¹ Winter dissolved and particulate zinc in the Indian Sector

² of the Southern Ocean: Distribution and relation to major

³ nutrients (GEOTRACES GIpr07 transect)

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18 Abstract

19 First winter measurements of dissolved zinc (dZn) and particulate zinc (pZn) are presented from seven 20 stations, between 41 and 58°S, occupied in July 2017 along the 30°E longitude in the Indian Sector of the 21 Southern Ocean. This unique spatial and seasonal dataset provided the opportunity to investigate Zn 22 biogeochemical cycling in a region which is extremely data scarce and during a period when conditions are 23 unfavourable for phytoplankton growth. Surface comparisons of our winter dZn and pZn to previous 24 measurements during spring and summer revealed that Zn seasonality is most pronounced at the higher 25 latitudes where higher dZn (and higher ratios of dZn to phosphate; dZn:PO4) and lower pZn in winter 26 reflect decreased biological uptake and preferential dZn resupply (relative to PO4) to surface waters through 27 deep winter mixing. The composition of pZn was majorly biogenic however localised lithogenic inputs 28 were attributed to potential hydrothermal activity and transport of continental sediment via Agulhas waters. 29 Calculated vertical attenuation factors (b values) for pZn (0.31) and phosphorus (P; 0.41) suggest that Zn 30 has a longer remineralisation length scale than P, providing a mechanism as to why dZn appears to be 31 remineralised deeper in the water column than PO4. Ratios of pZn to P (pZn:P) in surface waters increased 32 with latitude from 1.12 to 8.28 mmol mol⁻¹ due to increased dZn availability and the dominance of diatoms 33 (with high cellular Zn quotas) in the high latitude Antarctic Zone (AAZ). Interestingly, the high surface 34 pZn:P ratios in the AAZ did not change significantly with depth (in contrast to the northern stations where pZn:P increased with depth) suggesting the export of diatom cells below the winter mixed layer where 35

36 remineralisation and rigorous mixing may resolve the linear dZn to silicic acid (dZn:Si(OH)₄) correlation

37 $(dZn (nmol kg^{-1}) = 0.064 Si(OH)_4 (\mu mol kg^{-1}) + 0.690; r^2 = 0.93; n = 120)$ despite these elements being 38 located in separate components of the diatom cell. Additionally, elevated concentrations of dZn and Si(OH)_4 39 below 3000 m in the AAZ may reflect nutrient accumulation in bottom waters where northward flow is 40 inhibited by the Indian mid-Ocean ridge.

41 1. Introduction

42 Zinc (Zn) is an essential trace metal micronutrient for marine phytoplankton and its availability in the 43 surface ocean has potential implications for biological productivity which, in turn, influences the 44 biogeochemistry of non-metals such as carbon and nitrogen (Morel et al., 1991; Morel and Price, 2003). 45 While Zn is a critical component in numerous enzymes involved in metabolism, perhaps the most important 46 are the metalloproteins carbonic anhydrase and alkaline phosphatase, which enable efficient carbon fixation 47 and the uptake of dissolved organic phosphorous, respectively (Shaked et al., 2006; Twining and Baines, 48 2013). However, high concentrations of dissolved Zn (dZn; > 10 nmol kg⁻¹), which can be found in estuarine and coastal upwelling settings, are known to be toxic to some phytoplankton (Sunda and 49 50 Huntsman, 1996). On the other hand, low dZn concentrations ($< 0.01 \text{ nmol kg}^{-1}$), typical of localised 51 oligotrophic settings, could be potentially bio-limiting (Brand et al., 1983).

52 Owing to an increase in reliable data in the last few decades, and particularly since the onset of the 53 GEOTRACES era (Anderson, 2020), our understanding of the processes controlling the exchange between 54 dissolved and particulate trace metal phases are better constrained. In the surface ocean, assimilation of dZn by phytoplankton is the primary process resulting in the exchange between inorganic dZn (< 0.2 µm pore 55 size filtered seawater) and organic particulate Zn (pZn; $> 0.45 \mu m$ pore size filter) phases. Subsequent 56 57 processes, e.g., zooplankton grazing, result in the formation of larger particles which either sink and 58 contribute to the vertical export of pZn, or disaggregate (via remineralisation, desorption and dissolution) 59 back to smaller particles contributing a source of dZn to the surface ocean (Lam and Marchal, 2015). In 60 addition, scavenging (or adsorption) of dZn onto organic particle surfaces, and the resulting formation of 61 larger and more rapidly sinking particulates phase, may enhance export of Zn to deeper waters (John and 62 Conway, 2014). Similarly, authigenic Zn sulphide precipitation has been suggested in low oxygen 63 environments with implications for pZn export in localised regions (Conway and John, 2015; Janssen and 64 Cullen, 2015). As a result of the interplay between uptake and regeneration processes, an increase in the 65 dZn pool typically leads to a decrease in the pZn pool. However, the exchange between dissolved and particulate phases is not necessarily proportional considering that biological and physical processes lead to 66 67 differences in their residence times, i.e., years to thousands of years for dissolved phases compared with 68 days to months for particulate phases (Lam and Marchal, 2015).

69 Comparing the distribution of Zn to those of the major nutrients, silicic acid $(Si(OH)_4)$ and phosphate (PO_4) ,

- 70 has proven a useful tool in constraining processes governing Zn distribution. For example, there is a tight
- 71 correlation between global dZn and Si(OH)₄ distributions with water column concentrations of both species
- characterised by surface depletion and enrichments at depth (Bruland, 1980; Ellwood, 2008; Janssen et al.,

