigate the primary magnetic mineral assemblage (0 cmbsf), the altered assemblage in the lower half of the SMT (148 cmbsf), the altered assemblage well below the SMT in the methanic zone (458 cmbsf), and at the anomalous magnetic mineral horizon recognized by high values of HIRM and remanence sensitive parameters (i.e., M_r , M_r/M_s , H_c , and H_{cr}) and low S-ratio values (206 cmbsf) (Figures 3 and 5). The 0 cmbsf sample has a signature consistent with a primary detrital assemblage that consists of a range of magnetic domain states, including single domain and vortex state grains, while the 148 and 458 cmbsf samples display lower coercivities consistent with a mix of vortex state, multidomain, and possibly SP grains (Figure 6a). Compared with FORC diagrams from

446 other sulfidic sediments, neither the 148 nor the 458 cmbsf sample are consistent with FORC signatures

for pure SP/single domain greigite or hexagonal 3C pyrrhotite (Roberts et al., 2018b); however, if those

448 minerals are present they may not be easy to recognize if they exist in a mixture with the altered

primary detrital assemblage. Comparison of the primary detrital assemblage (0 cmbsf) and the altered
 assemblage (148 and 458 cmbsf) suggests that magnetic mineral dissolution in the SMT primarily

451 removed the finest single-domain to vortex state grains from the primary assemblage.

452 The 206 cmbsf sample has a FORC distribution that extends to higher H_c with a tighter 453 distribution around the H_u = 0 mT axis, but still contains some spread indicating magnetostatic 454 interactions. This FORC diagram likely represents a mixture of the altered detrital assemblage (± SP 455 grains) observed in the 148 cmbsf and 458 cmbsf samples and an anomalous higher coercivity magnetic 456 mineral assemblage. Compared with FORC diagrams from other sulfidic sediments, this assemblage is 457 consistent with a mixture of the altered detrital assemblage observed in the 148 and 458 cmbsf samples, 458 SP/single domain greigite, and another mineral with coercivity higher than typical examples of greigite 459 bearing samples (Roberts et al., 2018b). The FORC distribution is somewhat skewed to more negative H 460 values in the 40 – 80 mT H_c range and extends beyond the measured H_c range that could indicate fine 461 grained and/or authigenic sedimentary 3C pyrrhotite contribution to the coercivity distribution of the 462 206 cmbsf sample (Horng, 2018). We note that the 206 cmbsf has differences from past examples of 463 pyrrhotite interpreted to be present in diagenetic sediments (e.g., Horng, 2018; Kars and Kodama, 2015; 464 Larrasoaña et al., 2007), which we attribute to the mixture of detrital and authigenic minerals in the 465 sample. While hematite could also explain the high coercivities (Roberts et al., 2006), this would require a high weight percentage of hematite relative to the lower coercivity phases, which is inconsistent with 466 no apparent decrease ~670 °C in the high temperature $\chi_{\rm ff}$ curves (Carvallo et al., 2006; Liu et al., 2019). 467

468

469 4.2.4.2 Isothermal remanent magnetization (IRM) acquisition

IRM acquisition curves were measured at or near the same intervals at the FORC diagrams to
further investigate the coercivities of the remanence bearing grains. IRM values at each acquisition step
are reported after being normalized by the IRM acquired at 1 T, as IRMs measured above 1 T were
imparted using a different coil and have slightly lower values than those collected at 1 T and below.
There is no significant remanence acquisition from 2 to 5 T for any samples.

475 The 0, 148, and 450 cmbsf IRM acquisition curves are similar, which was expected with S-ratio 476 values from 0.96 to 0.99. The initial acquisition at 50 mT is slightly lower for the 0 cmbsf sample (0.50) 477 relative to the 148 cmbsf (0.54) and the 450 cmbsf sample (0.56), but changes with the 148 and 450 478 cmbsf samples with a slightly stronger IRM acquired between 100 and 300 mT (e.g., IRMs at 150 mT are 479 0.93, 0.88, and 0.91, respectively) (Figure 6). All three samples reach values of at least 0.99 by the 500 480 mT step. This is consistent with the altered assemblages in the 148 and 450 cmbsf samples coarsening 481 slightly at lower coercivities, but also including greater (but still low) concentrations of a higher 482 coercivity component that saturates above 300 mT but below 500-600 mT.

The 206 cmbsf sample is unique in that it has a lower S-ratio (0.87) and acquires IRM more slowly than the other samples, with only 0.26 acquired by 50 mT, 0.65 acquired by 150 mT, and 0.92 by 300 mT (**Figure 6**). However, the 206 cmbsf sample also reaches 0.99 by 500 mT, suggesting the high 486 coercivity values are driven by a mineral that saturates at fields greater than 300 mT, thus driving S-ratio

- and HIRM variability around this horizon, but lower than about 500-600 mT. Greigite is generally
- 488 reported as having coercivities like that of magnetite, saturating by around 300 mT, and single domain
- 489 greigite has been shown to acquire a remanence faster than the 206 cmbsf sample (normalized
- 490 remanence of about 0.90 at 150 mT; Roberts et al., 2011). Comparison of the S-ratio of metamorphic
- 491 monoclinic 4C pyrrhotite (0.96-1.00), sedimentary greigite nodules (0.98-1.00), and sedimentary
- authigenic hexagonal 3C pyrrhotite nodules (0.52-1.00), suggest the authigenic 3C pyrrhotite can be
 distinguished from other magnetic iron sulfides based on its higher coercivity (Horng, 2018; Horng and
- distinguished from other magnetic iron sulfides based on its higher coercivity (Horng, 2018; Horng and
 Roberts, 2018). So, the presence of greigite cannot explain the high coercivity of the 206 cmbsf sample,
- 495 but authigenic hexagonal 3C pyrrhotite could.

496

497 4.2.4.3 Magnetic susceptibility (χ_{tf}) as a function of heating in argon and ambient air

498 To further investigate if thermomagnetic curve features discussed in Section 4.2.2 and 499 presented in Figure 4 reflect magnetic phase transitions or mineral alteration upon heating, we 500 remeasured four samples (using fresh material from the same core horizon) for χ_{if} as a function of 501 heating to ~700 °C and cooling back to room temperature in ambient air, at or around the same intervals 502 as the FORC measurements (Figure 6c). The heating curves measured in ambient air are nearly identical 503 to those measured in argon atmospheres, while the cooling curves measured in ambient air at 504 temperatures less than ~450 °C are always greater than those measured in argon atmospheres. In the 0, 505 206, and 500 cmbsf ambient air thermomagnetic curves, a local $\chi_{\rm fr}$ maximum is introduced in the cooling 506 curve between ~200 and 300 °C, which suggests that heating in ambient air facilitates alteration of a 507 mineral phase, such as greigite or other iron sulfides (Chang et al., 2008; Roberts et al., 2011; Schwarz 508 and Vaughan, 1972; Sweeney and Kaplan, 1973), to an antiferromagnetic hexagonal pyrrhotite polytype, one that is different from the hexagonal 3C pyrrhotite with ferrimagnetic room temperature properties 509 510 (Horng, 2018, discussed elsewhere). This feature is present with similar amplitude in both the argon and 511 ambient air measurements at 100 cmbsf. The similarity of the heating curves support the earlier 512 interpretation in Section 4.2.2 that the features of the heating curves are related to magnetic phase 513 transitions and that our primary magnetic assemblage is dominated by magnetite, maghemite, and 514 titanomagnetite, which is modified during diagenesis.

515

516 4.2.4.4 Zero field cooled (ZFC) and field cooled (FC) remanence

517 To investigate the low temperature properties of the remanence bearing magnetic minerals we 518 measured ZFC and FC remanences at four intervals at or near where the FORC measurements were 519 made (Figure 6d). While these measurements are not typically considered useful for the identification or 520 characterization of potential authigenic minerals that lack low-temperature order-disorder transitions 521 like greigite (Roberts et al., 2011), they do offer an additional constraint on the magnetic mineral 522 assemblage present.

523 Low temperature data do not contain a clear phase transition at 30-34 K, as would be expected 524 for monoclinic 4C pyrrhotite (Rochette et al., 1990). However, this transition is not detected in 525 authigenic sedimentary hexagonal 3C pyrrhotite nodules, which have similar behavior to greigite in FC 526 and ZFC measurements (Horng, 2018; Horng and Roberts, 2018). A magnetization decrease is present in 527 all samples between 100 and 120 K, which is indicative of a Verwey transition in magnetite. Quantifying 528 the remanence loss using the δT_v parameter of Moskowitz et al. (1993) ($\delta T_v = LTSIRM_{80k}$ – 529 LTSIRM_{150K}/LTSIRM_{80K}; where LTSIRM is the field cooled low temperature saturation IRM) indicates a 530 similar remanence loss in the altered magnetic mineral assemblage for samples 100 and 450 cmbsf (δT_v 531 = 0.47 and 0.51, respectively) which is somewhat greater than for the primary magnetic mineral 532 assemblage sample at 0 cmbsf ($\delta T_v = 0.37$). The anomalous magnetic mineral assemblage at 206 cmbsf 533 has the smallest remanence decrease ($\delta T_v = 0.21$). Suppression of the Verwey transition in the primary 534 magnetic assemblage relative to the altered assemblage samples could be related to the presence of 535 oxidized maghemite, as recognized in the high temperature $\chi_{\rm f}$ curves (Özdemir et al., 1993). Even 536 greater suppression of the Verwey transition in the anomalous assemblage at 206 cmbsf that lacks 537 strong evidence for maghemite could be explained by the presence of populations of fine SD size 538 magnetic grains and/or greigite (Moskowitz et al., 1993) or authigenic hexagonal 3C pyrrhotite (Horng 539 and Roberts, 2018).

540 The remanence loss above the Verwey transition varies for each sample as quantified by the δM 541 parameter ($\delta M = (LTSIRM_{150k} - LTSIRM_{300k})/LTSIRM_{150k}$) (Hatfield et al., 2017; Moskowitz et al., 1998). The 542 gradual remanence decrease above the Verwey transition is smaller for the primary detrital assemblage 543 at 0 cmbsf ($\delta M = 0.29$) than for the altered assemblages in the 100 and 450 cmbsf samples ($\delta M = 0.45$ 544 and 0.48, respectively). This increase in δM is consistent with the high temperature χ_{μ} data that indicate 545 an increased relative proportion of titanomagnetite to stochiometric magnetite in the altered magnetic 546 mineral assemblages. The anomalous magnetic mineral assemblage at 206 cmbsf has an intermediate 547 remanence decrease above the Verwey transition ($\delta M = 0.37$), which is likely related to the mixture of 548 the altered magnetic mineral assemblage with another magnetic mineral phase. High temperature $\chi_{\rm fr}$ 549 behavior rules out significant stochiometric magnetite or titanomagnetite with a different composition 550 than other samples below the dissolution front. Alternative candidates could be a higher proportion of 551 iron sulfides like fine grained authigenic greigite or hexagonal 3C pyrrhotite within the altered magnetic 552 mineral assemblage, which have only a small decrease above the Verwey transition temperature (Chang 553 et al., 2009; Horng and Roberts, 2018).

ZFC remanence values are typically greater than FC remanence values for the primary (0 cmbsf) and altered samples (100 and 450 cmbsf), but less than FC remanence values for the anomalous sample at 206 cmbsf. Quantifying this difference using the R_{LT} parameter of Smirnov (2009) (R_{LT} = FC SIRM_{20K}/ZFC SIRM_{20K}) yields similar values for the 0, 100, and 450 cmbsf samples (R_{LT} = 1.04, 1.06, and 1.05, respectively) and a higher value for the anomalous magnetic mineral assemblage at 206 cmbsf (R_{LT} = 1.16), which is consistent with the presence of vortex state grains in the primary and altered assemblages and fine-grained single domain grains in the 206 cmbsf sample (Smirnov, 2009). Like the

561 206 cmbsf sample, greigite and hexagonal 3C pyrrhotite samples have FC curves greater than ZFC curves

562 (Chang et al., 2008; Horng and Roberts, 2018).

563

564 4.2.4.5 Low temperature and frequency dependence of magnetic susceptibility

565 To further investigate the low temperature properties of the magnetic mineral assemblages, we 566 measured $\chi_{If-MPMS}$ as a function of temperature below room temperature at a variety of frequencies for 567 three intervals representative of the primary detrital assemblage (0 cmbsf), altered detrital assemblage 568 (100 cmbsf), and anomalous assemblage (206 cmbsf) (Figure 6e). Frequency dependent magnetic susceptibility ($X_{fd} = (\chi_{1 Hz} - \chi_{31.6 Hz})/\chi_{1 Hz} \times 100$) is highest at 300 K for the 206 cmbsf sample and lowest for

- 570 the 0 cmbsf sample with the 100 cmbsf sample falling in between. These values and the increasing χ_{fd}
- values with temperature above 100 K suggest enrichment of SP and single domain grains in the altered
- 572 assemblage captured by the 100 cmbsf sample and the anomalous assemblage captured by the 206
- 573 cmbsf sample relative to the 0 cmbsf sample (Worm, 1998; Worm and Jackson, 1999). χ_{fd} peaks between 574 about 60 and 70 K can be attributed to thermally activated electron hopping in titanomagnetite (Carter–
- 575 Stiglitz et al., 2006; Hatfield et al., 2017; Walz et al., 1997).
- 576

577 4.2.5 Scanning Electron Microscope Observation of Fe-Ti Oxides and Fe Sulfides

578 Unpolished heavy mineral extract samples representative of the altered detrital assemblage at 579 148 and 450 cmbsf and the anomalous magnetic mineral assemblage at 206 cmbsf were observed to 580 assess the potential contributors to the sediment magnetism. While it is perhaps likely that magnetic 581 mineral inclusions in silicate minerals should be considered (e.g. Chang et al., 2016b; Hatfield et al., 582 2017), they could not be imaged with our unpolished samples. Fe-Ti oxides, when observed, have 583 evidence of surface pitting or other dissolution structures, often with iron sulfides present on the 584 mineral surface and in cavities (Figure 7a-c), similar to observations from other reducing sediments 585 (Canfield and Berner, 1987; Channell and Hawthorne, 1990; Nowaczyk, 2011). Other minerals were 586 completely encased by iron sulfides (Figure 7d). In elemental mapping, pitted regions of Fe-Ti oxides 587 often had higher Ti concentrations relative to Fe (Figure 7a). Pyrite framboids are common in all 588 samples, reaching tens of μ m in size and formed with pyrite crystals typically between 1 and 2 μ m 589 (Figure 7e-g).