73 2020; Middag et al., 2019; Wyatt et al., 2014; Zhao et al., 2014). The depths at which dZn and $Si(OH)_4$ 74 reach their greatest enrichment are deeper compared to the intermediate depth maxima displayed by global 75 PO₄ distributions (Bruland, 1980; Middag et al., 2018; Quay et al., 2015). The relationship between dZn, 76 $Si(OH)_4$ and PO₄ appears at odds considering that Zn and organic phosphorous (P) are associated with the 77 organic matter of phytoplankton cells (Twining et al., 2004, Twining et al., 2003), while a negligible amount 78 of Zn is incorporated into the siliceous frustules of diatoms (Ellwood and Hunter, 2000). It would therefore 79 be expected that dZn, like PO₄, is regenerated at shallower depths from the rapidly dissolvable organic 80 material and that there is no direct mechanism linking dZn and Si(OH)₄. Recently, two hypotheses have 81 sought to explain this phenomenon. The first suggests that the coupling of dZn and Si(OH)₄ is a 82 consequence of the natural interaction between ocean biogeochemistry and physical circulation through the 83 Southern Ocean hub (Vance et al., 2017). In sum, Antarctic Surface Waters (AASW) are preferentially 84 stripped of dZn and Si(OH)₄ relative to PO₄, consistent with the known uptake stoichiometry of diatoms in 85 this region. Physical circulation dictates that the dZn and Si(OH)₄ depleted surface waters are transported 86 to the low latitude thermocline thereby setting the low dZn, low Si(OH)₄ biogeochemical signature 87 throughout much of the upper ocean (Sarmiento et al., 2004). Furthermore, the export of diatom cells 88 beneath the winter mixed layer traps the majority of dZn and $Si(OH)_4$ in the deep ocean despite variances 89 in their regeneration length scales. The second hypothesis adopts a more mechanistic approach whereby a 90 secondary, deeper dZn source is inferred. The reversible scavenging model suggests that dZn is released at 91 shallow depths from decomposing organic particles at the same rate as PO₄, however dZn is then reversibly 92 scavenged (adsorbed) onto sinking organic particles resulting in its enhanced flux, relative to non-93 scavenged elements like PO₄, to the deep ocean (John and Conway, 2014; Weber et al., 2018).

94 Another aspect of Zn cycling requiring attention relates to seasonality. To date, the vast majority of Zn data 95 represent spring/summer seasons, when conditions are favourable for phytoplankton growth (e.g., increased light levels, shallow mixed layers). However, little is known about Zn cycling during winter months with 96 97 only two studies reporting winter Zn data for the Atlantic (Cloete et al., 2019) and Pacific (Ellwood, 2008) 98 Sectors of the Southern Ocean. Of particular interest is winter deep mixing, a seasonally constrained 99 physical supply mechanism whereby deeper mixing in winter compared to summer may tap into nutrient 100 rich subsurface water masses found below the summer stratified layer. Resupply of nutrients through deep 101 winter mixing may be critical in initiating and sustaining phytoplankton growth over the subsequent spring 102 and summer seasons, as shown previously for iron (Tagliabue et al., 2014). Furthermore, Subantarctic Mode 103 Water (SAMW), formed in the deep winter mixed layers of the Subantarctic Zone, is the primary 104 communication channel between the Southern Ocean and the global low latitude upper ocean (Broecker, 105 1991). Therefore, investigating Zn cycling in the source region for SAMW may have important and far 106 reaching implications as shown previously for Zn (Vance et al., 2017), Si(OH)₄ (Sarmiento et al., 2004) 107 and cadmium (Middag et al., 2018; Xie et al., 2015). This study aims to address these knowledge gaps by 108 contributing the first winter measurements of dZn and pZn from the Indian Sector of the Southern Ocean

109 (WOCE meridional section IO6S; GEOTRACES section GIpr07).

110 2. Methods

111 2.1. Sample collection

Seawater samples were collected on-board the SA Agulhas II polar research vessel during the 2017 Winter
 Cruise (28/06/2017–13/07/2017). The transect followed the World Ocean Circulation Experiment (WOCE)

- 114 I06S transect along the 30°E meridian in the Indian Sector of the Southern Ocean (Fig. 1A). Of the nine
- planned stations, seven, consisting of four deep (< 4500 m) and three shallow stations (> 1500 m), were
- sampled between 41°00'S and 58°30'S allowing observations to be investigated over important frontal
- 117 systems and in different water masses. Two stations (IO03 and IO09) were cancelled for trace metal
- 118 sampling due to intense storms present at the time of station occupation.
- 119 A vertical profile sampling method, using 24 internally Teflon-coated polyvinyl chloride (PVC) 12 L GO-
- 120 FLO sampling bottles (General Oceanics), was employed at all sampling stations. GO-FLO bottles were
- 121 mounted on a GEOTRACES compliant rosette frame (Cutter et al., 2017) housing a Seabird 9+ CTD
- 122 (conductivity, temperature and depth) recorder. A Kevlar hydrowire with internal signal cables allowed for
- 123 the transfer of data between the CTD and the on-board control room. The GO-FLO bottles were triggered 124 at pre-determined depths during the up cast. Directly upon recovery of the rosette, the GO-FLO bottles
- 125 were covered in a PVC plastic wrap in addition to their ends being covered in PVC shower caps, and were
- 126 transported into a class 100 clean lab for sub-sampling. Samples for dissolved trace metal determination
- 127 were collected in 125 mL acid-cleaned low density polyethylene (LDPE, Nalgene) bottles after online
- 128 filtration through 0.2 μm Sartobran filters and under slight nitrogen (N₂, 99.9999% purity, BIP technology)
- 129 overpressure. Samples were acidified (pH = 1.7) on-board under a laminar flow hood using hydrochloric
- 130 acid (Ultrapur® HCl, Merck) and stored for later analysis. Thereafter, previously cleaned filters (25 mm
- 131 diameter, Supor, 0.45 µm pore size) were mounted on acid-washed Swinnex (Millipore) filter holders which
- 132 were then attached to the GO-FLO spigots and used for filtration (recorded between 5 and 10 L) of samples
- 133 for particulate trace metal determination (Planquette and Sherrell, 2012). After filtration, filter holders were
- 134 removed and disassembled before the sampled filter was transferred to an acid-washed polystyrene Petri-
- 135 dish using clean plastic forceps. Sampled filters were subsequently frozen at -20 °C for transport back to
- 136 land. All handling of filters, filter holders and samples was done under a laminar flow hood.