590 In a single instance, a platy iron sulfide mineral was observed in the 148 cmbsf sample with 591 crystals about $1.5 - 2 \mu m$ in diameter (Figure 7h). We consider this tentative visual evidence for the 592 presence of fine-grained hexagonal 3C pyrrhotite in Core JKC36. We also observed a single instance of 593 an iron sulfide mineral framboid composed of crystals with a unique shape (Figure 7i-k). The crystals are 594 octahedral in shape and are about 0.5 - 1 μ m in size. We consider this visual evidence for the presence 595 of fine-grained greigite (likely on the border between single domain and vortex state; Roberts et al., 596 2011) in Core JKC36. However, magnetostatic interactions in greigite framboids could make their 597 magnetic signature appear like that of a larger, multi-domain particle (Valdez-Grijalva et al., 2020). For 598 both these minerals, the only major elements present are Fe and S, but we are unable to quantify the 599 ratio of the two elements for these unpolished samples. Given the presence of these iron sulfides with 600 large enough sizes to be imaged using an SEM, it is likely that there are additional, smaller iron sulfides 601 that are below the imaging capabilities of the SEM system.

602

603 4.2.6 Magnetic Mineralogy of Potential Source Rocks

604To compare the Core JKC36 magnetic properties, particularly for the anomalous magnetic

605 mineral assemblage at 206 cmbsf, with potential source rock magnetic properties, magnetic

606 measurements from PRR rock samples that were available in the Joinville Island region were

- 607 investigated and indicate a wide range of magnetic grain sizes in the regional bedrock(**Figure 8**). The
- samples with the highest M_s values are gabbros from small nearby islands. A Heroina Island gabbro (e.g.
- Hamer and Hyden, 1984; Watts et al., 1984) has fine magnetic mineralogy ($M_r/M_s = 0.38$, $H_{cr}/H_c = 1.66$,

- 610 $H_{cr} = 79.41$, S-ratio = 0.99). A Wideopen Island gabbro has coarser magnetic mineralogy ($M_r/M_s = 0.06$,
- 611 $H_{cr}/H_c = 2.63$, $H_{cr} = 6.86$, S-ratio = 1.04) (Figures 1 and 8). χ_{lf} as a function of heating from room
- 612 temperature to ~700 °C in an argon atmosphere indicates that the magnetic mineralogy of these
- samples is dominated by low titanium Fe-Ti oxides with χ_{if} decreases between 500 and 580 °C for the
- 614 two samples (Figure 8).

615 Sedimentary and volcanic rocks from the Nordenskjöld Formation (e.g. Elliot, 1967;

- 616 Farquharson, 1983), Northern Joinville Island, have weaker M_s and relatively coarse magnetic grain size
- 617 (Figures 1 and 8). χ_{lf} curves as a function of heating from room temperature to ~700 °C are dominated by
- 618 steady declines that can be attributed to paramagnetic minerals and significant alteration creating
- 619 higher $\chi_{\rm lf}$ above ~400 °C (Figure 8). One of the stronger Nordenskjöld Formation samples and the sample
- 620 with the finest magnetic mineralogy ($M_r/M_s = 0.19$, $H_{cr}/H_c = 2.32$, $H_{cr} = 34.64$, S-Ratio = 0.99) is an 621 andesite from King Point which has a steady magnetization decline between 500 and 600 °C, which
- 622 indicates a magnetic mineral assemblage consisting of low titanium Fe-Ti oxides (Figure 8).

623 A diorite sample from Mt. Alexander, southern Joinville Island (e.g. Grunow, 1993), has a 624 relatively fine magnetic mineralogy ($M_r/M_s = 0.21$, $H_{cr}/H_c = 2.13$, $H_{cr} = 38.77$, S-ratio = 0.82) (Figures 1 625 and 8). This sample has an abrupt $\chi_{\rm fr}$ increase at ~235 °C and abrupt $\chi_{\rm fr}$ decrease at 335 °C and 580 °C, 626 which indicate the cooccurrence of ferrimagnetic monoclinic 4C pyrrhotite and antiferromagnetic 627 hexagonal pyrrhotite polytypes (e.g. Rochette et al., 1990) and magnetite (Figure 8). Like the 628 Nordenskjöld Formation samples, the Mt. Alexander Diorite experiences significant alteration at high 629 temperatures, with χ_{lf} at room temperature after heating and cooling one order of magnitude larger 630 than the initial $\chi_{\rm f}$ measurement. While there is direct evident for pyrrhotite in this potential source rock, 631 the evidence is for pyrrhotite polytypes that are different from the hexagonal 3C pyrrhotite with room 632 temperature ferrimagnetic properties (Horng, 2018) that is most consistent with our rock magnetic data from Core JKC36 (see Figure 9 and discussion below). 633

634

635 5. Discussion

636

637 5.1 Magnetic Mineral Dissolution

638 Core JKC36 experiences significant detrital magnetic mineral dissolution through the SMT. 639 However, not all detrital iron-titanium-oxides are dissolved (Figures 4 and 6). Ferrimagnetic 640 (titano)magnetite steadily dissolves as $[S^2]$ increases near the top of the SMT in the upper 30 cm of the 641 core (Figure 3) with dissolution preferentially removing stochiometric magnetite to titanomagnetite (Figure 4) and finer (titano)magnetite to coarser (titano)magnetite (Figures 5). This is consistent with 642 previous observations and is likely related to stochiometric magnetite having a higher ratio of Fe³⁺ to Fe²⁺ 643 644 than titanomagnetite, and finer magnetic minerals having a higher surface area to volume ratio (Canfield 645 and Berner, 1987; Dillon and Bleil, 2006; Karlin, 1990). In the upper 30 cm, ferrimagnetic iron-titanium 646 oxide dissolution is accompanied by dissolution of iron-bearing paramagnetic minerals, which is tracked 647 by a decline in χ_{hf} , but not the dissolution of higher coercivity minerals (likely hematite) that are part of 648 the primary detrital assemblage, as indicated by no change in HIRM (Figure 3).

- 649 A dissolution front occurs between 30 and 32 cmbsf, where there is a large drop in parameters
- 650 sensitive to the concentration of (titano)magnetite (Figure 3). This dissolution front occurs in the upper
- 651 SMT, when sulfide is only at around one-third of its maximum observed value. Below the dissolution
- 652 front, (titano)magnetite dissolution is limited, likely due to the exhaustion of (titano)magnetite particles
- that are susceptible to dissolution. High coercivity detrital minerals that contribute little to magnetic
- 654 remanence relative to (titano)magnetite begin to dissolve below the dissolution front and continue to
- dissolve through the entire SMT where sulfide is elevated and below the depth of sulfate exhaustion.
- These high-coercivity detrital components, likely hematite, are probably a source of ferric iron
- 657 throughout the SMT for microbial respiration.
- 658

659 5.2 Authigenic Magnetic Minerals 1: Superparamagnetic Enrichment

660 As previously observed in diagenetic sediments (Tarduno, 1995), there is clearly a SP mineral 661 enrichment within the SMT and in sediments below the SMT, as seen through H_c , H_{cr}/H_c , and frequency 662 dependent $\chi_{If-MPMS}$ (Figures 5 and 6). This SP enrichment occurs in three stages, as follows.

663 1. Moving down core, the first signs of SP enrichment, as observed in the difference between the 664 in-field H_c and remanence parameter H_{cr} and supported by frequency dependent $\chi_{\text{If-MPMS}}$ (Figures 665 5 and 6e), occur just below the dissolution front in Zone 2a (33-87 cmbsf) where [Fe²⁺] is 666 depleted to background values, $[SO_4^{2-}]$ is present, and $[S^{2-}]$ is increasing downcore as it diffuses 667 upward from the reduction of sulfate at around 80 cm (Figures 2). While not clear, SP 668 enrichment is possibly also occurring above the dissolution front near the top of the SMT and is 669 obscured by the relative coarsening of the detrital assemblage (Figure 5). SP enrichment in 670 sulfate reducing sediments has previously been attributed to enrichment of fine-grained 671 ferrimagnetic iron sulfides like greigite (Rowan et al., 2009). However, it is generally thought 672 that preservation of minerals that form as intermediaries on the path to pyrite, like greigite, 673 require sulfide limitation relative to iron to preserve the intermediary mineral (e.g. Kao et al., 674 2004), which is not the case here as S^2 is accumulating in the porewater during sulfate reduction, suggesting its production from sulfate is greater than its consumption from reacting 675 676 with iron, and Fe^{2+} is not present in significant concentrations. While the nature of the Zone 2a 677 SP enrichment is not clear at this time, we could be observing transient intermediary 678 ferrimagnetic iron sulfides.

- 679 2. Below in Zone 2b (87-131 cmbsf), SP minerals are depleted relative to the overlying and 680 underlying sediments, as indicated by lower H_{cr}/H_c (Figure 5). In this zone, sulfide decreases 681 downcore in a concave-up shape, which suggests a continuous zone of consumption. 682 Meanwhile, [Fe²⁺] increases downcore to a local maximum near the base of Zone 2b, likely from 683 iron reduction near the base of the SMT perhaps due to microbial iron reduction coupled to 684 AOM (e.g. Beal et al., 2009; Riedinger et al., 2014). In Zone 2b, we also observe a χ_{hf} increase 685 relative to Zone 2a sediments (Figure 3), which indicates that pyrite is being created and 686 preserved at rates higher than iron-bearing paramagnetic minerals are being dissolved. The lack 687 of SP minerals in Zone 2b is almost certainly the result of sufficient $[Fe^{2+}]$ and $[S^{2-}]$ in the 688 porewater to let stable pyrite form without the preservation of SP ferrimagnetic iron sulfide 689 intermediaries.
- 690 3. Near the top of Zone 3a (> 131 cmbsf), around the base of the SMT, we observe variable SP 691 enrichment that is present to the base of the core, based on the range of H_{cr}/H_c variations

692 relative to Zone 2 (Figure 5). The top of Zone 3a is marked by a downcore decrease in sulfide to 693 negligible values below the SMT, which indicates consumption of sulfide that outpaces 694 production or diffusion now that we are far from where SO_4^{2-} is depleted. Fe²⁺ decreases 695 downcore which indicates dissolved iron production at rates faster than consumption by S²⁻ near 696 the base of the SMT. This likely creates conditions favorable for a [S²⁻]-limitation relative to 697 $[Fe^{2+}]$, which allows preservation of ferrimagnetic iron sulfides that are intermediaries on the 698 pyrite formation pathway. These SP minerals are then likely preserved in the underlying methanic sediments when $[S^2]$ and $[Fe^{2+}]$ are depleted and at background levels. While we 699 700 cannot rule out the continued production of SP iron sulfides below this horizon near the base of 701 the SMT, we consider it likely that most of the SP minerals observed in the methanic sediments 702 beneath the SMT were formed at this horizon near the base of the SMT when porewater 703 conditions were favorable for their formation.

704

5.3 Authigenic Magnetic Minerals 2: Remanence-Bearing Iron Sulfides

706 Horizons with high concentrations of high coercivity minerals tracked by HIRM occur at a few 707 intervals, notably at 206 and 442 cmbsf (Figure 3). These horizons, which only occur at the bottom of the 708 SMT and below, have sharp bases that grade upward from high concentrations at the base to lower 709 concentrations above (Figure 3). High temperature $\chi_{\rm f}$ heating curves do not indicate any anomalous 710 magnetic minerals in these horizons (Figure 4). Perhaps this is not surprising because while these layers 711 are easy to pick out using remanence parameters, like M_{r} , HIRM, and H_{cr} , they cannot be distinguished 712 stratigraphically using in-field concentration dependent magnetic parameters like χ_{if} and M_s (Figures 3, 713 5). These layers produce a looping pattern on a Day Plot (Figure 5), which has been interpreted as 714 diagnostic of greigite formation, with SP enrichment first pushing values to the right on the Day Plot and 715 then continued growth to stable single domain sizes pushing values toward the upper left-hand corner 716 (Rowan et al., 2009). However, the high remanence coercivity, tracked by H_{cr} values, and saturation 717 above 300 mT, tracked by low S-ratio values, is inconsistent with observations of sedimentary greigite 718 (Horng, 2018; Roberts et al., 2011). Thus, while authigenic greigite is likely present in the assemblage as 719 observed in SEM images (Figure 7), it cannot be the only mineral added to the altered detrital 720 assemblage observed elsewhere in the core below the SMT. We also consider significant hematite 721 concentrations unlikely because we see no indication of hematite in the high temperature χ_{if} curves 722 given the amount of hematite required to lower the S-ratio to 0.87 (Frank and Nowaczyk, 2008).

723 The magnetic mineral that is most likely to explain our observation of high-coercivity horizons, 724 like the one that exists at the base of the SMT around 200 cm, is authigenic hexagonal 3C pyrrhotite. 725 Recent detailed documentation of rock magnetic properties from sedimentary hexagonal 3C pyrrhotite 726 nodules have challenged previous thoughts on authigenic pyrrhotite formation in sediments (Horng, 727 2018; Horng and Roberts, 2018). Comparison of our H_{cr} and S-ratio data with the observations of Horng 728 (2018) suggest that the high coercivity layers in Core JKC36 are likely a mixture of the altered detrital 729 assemblage and authigenic remanence bearing greigite and hexagonal 3C pyrrhotite (Figure 9). This is 730 consistent with the low temperature remanence data, IRM acquisition data, FORC diagrams, and SEM observations (Figures 6 and 7). While monoclinic 4C pyrrhotite and an antiferromagnetic pyrrhotite 731 732 polytype is observed in the regional diorite bedrock sample, PRR16328 (Figure 8), the diorite's magnetic 733 properties are not similar to the magnetic properties of the high coercivity mineral horizon at 206 cmbsf 734 in JKC36 (Figure 9). This difference is expected because the pyrrhotite minerals found in the diorite have

735 different magnetic properties than authigenic hexagonal 3C pyrrhotite (Horng, 2018; Horng and Roberts, 736 2018). The relationship between H_{cr} and S-ratio in Zone 1 is different than the relationship in Zones 2 737 and 3 (Figure 9c-d), which suggests that the magnetic minerals driving S-ratio and HIRM variability are 738 largely different above the dissolution front versus below—which likely indicate that while in Zone 1 739 HIRM and S-ratio reflect the preferential dissolution of detrital (titano)magnetite relative to hematite, in 740 Zones 2 and 3 HIRM and S-ratio largely reflect the addition of authigenic hexagonal 3C pyrrhotite. The 741 trend of these samples in Zone 3 towards where the sedimentary 3C pyrrhotite nodules of Horng (2018) 742 plot is different from the trends that seem more typical of detrital assemblages, as observed above the 743 dissolution front in Zone 1; in more terrigenous Antarctic Peninsula Fjord glacial marine sediments with 744 less extreme diagenetic conditions (Reilly et al., 2016); in lake sediments that are interpreted to be a 745 mixture of magnetite and hematite (Stober and Thompson, 1979); and the regional bedrock samples 746 (Figure 9).