137 2.2. Dissolved zinc (dZn) determination

138 2.2.1. sea*FAST* and ICP-MS

Seawater samples were first preconcentrated offline by Solid Phase Extraction (SPE) on a seaFAST SC-4 DX module (Jackson et al., 2018; Rapp et al., 2017) prior to quantification by ICP-MS (Agilent 7900). During preconcentration, 10 mL of seawater was buffered to a pH of 6.0 ± 0.2 with an ammonium acetate buffer. The buffered solution was then loaded onto a column containing a high affinity metal chelating resin (Nobias PA1) where the Zn ions were bound to the resin and separated from the seawater matrix elements, e.g., Sodium (Na), Magnesium (Mg) and chlorine (Cl) which passed through the column. The Zn ions were subsequently eluted from the resin column in low volumes (250 µL) resulting in a preconcentration factor

- of 40. After preconcentration, the analyte was introduced into the ICP-MS using a low self-aspirating
- 147 perfluoroalkoxy (PFA) nebulizer with a flow rate of 0.2 mL min⁻¹. Isotopes of 66Zn were measured using
- 148 the instruments Octopole Reaction System (ORS) in helium (He) collision mode to eliminate plasma and
- 149 matrix based interferences although the latter was extensively reduced by the seaFAST matrix removal
- 150 system. Online internal standard addition for drift correction was not possible using the self-aspirating
- 151 nebulizer. Instead, instrument drift was monitored by running a multi-element standard (MES; verified by
- 152 Inorganic Ventures) every 6 samples. Where drift exceeded 5% relative to the starting concentration of the
- 153 MES for a specific element, a drift correction was applied using Eq. (1)
- 154 $2*Conc_{MES_start}/(Conc_{MES_a}+Conc_{MES_b})*Conc_{sample}$ Eq.

1

- 155 where *a* and *b* are the MES before and after each set of 6 samples.
- 156 2.2.2. Accuracy and precision

157 The accuracy of the dZn analyses was verified by way of comparison with GEOTRACES reference 158 seawater and NASS-7 certified reference material (Table 1). For the GEOTRACES SAFe D2, GSC and 159 GSP reference seawater, our dZn values were within analytical uncertainty confirming the method's 160 accuracy over oceanographically relevant concentrations. For the NASS-7, our dZn value was in good agreement with the certified value. To monitor ICP-MS precision, internal reference seawater (0.2 µm 161 162 filtered) was collected from 55°S; 28°E during our transect (Winter Indian Southern Ocean Seawater; WISOS). The WISOS internal reference seawater was placed in the analysis sequence and results compared 163 against each other and the calibrated mean (Table 1). The calibrated mean concentration $(9.67 \pm 0.23 \text{ nmol})$ 164 kg^{-1}) was established by replicate analysis (n = 10) of the WISOS in conjunction with the SAFe and NASS-165 7 reference materials. The in-sequence analysis of the WISOS (n > 30) yielded a value of 9.63 \pm 0.24 nmol 166

- 167 kg^{-1} confirming the method's precision.
- 168 2.2.3. Blanks and limits of detection
- 169 The instrument (ICP-MS) blank was quantified by introducing a solution of un-preconcentrated 2% HNO₃,
- identical to the seaFAST eluent, in ultra-pure deionized water. The instrument blank was 0.07 ± 0.01 nmol
- 171 kg^{-1} for Zn (Table 1). To determine the blank contribution from the method, a solution of HNO₃ (ultrapur®,
- 172 Merck) diluted to 2% with ultra-pure deionized water, the same composition as the eluent used in sample
- 173 preconcentration, was analysed. The blank was subjected to the same preconcentration procedure as the
- seawater samples. The method blank was 0.09 ± 0.01 nmol kg⁻¹ and was subtracted from sample values.
- 175 The limit of detection (LOD), calculated as three times the standard deviation of the preconcentrated blank,
- 176 was $0.02 \text{ nmol kg}^{-1}$.
- 177 2.2.4. Data processing

178 Samples for dZn were analysed in duplicate. The final value represents the mean of duplicate

- measurements. In cases where the percentage relative standard deviation (%RSD) between the duplicate measurements was > 10% (n = 13), one of the values was deemed a suspected outlier and not used further.
- 181 This was determined by curve fitting the data points based on the values above (shallower depth) and below

- 182 (deeper depth) the suspect value as well as by comparison with other parameters (salinity, temperature, and
- 183 nutrients) measured from the same GO-FLO bottle.

184 2.3. Phosphate (PO₄) and silicic acid (Si(OH)₄) determination

185 Macronutrient concentrations were described previously (Weir et al., 2020). Here, we briefly summarise

the analytical methods. Seawater samples for PO_4 and $Si(OH)_4$ analysis were collected from GO-FLO bottles, filtered immediately after collection using a 0.2 µm pore size syringe filter into 50 mL Falcon®

tubes and frozen at -20 °C. Macronutrient analysis was done at the Marine Biogeochemistry Laboratory at

189 the University of Cape Town (MBL-UCT). PO₄ was determined manually by colorimetric method

190 (Grasshoff, 1983) with an analytical error of $\pm 0.06 \,\mu\text{mol kg}^{-1}$. Si(OH)₄ was analysed using a Lachat Quick-

191 Chem flow injection analyser (Wolters, 2002) with an analytical error of $\pm 0.2 \,\mu\text{mol kg}^{-1}$.

192 2.4. Particulate Zn (pZn) and phosphorous (P) determination

Samples for particulate trace metals were analysed at the Université de Bretagne Occidentale (UBO).
Details of the analytical procedure have been previously described (Planquette and Sherrell, 2012) and are

195 presented briefly here. The pZn samples were processed and analysed 6 months after sample collection.