747 The largest of the high-coercivity layers occurs at the base of the SMT with its peak at 206 748 cmbsf, where [S²⁻] is almost completely depleted and below where [Fe²⁺] accumulates in the porewater 749 at the base of the SMT (Figure 3). We therefore hypothesize that this layer forms transiently at the 750 porewater transition where downward diffusing [S²⁻] is limited but there is still a source of [Fe²⁺], and 751 thus porewater conditions are favorable for the growth of remanence-bearing hexagonal 3C pyrrhotite 752 and greigite. The source of $[Fe^{2+}]$ following the complete reduction of sulfate may be AOM coupled to 753 microbial iron reduction (Beal et al., 2009; Riedinger et al., 2014). Microbial use of a ferric iron source 754 such as hematite could in part explain the slower dissolution rates of high-coercivity iron oxide 755 components of the detrital assemblage. In this case, the JKC36 206 cmbsf remanence-bearing layer is 756 created by the balance of [Fe²⁺] production by AOM coupled to microbial iron reduction and downward 757 [S²⁻] diffusion.

758 The formation of this transient remanence bearing iron sulfide layer at a redox boundary is 759 much like the formation of transient manganese oxide horizons near the top of the manganese 760 reduction zone, which are created by the balance of oxygen diffusing downward and Mn²⁺ diffusing 761 upward (Froelich et al., 1979). Thus, analogous to preserved manganese oxide layers (Froelich et al., 762 1979), we predict that electron donor and/or organic carbon flux perturbations to steady state 763 diagenesis through the SMT should lead to preservation of horizons with high concentrations of 764 remanence-bearing hexagonal 3C pyrrhotite and greigite within the methanic zone such as the smaller 765 layer that peaks at 442 cmbsf (Figures 3 and 5).

766

767 5.4 Non-Steady State Diagenesis in Relation to Paleoenvironmental Conditions

768 There is evidence in Core JKC36 for non-steady state diagenesis over the time the sediments 769 were deposited, including a shift in magnetic mineral assemblage properties from Zone 3a to 3b around 770 325 cmbsf and preservation of a horizon with high concentrations of the remanence-bearing iron 771 sulfides, greigite and hexagonal 3C pyrrhotite, around 442 cmbsf (Figures 3, 5, and 9). Kyrmanidou et al. 772 (2018) observed an anomaly in the diatom assemblage over this time as well in the ratio of 773 Chaetocerous subg. Hyalochaete resting spores to cells (Figure 10). High concentrations of Chaetocerous 774 subg. Hyalochaete resting spores are generally interpreted as indicating high primary productivity during 775 the spring bloom (Buffen et al., 2007; Crosta et al., 1997; Leventer, 1991; Leventer et al., 1996, 2002). 776 Chaetocerous subg. Hyalochaete resting spores form as a survival mechanism when nutrients are

777 depleted after high primary productivity and low light during the polar winter following a large spring

bloom (Leventer et al., 1996; Crosta et al., 1997). A peak in the ratio of resting spores to cells between

about 150 and 275 cmbsf suggests that conditions in the late Holocene (~250-400 cal yrs BP) supported

780 higher primary productivity for a brief time.

781 If this primary productivity perturbation also led to increased organic matter export from the 782 surface ocean to the seafloor, the increased organic matter would fuel increased respiration of 783 sedimentary microbial communities and place a greater demand on electron acceptors, causing a shift in 784 the porewater geochemistry. To illustrate this idea, we offset the modern SMT to the core depth just 785 below the start of this primary productivity perturbation to conceptualize what the pore water profile 786 might have looked like at this time (Figure 10). This is a simplification that assumes other factors, such as 787 sedimentation rates, remain constant, which otherwise could drive non-steady state diagenesis itself. 788 However, this scenario is consistent with our observation that the depth of the transition from Zones 3a 789 to 3b between 300 and 350 cmbsf, which is marked by a M_s increase and H_{cr} decrease, is about where 790 we might expect the dissolution front to be at this time (solid gold line in Figure 10). Moreover, the 791 location of the deeper horizon of high concentration of remanence-bearing iron sulfide minerals around 792 442 cmbsf is around the base of this shifted SMT (dashed gold line in Figure 10). Therefore, we 793 hypothesize that this primary productivity perturbation increased the organic carbon supply to the 794 seafloor, which altered the porewater geochemistry including increasing the sulfide, leading to greater 795 magnetic mineral dissolution in Zone 3a relative to Zone 3b, and preserving a horizon of remanence-796 bearing iron sulfides at the base of the pre-perturbation SMT.

797

798 6. Conclusions

We make new observations of magnetic mineral diagenesis in relation to porewater geochemistry in a sedimentary record with a shallow SMT from Perseverance Drift, Northwestern Weddell Sea. While more work is needed to investigate the kinetics of these processes, the following conclusions are reached for our work.

- A (titano)magnetite dissolution front associated with elevated [S²⁻] in the upper portion of the
 SMT occurs when [S²⁻] is about one-third of its peak concentration, almost 1 meter
 stratigraphically above where sulfide reaches peak concentrations in the sediment porewater.
- During magnetic mineral dissolution, magnetite is preferentially dissolved relative to
 titanomagnetite. While magnetite is exhausted in the upper part of the SMT, dissolution of
 minerals with coercivity greater than 300 mT (likely hematite) continues through the entire SMT.
 This could be an important source of ferric iron within the SMT.
- 810 SP enrichment and depletion in this SMT occurs in three phases. First, when [Fe²⁺] is negligible 811 relative to $[S^{2-}]$ in the upper SMT where $[SO_4^{2-}]$ decreases downcore, SP minerals are enriched. SP 812 concentrations decrease in the zone below peak [S²⁻] concentrations, where the concave-up 813 shape of the sulfide profile indicates consumption, and $[Fe^{2+}]$ increases to above background 814 concentrations. These conditions are likely more favorable for pyrite formation than 815 ferrimagnetic iron sulfide formation. At the base of the SMT, where [Fe²⁺] is being produced or 816 diffused faster than it is consumed and [S²] concentrations are low, SP minerals are enriched 817 again and preserved in the underlying methanic sediments.