196 Filters containing the samples were acid reflux digested at 130 °C in acid-cleaned Savillex vial. Archive

197 solutions were stored in 3 mL of 0.12 M HNO₃ (Ultrapur[®] grade), of which 250 μL was diluted up to 2 mL

198 for analysis by sector field inductively coupled plasma mass spectrometry (SF-ICP-MS, Element XR

199 Thermo Scientific). Samples were spiked with 1 μ g L⁻¹ indium (In) as an internal standard to correct for

- 200 instrument drift. Three certified reference materials (PACS 3, MESS 4 and BCR 414) were processed as
- samples and analysed for pZn and P to assess the accuracy of the methodology (Table 2). The certified
- reference materials yielded mean percentage recoveries of 108%, 100% and 97% for pZn, and 100%, 106%
- and 95% for P (PACS 3, MESS 4 and BCR 414, respectively). Filter blanks (1.51 pM and 0.08 nM for pZn
- and P, respectively) were determined by digesting and analysing an acid-washed filter. The detection limits,
- 205 defined as three times the standard deviation of the blanks, for pZn and P were 2.82 pmol kg^{-1} and 0.03

206 nmol kg⁻¹, respectively (n = 5).

207 2.5. Estimating pZn composition

The lithogenic fraction of the total pZn (pZn_{lith}) was calculated by multiplying the particulate aluminium (pAl) concentration (Van Horsten et al., 2021) with the Zn/Al upper continental crust (UCC) ratio of 0.00163 (Rudnick and Gao, 2013). The non-lithogenic fraction (assumed to be the biogenic fraction) of the

- 210 0.00103 (Rudnick and Gao, 2013). The non-inflogenic fraction (assumed to be the biogenic fraction) of the
- 211 total pZn (pZn_{bio}) was then calculated by subtracting the lithogenic fraction of pZn from the total pZn (Eq.
- 212 (2)). Unless otherwise stated, pZn in this manuscript refers to pZn_{bio} .
- 213 $pZn_{bio} = [pZn] ([pAl] \times (Zn/Al)_{UCC})$

Eq. 2

214 2.6. Calculation of vertical attenuation factors (*b* values)

215 The remineralisation length scale of an element is determined by the attenuation of the downward particle

216 flux settling gravitationally and is element specific (Boyd et al., 2017). The attenuation of Zn with depth

217 was modelled using the Martin equation (Martin et al., 1987), originally developed for particulate organic

- 218 carbon, modified for Zn (Eq. (3)), P and cadmium accordingly (Ellwood et al., 2020). Particulate cadmium
- 219 data are from Cloete et al. (2021).

220
$$[pZn_z] = [pZn_{MLD}](z/MLD)^{-b}$$
 Eq. 3

222 Where $[pZn_z]$ is the pZn concentration at depth z, $[pZn_{MLD}]$ is the pZn concentration nearest the base of the

223 Mixed Layer Depth (MLD) at each station and b is the vertical attenuation factor. Model fits of the

- 224 particulate data were optimised by minimising the square of the differences between observational data and
- the model output.

221

226 2.7. Extended optimum multiparameter (eOMP) analysis

227 An extended optimum multiparameter (eOMP) analysis (Karstensen and Tomczak, 1998) was used to solve 228 the water mass structure of the section. Briefly, eOMP analyses consider that the observed physicochemical 229 properties of a given water sample can be formulated as a linear combination of a finite number of water 230 masses represented by the so-called source water types (SWTs; Tomczak, 1999). The SWTs are points in the n-dimensional parameter space, where n is the number of properties that characterize SWTs. In this 231 work, the SWTs are characterised by potential temperature, salinity, oxygen, nitrate, PO_4 and $Si(OH)_4$ 232 233 (Table 3). Once the SWTs and their physicochemical properties are defined, the goal of an eOMP analysis 234 is to find the fractions of each SWT (Xi) in each water sample and does so by a least-square method 235 constrained to be positive definite:

236
$$d = G * X_i + \Delta O_{2bio}/R + \epsilon$$
 Eq. 4

where d is the observed property in a water sample, G is the matrix containing the properties defining the SWTs, X_i is the relative contributions of each SWT to the sample, ΔO_{2bio} accounts for the changes in oxygen due to the synthesis and/or remineralisation of the organic matter, R are the Redfield-like stoichiometric ratios, and ε is the residual. In this study we used an R of 12 for Si(OH)₄ (Castro et al., 1998; Perez et al., 1993), 175 for PO₄ and 10.5 for nitrate (Anderson and Sarmiento, 1994; Takahashi et al., 1985).

The eOMP analysis was used to locate the realm of Antarctic Bottom Water (AABW), and therefore the eOMP analysis was restricted to water samples > 1000 m. For this purpose, only four SWTs were selected: Upper Circumpolar Deep Water (UCDW), Lower Circumpolar Deep Water (LCDW), North Atlantic Deep Water (NADW), and AABW. In this work, we selected the properties for UCDW from Carter et al. (2014); and those of LCDW, NADW, and AABW from Liu and Tanhua (2021). The eOMP analysis was reliable since it explained 99% of the changes in the conservative tracers (Table 3).

248 3. Results

249 3.1. Description of the study area

250 3.1.1. Frontal positions

251 The Southern Ocean is dominated by the uninterrupted, eastward flowing Antarctic Circumpolar Current 252 (ACC). Another major current, the Agulhas current (AC) flows southward along the east coast Africa before 253 retroflecting at the southern tip of the continent due to shear interactions with the ACC. The resulting 254 Agulhas Return Current (ARC) flows eastward and intersects the 30°E section at ~40°S (Barrett et al., 255 2018). A number of important oceanographic fronts run parallel with the ACC and divide the Southern Ocean into distinct biogeochemical zones (Fig. 1A). Frontal positions were determined previously (Weir et 256 257 al., 2020) following temperature, salinity and oxygen criteria (Belkin and Gordon, 1996; Orsi et al., 1995; 258 Pollard et al., 2002). To the north of the transect, the Subtropical Front (STF) at 42.4°S separated the 259 Subtropical Zone (STZ) to the north from the Subantarctic Zone (SAZ) to the south. Likewise, the Subantarctic Front (SAF) at 46.2°S separated the SAZ in the north from the Polar Frontal Zone (PFZ) to 260 261 the south which, in turn, was bounded in the south by the Antarctic Polar front (APF) at 49.3°S. Further 262 south, the Antarctic Zone (AAZ) extended until the Southern Boundary (SBdy) at 58.5°S, incorporating the 263 Southern Antarctic Circumpolar Current Front (SACCF) at 56.5°S. The Marginal Ice Zone (MIZ), defined as 30% ice cover, was encountered at 61.7°S (de Jong et al., 2018), south of the southernmost sample 264 265 station.