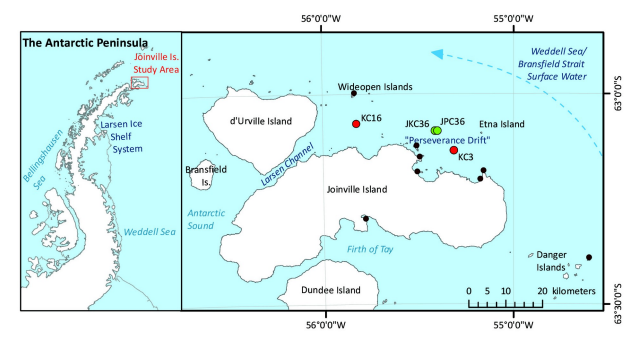
- 818 Horizons with high concentrations of high coercivity minerals that contribute significantly to the 819 bulk sediment remanent magnetization form at the base of the SMT where [S²⁻] is limited and 820 $[Fe^{2+}]$ is produced by iron reduction. During steady-state diagenesis, this horizon is likely 821 transient and moves with the SMT as sediment accumulates. Non-steady state diagenetic 822 changes driven by carbon flux or sedimentation rate changes could preserve these horizons in 823 the underlying methanic sediments. The magnetic properties of these horizons are consistent 824 with a mixture of an altered detrital magnetic mineral assemblage, greigite, and hexagonal 3C 825 pyrrhotite.
- Changes in the altered magnetic mineral assemblage preserved in the methanic zone are
 consistent with changes in past porewater conditions and the nature of the SMT at the studied
 site. This change to more extreme magnetic mineral diagenesis occurred following increased
 primary productivity at Perseverance Drift and was likely a response to increased export
 productivity to the seafloor. Non-steady state changes to porewater geochemistry likely led to
 preservation of a high concentration of remanence-bearing iron sulfide horizon at depth in the
 methanic zone that originally formed at the base of the paleo-SMT.
- 833

834 Acknowledgements

835 We thank the former Raytheon Polar services (now Lockheed Martin) shipboard team, the 836 LARISSA science team, and the captain and crew of the R/V Nathaniel B. Palmer for their efforts during 837 NBP1203. We thank A. Grunow, curator of the Polar Rock Repository (PRR), for her assistance and 838 support in identifying and supplying bedrock samples. This research used samples and data provided by 839 PRR and the Byrd Polar Research Institute, Ohio State University. The PRR is sponsored by the National 840 Science Foundation Office of Polar Programs (NSF-OPP). This research was supported by a visit to the 841 Institute of Rock Magnetism (IRM) by S.A. Brachfeld. The IRM is a US National Multi-user Facility 842 supported through the Instrumentation and Facilities program of the National Science Foundation, Earth 843 Sciences Division, and by funding from the University of Minnesota. This research was funded by NSF-844 OPP grant 0732917 to M.L. McCormick and NSF-OPP grant 0732605, NSF-MRI grant 0521069, and NSF-845 MRI grant 0619402 to S.A. Brachfeld. B.T. Reilly thanks the Scripps Institution of Oceanography for 846 support. Data are archived with the Magnetics Information Consortium (MagIC) (DOI: 847 10.7288/V4/MAGIC/16906). We thank Andrew Roberts and Myriam Kars for helpful and insightful 848 reviews.

- 849
- 850

851 Figures



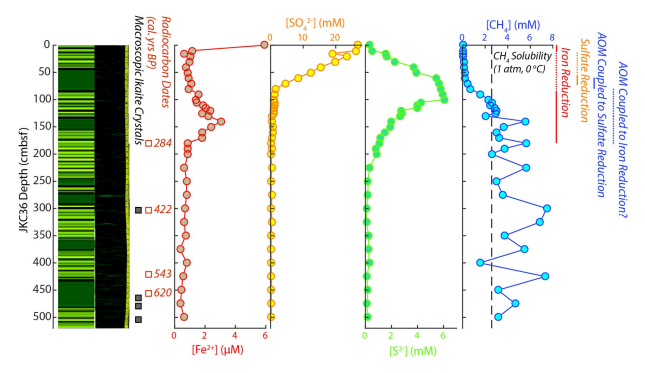
853 Figure 1. Perseverance Drift regional map. Left: Map of the Antarctic Peninsula with the Joinville Island

854 Study Area indicated with a red rectangle. Right: Location of Perseverance Drift, north of Joinville Island,

and sediment cores from Cruises LMG0404 (red) and NBP1203 (green) indicated. Polar Rock Repository

856 sample locations are also indicated (brown). Core JKC36 is the focus of this study.

857





859 Figure 2. JKC36 stratigraphy and porewater geochemistry. From left to right: lithologic log indicating

860 laminated (light green) versus bioturbated (dark green) intervals of JKC36 (Kyrmanidou et al., 2018);

861 photograph of JKC36 (dark areas are where samples were taken for microbiology and porewater

862 geochemistry immediately after recovery); brown squares indicate locations where macroscopic ikaite

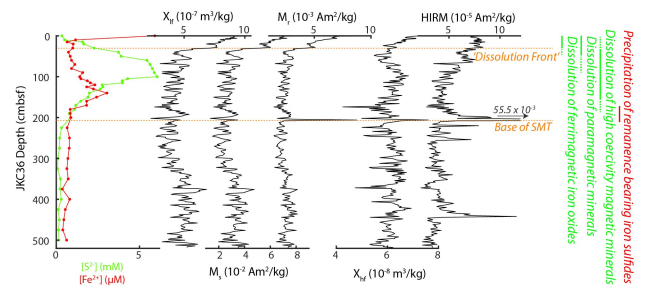
863 crystals were found; open orange squares indicate radiocarbon dates (corrected and calibrated in CALIB

864 version 7.1 using $\Delta R = 860$ yrs and the MARINE 13 dataset; Kyrmanidou et al., 2018); porewater

dissolved ferrous iron (Fe²⁺ in red), sulfate (SO₄²⁻ in gold), sulfide (S²⁻ in green), and methane (CH₄ in

866 blue). Methane solubility at 1 atmosphere and 0 °C indicated with vertical dashed black line.

867 Interpretation of porewater geochemistry included to the right (AOM = anaerobic oxidation of 868 methane).



871 **Figure 3. Concentration dependent magnetic parameters.** From left to right: porewater S²⁻ (green) and

872 Fe²⁺ (red) concentrations; low-field magnetic susceptibility (χ_{lf}); saturation magnetization (M_s); saturation

873 remanence (M_r); high-field susceptibility (χ_{hf}); and the hard isothermal remanent magnetization (HIRM).

874 The dissolution front around 30 cmbsf, where there is a sharp drop in χ_{lf} , M_s , and M_r , and the base of the

sulfate-methane transition (SMT) around 200 cmbsf, where S²⁻ returns to background levels are

876 indicated with horizontal dashed gold lines.