266 3.1.2. Water mass characterisation

267 Numerous water masses were sampled throughout the study (Fig. 1B; water mass definitions found in 268 Anilkumar et al., 2006; Orsi et al., 1995). The hydrography of the Southern Ocean is largely defined by the 269 upwelling and ventilation of deep waters at the higher latitudes. As a consequence of wind-driven forcing, 270 nutrient-rich UCDW, identified at a potential temperature (θ) of 2.0–2.5 °C and potential density (σ_{θ}) of 27.2–27.8 kg m⁻³, upwells south of the APF. Near the surface, a portion of the upwelled water moves south, 271 as AASW ($\theta \le 2.5$ °C), where continued cooling and sea-ice formation increase water density initiating 272 subduction and eventual formation of cold ($\theta = -0$ °C; $\sigma_{\theta} = 27.8-27.9$ kg m⁻³) AABW, which flows 273 274 northward along the basin floor (Orsi et al., 1999). In the upwelling region, the remaining portion of 275 upwelled UCDW flows northward via Ekman drift. Here, surface waters warm rapidly toward the north forming Subantarctic Surface Water (SASW; $\theta = 7-12$ °C) and Subtropical Surface Water (STSW; $\theta \ge 12$ 276 277 °C). Subducting northward below the surface water masses were Antarctic Intermediate Water (AAIW; θ = 2.5–5.0 °C, σ_{θ} = 26.7–27.4 kg m⁻³) and SAMW (θ = 12–15 °C, σ_{θ} = 26.2–26.6 kg m⁻³). Intermediate and 278 279 mode waters are the chief communicators between the major ocean basins exporting their biogeochemical 280 signatures to the lower latitude oceans (Gordon et al., 1992). Deeper in the water column, NADW, identified chiefly by a deep salinity maximum (S > 34.80; van Aken et al., 2004) was generally characterised by lower 281 concentrations of all major nutrients in relation to the overlying UCDW and underlying LCDW ($\theta = 1.5$ -282 283 2.0 °C; $\sigma_{\theta} = 27.8 - 27.9 \text{ kg m}^{-3}$).

- 284 3.1.3. Surface mixed layer and remineralisation depth
- 285 The Surface Mixed Layer (SML) at each station was defined as the area between the sea surface and the
- 286 MLD, which was determined as the depth between 10 m and 400 m at which the Brunt Väisälä frequency
- (BVF) squared reach a maximum (Weir et al., 2020). Along this transect, the STZ had the shallowest MLD
- 288 (122 m) while the SAZ was characterised by the deepest MLDs observed (190–220 m). Further south, the
- MLD generally increased from 134 m in the PFZ to 125–161 m in the AAZ. The remineralisation depth
- 290 was approximated at 500 m for all stations based on the observation that at this depth, vertical profiles of P
- 291 (and pZn) converged on low values (Fig. S1), representative of the deep water values.
- 3.2. Distribution of dissolved elements
- 293 3.2.1. dZn
- 294 At each station, dZn (Figs. 2A, S1 and S2) was lowest in surface waters and increased to local maxima in the deepest sample. The lowest measured dZn was in STSW (0.44 \pm 0.28 nmol kg⁻¹) while highest 295 concentrations were observed in AABW (8.67 ± 0.42 nmol kg⁻¹), sampled south of the Indian ridge (located 296 297 broadly between 48°S and 52°S) below 3000 m depth. Surface water dZn concentrations increased 298 southwards by approximately 8-fold between STSW in the north and AASW $(3.37 \pm 0.28 \text{ nmol kg}^{-1})$ to the south. Concentrations of dZn in AAIW was 3.67 ± 1.79 nmol kg⁻¹ while dZn in UCDW was 5.89 ± 1.06 299 nmol kg⁻¹. In general, NADW was characterised by slightly lower dZn concentrations relative to 300 301 surrounding water masses. For example, in the PFZ (St. IO05; 48° S), dZn in NADW was 4.91 ± 0.20 nmol 302 kg^{-1} , 5.14 ± 0.89 nmol kg^{-1} in the overlying UCDW and 6.62 ± 0.03 nmol kg^{-1} in the underlying LCDW. In deep waters (> 2500 m) north of the Indian ridge (St. IO05 and IO06), dZn was 5.71 ± 1.18 nmol kg⁻¹ 303
- 304 while to the south (St. IO04 and IO02) dZn was 8.51 ± 0.37 nmol kg⁻¹.
- 305 3.2.2. PO₄ and Si(OH)₄
- 306 Concentrations of PO_4 and $Si(OH)_4$ (Fig. S1) have been previously reported (Weir et al., 2020) and are
- briefly described here. Surface (25 m) PO₄ increased southward from 0.22 to 1.78 μ mol kg⁻¹ while Si(OH)₄ increased from 2.93 to 46.8 μ mol kg⁻¹. At each station PO₄ increased with depth until a mid-depth maximum
- 309 (between 2.20 and 2.67 μ mol kg⁻¹ at 41 and 48°S, respectively) located within the southward upwelling
- 310 UCDW. In contrast to PO₄, vertical profiles of Si(OH)₄ increased to maximum concentrations at the deepest
- 311 depth sampled (like dZn) and reached a maximum concentration of 132 μ mol kg⁻¹ in AABW (3500 m) at
- 312 50°S.

313 3.3. Distribution of particulate elements

- 314 3.3.1. pZn
- 315 Depth profiles of pZn (Figs. 2B, S1 and S2) typically had highest concentrations near the surface,
- 316 particularly in the AAZ where surface water pZn was 0.106 ± 0.029 nmol kg⁻¹. To the north, surface water
- $317 \qquad pZn \ was \ 0.068 \pm 0.009 \ nmol \ kg^{-1} \ in \ the \ PFZ, \ 0.037 \pm 0.012 \ nmol \ kg^{-1} \ in \ the \ SAZ \ and \ 0.021 \pm 0.006 \ nmol \ nmol \ kg^{-1} \ in \ the \ SAZ \ and \ 0.021 \pm 0.006 \ nmol \ nmol \ kg^{-1} \ in \ the \ SAZ \ and \ 0.021 \pm 0.006 \ nmol \ kg^{-1} \ in \ the \ SAZ \ and \ 0.021 \pm 0.006 \ nmol \ kg^{-1} \ in \ the \ SAZ \ and \ 0.021 \pm 0.006 \ nmol \ kg^{-1} \ in \ the \ SAZ \ and \ 0.021 \pm 0.006 \ nmol \ kg^{-1} \ in \ the \ SAZ \ and \ 0.021 \pm 0.006 \ nmol \ kg^{-1} \ in \ the \ SAZ \ and \ 0.021 \pm 0.006 \ nmol \ kg^{-1} \$
- 318 kg⁻¹ in the STZ. There was no clear trend in intermediate and deep water pZn distributions. Below 1000 m
- depth, pZn was 0.033 ± 0.016 nmol kg⁻¹ south of the APF and 0.022 ± 0.011 nmol kg⁻¹ north of the APF.

- 320 The composition of pZn was dominated by pZn_{bio} (pZn (nmol kg⁻¹) = 0.960 pZn_{bio} (nmol kg⁻¹) + 0.004; r²
- 321 = 0.99; n = 120). However, there were localised areas where pZn_{lith} contributed significantly to pZn (Fig.
- 322 2C). For example, below the surface layer (> 250 m) in the STZ, pZn_{lith} was 0.007 \pm 0.001 nmol kg⁻¹
- equating to between 16 and 68% of total pZn. In the SAZ, pZn_{lith} was 0.004 ± 0.001 nmol kg⁻¹ below the
- surface layer (10–45% of pZn). In the AAZ, pZn_{lith} was low (0.002 \pm 0.001 nmol kg⁻¹) throughout the water
- 325 column with the exception at the deepest sample at 50° S, above the mid-ocean ridge, where pZn_{lith} was 0.01
- 326 nmol kg⁻¹ (30% of pZn).

327 3.3.2. P

The distribution of P (Fig. S1) was characterised by high surface concentrations, rapid decreases through subsurface water masses and low (< 2.5 nmol kg⁻¹) concentrations below 1000 m depth. Highest surface water (25 m) P concentrations were measured in the PFZ (27.5 nmol kg⁻¹) and SAZ (15.9–24.7 nmol kg⁻¹) and were lower in the AAZ (9.43–17.0 nmol kg⁻¹) and STZ (15.4 nmol kg⁻¹).

332 4. Discussion

4.1. Seasonal insights into Zn cycling

The data presented here contributes the first winter measurements of dZn and pZn from the 30°E longitude 334 in the Indian Sector of the Southern Ocean. For comparative purposes, a global compilation of dZn and pZn 335 measurements (Fig. S3) was created from available data in the GEOTRACES Intermediate Data Product 336 337 (IDP) 2017 (Schlitzer et al., 2018; see figure caption for dataset references). While the compilation demonstrates the usefulness of the GEOTRACES IDP, we focus our comparison on the Southern Ocean 338 339 where we compared our surface dZn (Fig. 3A) and pZn (Fig. 3B) to a compilation of measurements 340 (including data not in the IDP) covering all three Southern Ocean Sectors and multiple seasons in order to 341 assess seasonal variations and identify potential drivers thereof (Barrett et al., 2018; Butler et al., 2013; 342 Cloete et al., 2019; Coale et al., 2005; Croot et al., 2011; Ellwood, 2008; Ellwood et al., 2020; Janssen et 343 al., 2020; Löscher, 1999; Sieber et al., 2020; Wang et al., 2018). We acknowledge the limitations of such a 344 comparison where, for example, shifts in frontal positions, changes in ambient nutrient availability and 345 differences in the resident phytoplankton community between sample locations could potentially influence Zn distributions. Nevertheless, the comparisons do yield interesting observations regarding Zn seasonality. 346 There is a fairly consistent latitudinal trend for surface dZn whereby concentrations were lowest north of 347 ~46°S, increased southward until maxima in the upwelling zone, between 52°S and 56°S, and were 348 349 generally lower in surface waters south of 56°S. Seasonal dZn differences were least apparent in the region north of 46°S where our winter Indian Sector dZn (0.46 ± 0.16 nmol kg⁻¹; n = 3) was comparable to winter 350 $(0.33 \pm 0.16 \text{ nmol kg}^{-1}; n = 3)$ and summer Pacific Sector data $(0.34 \pm 0.23 \text{ nmol kg}^{-1}; n = 3)$. Interestingly, 351 winter dZn in the Pacific Sector remained low $(0.28 \pm 0.11 \text{ nmol kg}^{-1}; n = 6)$ between 46°S and 52°S while 352 353 dZn increased significantly during our transect (and for all other datasets). The low dZn in the Pacific Sector coincided with low Si(OH)₄ ($< 3.49 \mu$ mol kg⁻¹) which extended until the SAF at $\sim 52^{\circ}$ S, much further south 354 compared to the position of the SAF during this study (~46°S), illustrating the impact of frontal positions 355