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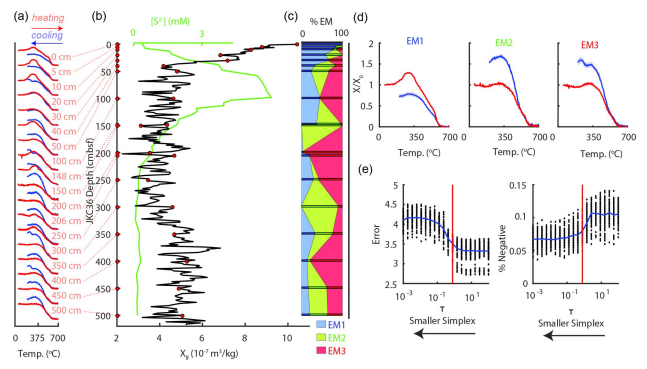
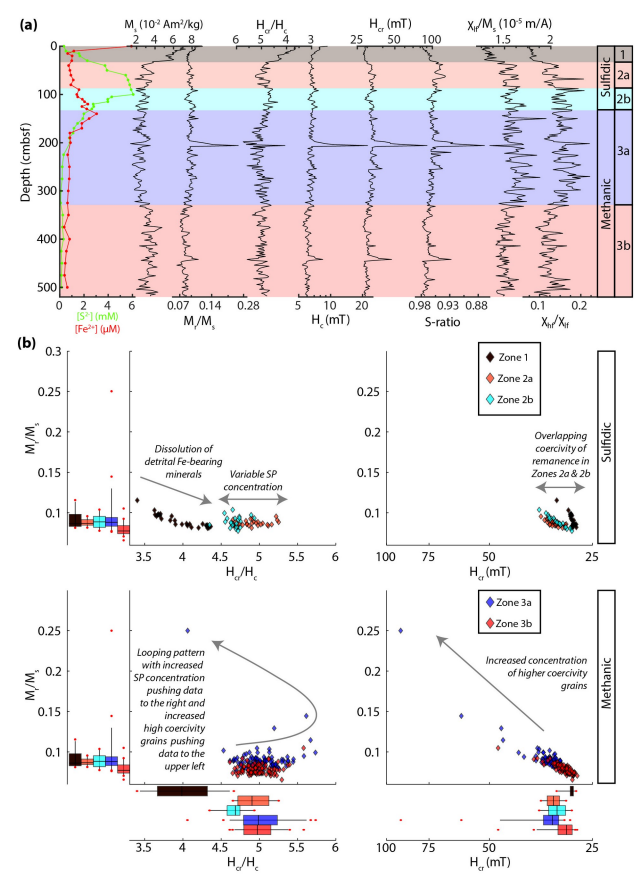


Figure 4. JKC36 high temperature magnetic susceptibility. (a) Magnetic susceptibility (χ_{if}) as a function 879 880 of temperature for 18 horizons in JKC36. Red curves are measurements made while heating from room 881 temperature to about 700 °C and blue curves are measurements made while cooling from about 700 °C 882 back to room temperature. (b) Room temperature χ_{μ} (black) compared with the S²⁻ concentration in 883 JKC36 porewater (green). Horizons with $\chi_{\rm ff}$ measurements as a function of temperature are indicated 884 with red dots. (c) Percent endmember 1 (blue), 2 (green), and 3 (red) from endmember modeling of the 885 temperature dependent χ_{if} curves. (d) End member temperature dependent χ_{if} curves (red = heating; 886 blue = cooling), with $\pm 1 \sigma$ (shading). (e) End member model selection (vertical red line; $\tau = 0.7848$) using 887 error (left) and shape constraints (right) for a range of regularization parameters (τ ; see Reilly et al. 888 (2016) for discussion).

889



- 891 Figure 5. JKC36 rock magnetic parameters. (a) From left to right: porewater S²⁻ (green) and Fe²⁺ (red)
- 892 concentration; saturation magnetization (M_s); ratio of saturation remanence to saturation magnetization
- 893 (M_r/M_s) ; ratio of the coercivity of remanence to the bulk coercivity (H_{cr}/H_c) ; bulk coercivity (H_c) ;
- 894 coercivity of remanence (H_{cr}); the S-ratio; ratio of low-field magnetic susceptibility to saturation
- 895 magnetization (X_{if}/M_s) ; ratio of high-field magnetic susceptibility to low field magnetic susceptibility (X_{if}/M_s)
- 896 X_{If}); and geochemical/magnetic zones discussed in the text. The dissolution front occurs at 33 cmbsf at
- the base of Zone 1. The base of the SMT is in Zone 3a around 200 cmbsf. **(b)** Day Plots (Day et al., 1977)
- and M_r/M_s plotted against H_{cr} . Box and whisker plots indicate the mean, the 84% percentile (box), and
- 899 the 95% percentile (whiskers). Data that plot outside of the 95% interval are indicated with red dots.

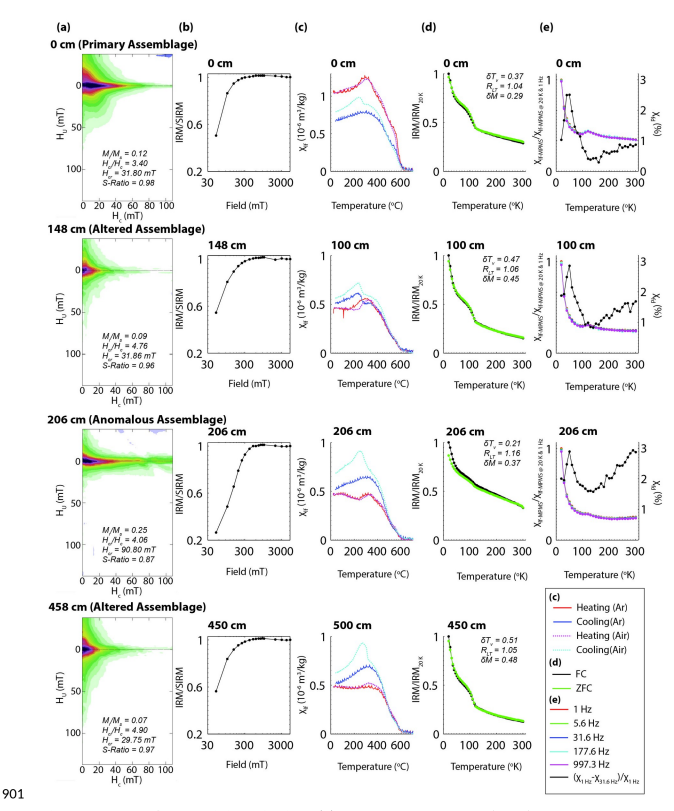
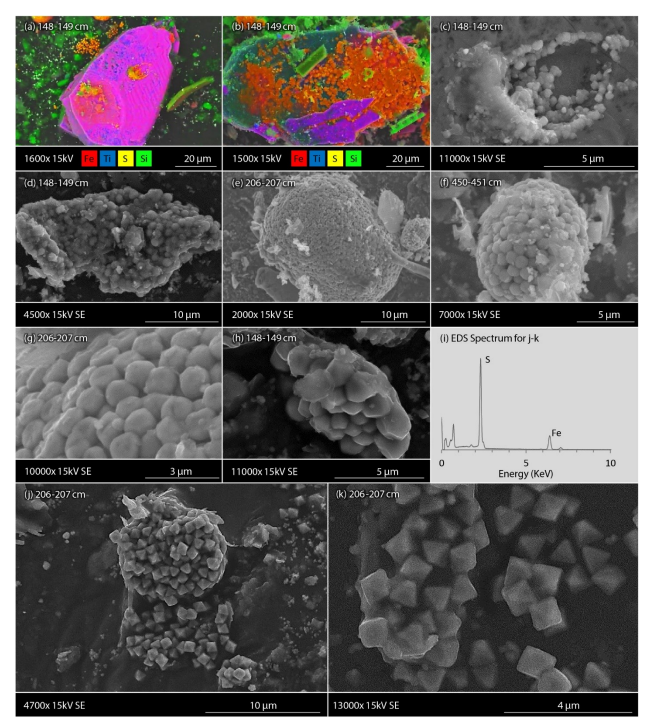


Figure 6. JKC36 rock magnetic investigations. (a) First order reversal curve (FORC) diagrams with
 hysteresis and S-ratio parameters listed. (b) Isothermal remanent magnetization (IRM) acquisition