356 and nutrient availability on dZn distributions. Further south, in the upwelling zone, seasonal dZn differences were more apparent. For example, at 56°S, our winter Indian Sector dZn value (4.19 nmol kg⁻¹) was slightly 357 higher than the spring Atlantic Sector (3.80 nmol kg⁻¹) and significantly higher compared to summer 358 359 measurements in the Atlantic (2.81 nmol kg⁻¹) and Pacific (0.11 nmol kg⁻¹) Sectors. The same trend was evident for ratios of dZn to PO_4 (dZn:PO₄; Fig. S4) at 56°S whereby ratios were highest in winter (2.17) 360 mmol mol⁻¹) and decreased in spring (2.02 mmol mol⁻¹) and summer (1.73 and 0.65 mmol mol⁻¹; Atlantic 361 362 and Pacific Sectors, respectively). As a result of unfavourable growth conditions for phytoplankton during winter (e.g., lower light levels and deeper mixed layers than in spring/summer), the higher dZn (and 363 364 dZn:PO₄) in winter likely reflects decreased biological uptake by diatoms as well as deep winter mixing 365 whereby dZn is preferentially resupplied (relative to PO_4) to surface waters through access to dZn-enriched waters below the spring/summer mixed layers. The source of excess Zn in subsurface waters is 366 predominantly a water mass characteristic reflecting preformed concentrations and non-local dZn 367 368 accumulation from remineralisation throughout the ocean. In an attempt to assess the relative importance 369 of biological uptake and deep winter mixing in driving the dZn seasonal signal, we calculated the theoretical 370 dZn uptake (at three locations; in the SAZ, PFZ and AAZ) based on the observed PO₄ seasonal signal i.e. 371 the difference between summer (Sieber et al., 2020) and winter (this study) concentrations averaged over 372 the mixed layer, and the summer dZn/PO₄ uptake ratio (calculated from the slope of regression between 373 dZn and PO₄ in the winter mixed layer). We then compared the theoretical dZn uptake value to the measured 374 dZn seasonal signal (as calculated for PO₄) assuming the difference to reflect additional dZn supply from a 375 combination of deep winter mixing, lateral advection or remnants from the previous growth season. We calculate an expected dZn uptake of 0.22, 0.24 and 0.57 nmol kg⁻¹ in the SAZ, PFZ and AAZ respectively 376 377 (Table S1), compared to measured winter-summer dZn differences of 0.79, 0.58 and 0.87 nmol kg^{-1} . Ultimately results suggest that, at high latitudes (AAZ), differences in biological uptake predominantly 378 379 drive the seasonal dZn signal while at lower latitudes (PFZ and SAZ), deep winter mixing, lateral advection 380 and remnant dZn signatures become progressively more important.

- 381 For pZn (Fig. 3B), comparable data was more scarce than for dZn. North of 50°S, our winter surface pZn
- 382 $(0.05 \pm 0.04 \text{ nmol kg}^{-1})$ was comparable to corresponding summer measurements $(0.09 \pm 0.02 \text{ nmol kg}^{-1})$;
- 383 n = 2). Between 50°S and 58°S (the southern extent of our transect) our winter pZn remained low, below
- 384 0.12 nmol kg⁻¹, while summer pZn increased to 0.28 ± 0.12 nmol kg⁻¹ reflecting the increased uptake of
- 385 Zn under favourable growth conditions. South of 58°S, summer pZn remained high (0.48 ± 0.22 nmol kg⁻¹;
- 386 n = 7) while spring pZn was lower (0.21 ± 0.12 nmol kg⁻¹; n = 5) yet still higher than winter pZn slightly
- 387 to the north. A southward increasing trend was also evident in ratios of pZn to P (pZn:P; Fig. S4) although
- 388 the magnitude of the pZn:P increase varied between seasons as well as location suggesting Zn uptake is not
- fixed to that of P.

390 4.2. Factors controlling Zn uptake in surface waters

391 The phytoplankton community present during this transect was active with winter biomass accumulating 392 to levels similar to the lower end of comparable summer measurements (Weir et al., 2020). The uptake of 393 Zn by phytoplankton in surface waters is related primarily to the availability of Zn (Middag et al., 2019; 394 Morel et al., 1994; Sunda and Huntsman, 1995) but may also be influenced by the availability of other 395 metals such as iron (Fe). Interestingly, both increases (Twining et al., 2004) and decreases (Cullen et al., 2003) in the pZn:P ratio of resident phytoplankton communities have been observed following Fe addition, 396 397 suggesting varied cellular responses to Fe-stimulated growth. In an effort to investigate the drivers of Zn 398 uptake during this study, we compare dZn, pZn and pZn:P averaged over the SML (Fig. 4). In addition, 399 Si(OH)₄ and dissolved Fe (dFe; see Table S2 for data validation) are also shown. There was a strong southward increase in dZn concentrations, which increased from 0.20 nmol kg^{-1} in the STZ to a maximum 400 of 3.66 nmol kg⁻¹ in the AAZ, driven by upwelling of deep waters enriched in Zn and other nutrients, e.g., 401 Si(OH)₄ (Vance et al., 2017). Similarly, P and pZn:P ratios increased southward. For example, pZn:P 402 403 increased from 0.97 mmol mol⁻¹ in the STZ to a maximum of 8.58 mmol mol⁻¹ in the AAZ. In contrast, dFe concentrations were > 1 nmol kg⁻¹ north of the APF and were lowest in the AAZ (0.04 \pm 0.01 nmol 404 kg^{-1}). 405

The clear association between dZn and pZn with latitude suggests Zn availability is the primary driver of 406 407 Zn uptake. However it must be noted that Zn uptake rates are related to the free Zn concentration rather than the total Zn concentration (Baars and Croot, 2011; Brand et al., 1983; Saito and Goepfert, 2008). 408 409 Organic complexation may potentially reduce biological uptake in surface waters of the STZ and SAZ where dZn is low and the concentration of Zn-binding ligands may saturate dZn (Baars and Croot, 2011). 410 411 In the PFZ and AAZ however, the high surface dZn saturates ligand concentrations equating to high 412 bioavailable dZn. Our pZn:P ratios (7.68 \pm 1.14 mmol mol⁻¹) from the diatom dominated AAZ (Fig. S5) were higher compared to previous measurements along this transect (3.00 \pm 0.90 mmol mol⁻¹; Barrett et 413 414 al., 2018), where a more diverse phytoplankton assemblage was likely present, and compared more similarly to diatom dominated assemblages from elsewhere in the Southern Ocean (8.10-14.90 mmol 415 416 mol⁻¹; Barrett et al., 2018; Cullen et al., 2003; Twining et al., 2004). Diatoms require Si(OH)₄ to build their siliceous frustules and the high Si(OH)₄ in the AAZ (26–48 μ mol kg⁻¹), compared to the northern zones (2– 417 5 μ mol kg⁻¹), allowed diatoms to dominate the phytoplankton community assemblage. While Si(OH)₄ was 418 highest in the AAZ, dFe was lowest and likely limited diatom growth. There is evidence that suggests 419 420 cellular pZn:P ratios are influenced by Fe availability and our data indicate higher pZn:P under low Fe 421 conditions, possibly via reduced growth rates and accumulation of Zn (and other metals) relative to organic 422 biomass (Cullen et al., 2003). The low Si(OH)₄ in surface waters close to and north of the APF allowed 423 flagellates to out compete diatoms (Weir et al., 2020). The pZn:P of flagellates in the Southern Ocean does 424 show latitudinal variation likely linked to Zn availability (i.e., higher pZn:P in the high dZn waters south 425 of the APF) however, the pZn:P of flagellates was at least four times less than co-occurring diatoms (Twining and Baines, 2013) suggesting flagellates have lower P normalised Zn quotas and providing further
reasoning behind the low pZn:P at the northern stations. Therefore, in addition to the primary control
exerted by Zn availability, the availability of dFe and changes in phytoplankton taxonomy may influence
pZn:P ratios in surface waters.