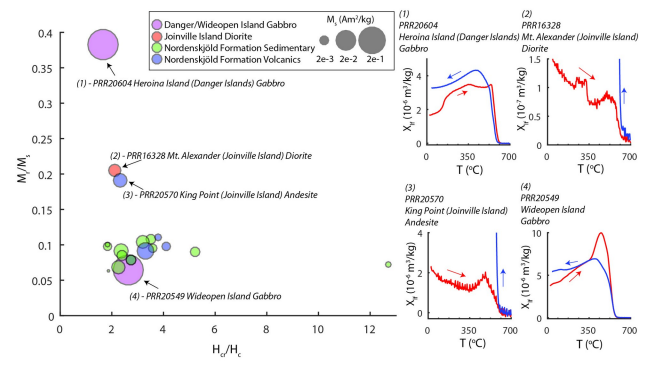
 904 curves normalized by the IRM at 1000 mT (SIRM). (c) Comparison of magnetic susceptibility ($\chi_{\rm f}$) as a

- 905 function of temperature when measured in an argon atmosphere (solid line) compared to ambient air
- 906 (dashed lines). (d) Comparison of field cooled (FC) and zero field cooled (ZFC) IRMs. (e) Low temperature
- magnetic susceptibility ($\chi_{If-MPMS}$) measured with five frequencies (colored lines). Frequency dependent
- 908 low temperature magnetic susceptibility (χ_{fd} ; black line).

909



- 911
- 912 Figure 7. JKC36 SEM images. (a-d) Examples of dissolution of iron bearing minerals and growth of
- 913 authigenic iron sulfides. (e-g) Examples of pyrite framboids. (h) Example of an iron sulfide mineral with
- 914 platy mineral structure that is tentatively interpreted as hexagonal 3C pyrrhotite. (i) EDS spectrum for
- 915 iron sulfide mineral displayed in (j-k). (j-k) Example of iron sulfide mineral with octahedral crystal
- 916 structure consistent with greigite.
- 917



918

919 Figure 8. Potential source rocks. (Left) Day plot (Day et al., 1977) for Polar Rock Repository (PRR)

samples from the Joinville Island region. The size of the circle is proportional to the M_s of the sample. All

921 Nordenskjöld Formation samples are from Northern Joinville Island, while the Joinville Island Diorite is from

southern Joinville Island (Figure 1). (Right) Selected $\chi_{\rm ff}$ measurements as a function of heating to ~700 °C

and cooling back to room temperature in an argon atmosphere. Samples not plotted have low χ_{lf} and

924 were dominated by paramagnetic contributions and significant alteration above 400 °C, but all data are

925 available in the MagIC Database (see Acknowledgements).

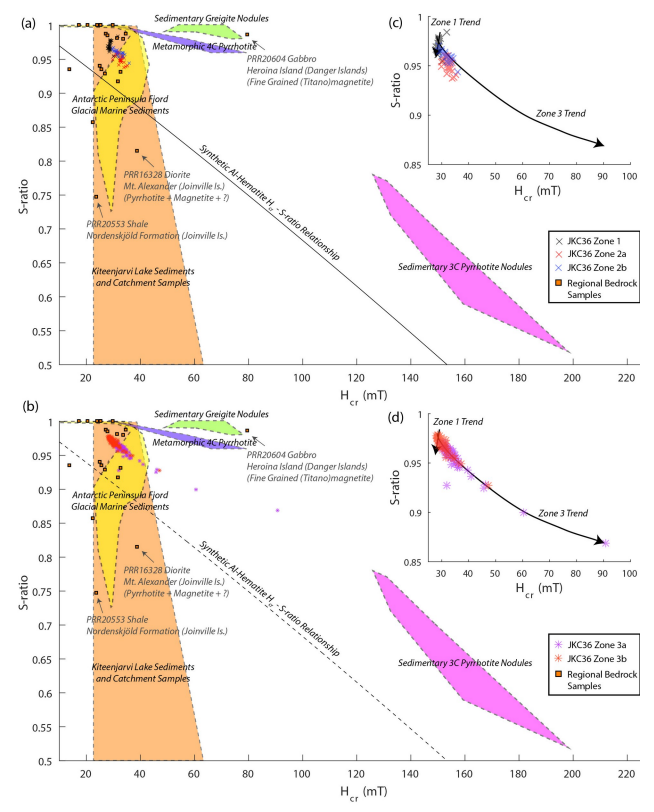
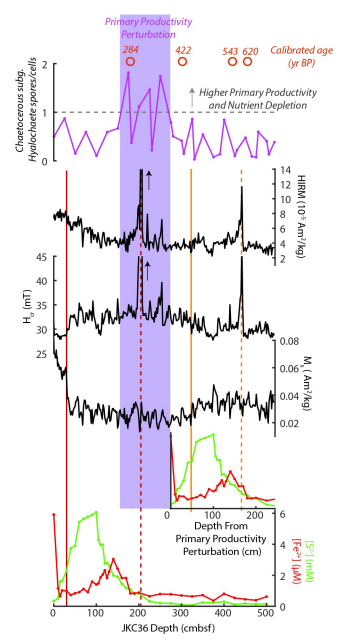


Figure 9. Comparison of Coercivity of Remanence (H_{cr}) and S-ratio. Core JKC36 H_{cr} and S-ratio values for
 (a) sulfidic sediments and (b) methanic sediments compared with regional bedrock samples (squares),

- values from glacial marine sediments in Barilari Bay, Western Antarctic Peninsula (yellow shading; Reilly
- et al., 2016), mixed hematite and magnetite magnetic mineral assemblages from Kiteenjarvi Lake and its
- catchment (orange shading; Stober and Thompson, 1979), and ranges for sedimentary hexagonal 3C
- pyrrhotite nodules (pink shading), sedimentary greigite nodules (green shading), and metamorphic rock
- monoclinic 4C pyrrhotite (purple shading) (Horng, 2018). The relationship between H_{cr} and S-ratio for
- synthetic Al-hematite samples is also plotted (black dashed line; Liu et al., 2007). **(c-d)** Expanded view of
- 936 the Core JKC36 data, annotated with trend lines for the Zone 1 and Zone 3 samples.





939 Figure 10. Comparison to Past Primary Productivity. From bottom to top: Porewater sulfide and iron 940 concentrations measured in Core JKC36; porewater data shifted 275 cm to represent what porewater 941 conditions may have looked like before a late Holocene primary productivity increase; M_s; H_{cr}; HIRM; ratio of Chaetocerous subg. Hyalochaete resting spores to cells (Kyrmanidou et al., 2018); and 942 943 radiocarbon dates (ΔR = 860 yrs; Kyrmanidou et al., 2018). A primary productivity perturbation based on 944 diatom data is highlighted in purple. The red solid vertical line indicates the modern dissolution front 945 while the red dashed vertical line indicates the modern base of the SMT where high concentrations of 946 remanence-bearing iron sulfides accumulate. Vertical orange solid and dashed lines indicate the hypothetical locations of the same two horizons about 400 cal. yrs ago prior to a perturbation caused by 947 948 increased primary productivity. Note the vertical orange solid line is at about the depth separating Zones

- 949 3a and 3b (Figure 5) and the vertical orange dashed line is where there is an enrichment of high
- 950 coercivity minerals at 442 cmbsf.

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