4.3. Differences in remineralisation length scales between zinc, phosphorus andcadmium

Vertical attenuation factors (b values) for pZn, P and particulate cadmium (pCd) were calculated for full 432 433 depth profiles and to 1000 m (Table 4) in order to investigate the relative remineralisation length scales for these elements. For the full depth profiles, b values were slightly higher than for < 1000 m with the 434 435 exception of pZn at 41°S and 43°S where the exclusion of high pZn concentrations (0.042 and 0.048 nmol kg^{-1} , respectively) measured at 1500 m caused increased b values. For simplicity, we focus on b values for 436 the full dataset. For pZn, a b value of 0.33 was obtained however there was significant latitudinal variation 437 438 between zones with lower values in the STZ and SAZ (0.18–0.24) and higher values in the PFZ and AAZ 439 (0.27–0.50). Importantly, at all stations the b value for pZn was lower than that of P and pCd. Overall, our 440 b values for pZn, P (0.41) and pCd (0.46) suggest a longer (slower) remineralisation length scale for pZn 441 compared to P and pCd, in agreement with the sequence of Ellwood et al. (2020). Scaled to P, our b value 442 for pZn is higher compared to Ellwood et al. (2020) and compares more similarly to Boyd et al. (2017). 443 The differences in remineralisation length scales for these elements are likely related to their specific 444 physiological association within the cell. For example, Zn is a constituent of a host of metalloproteins (Zn-445 finger proteins) such as carbonic anhydrase, alkaline phosphatase, and RNA polymerase (Twining and Baines, 2013) whereas P is located in ATP, DNA, RNA and phospholipid membranes (Raven, 2013 and 446 447 references therein) ultimately leading to differences in lability between Zn and P. As remineralisation 448 reflects the release of these elements from sinking organic matter (i.e., from particulate to dissolved phase), longer length scales translate to a deeper release and should be reflected in the depth profiles of pZn to P 449 450 (pZn:P) and dZn to major nutrients (dZn:PO₄ and dZn:Si(OH)₄).

451 4.3.1 pZn:P

452 Depth profiles of pZn:P generally showed increases with depth, particularly in the upper water column (Fig. 453 5). The increase was most pronounced in the STZ (Fig. 5A) where pZn:P was low at the surface (0.89 mmol 454 mol⁻¹) and increased more than 5-fold by 500 m. This factor was between 4 and 5 in the SAZ (Fig. 5B–C), 3 in the PFZ (Fig. 5D) and approximately 1 throughout the AAZ (Fig. 5E-G) indicating progressively 455 weaker gradients toward the south. Previously, pZn:P ratios from the Southern Ocean Time Series (SOTS), 456 located in the SAZ, showed a pZn:P depth increase of approximately 4-fold (Ellwood et al., 2020), in 457 agreement with our SAZ data. The trend of increasing pZn:P ratios with depth is consistent with the lower 458 459 b value (longer remineralisation length scale) for Zn than P and is clearly illustrated when comparing the 460 decreases in pZn and P (as a percentage of their respective maximum) with depth (Fig. 5). The pZn and P 461 maxima were generally at the same (or very similar) depths. However, below the concentration maximum,

462 P decreased more rapidly than pZn (by 500 m, P decreased to values between 9 and 26% of the P maximum 463 whereas pZn decreased to values between 20 and 59% of the pZn maximum). In the AAZ, pZn:P ratios did not increase significantly with depth, suggesting more similar remineralisation length scales for these two 464 elements in this region. This is consistent with P-normalised b values (Table 4) in the AAZ which were 465 466 closer to 1 (0.76–0.97) than at the stations north of the APF (0.33–0.61). The weaker pZn:P depth gradient was the result of high ratios at the surface, likely driven by diatoms with elevated pZn:P cellular 467 468 stoichiometries. Our results are similar to previous observations showing no significant changes in pZn:P ratios with depth from a diatom dominated phytoplankton assemblage (Twining et al., 2014) and highlight 469 470 the possible taxonomic effect on Zn cycling. Reversible scavenging has also been suggested to play an 471 important role in Zn cycling (John and Conway, 2014). The increase in pZn:P with depth at the northern 472 stations potentially indicates a role for reversible scavenging in influencing Zn cycling whereby pZn is 473 remineralised at the same rate as P but is then rapidly adsorbed onto organic particles resulting in an 474 apparent deeper regeneration than P (Weber et al., 2018). However, pZn:P does not show significant increases with depth in the AAZ, where we would expect the strongest scavenging signal as there is more 475 476 Zn available. Our data therefore suggests that scavenging might be a less important component of Zn 477 cycling at the higher latitudes.

478 4.3.2 dZn:PO₄ and dZn:Si(OH)₄

479 Differences in the remineralisation length scales of Zn and P were observed in the dissolved phase data as 480 well. A plot of dZn vs PO₄ shows a curvilinear relationship whereby the change in dZn concentration 481 relative to PO₄ ($\Delta dZn/\Delta PO_4$) increases with depth (Fig. 6A) reflecting the deeper release of dZn compared 482 to PO₄ from sinking organic matter. In contrast to dZn:PO₄, a plot of dZn vs Si(OH)₄ showed a strong linear relationship (Fig. 6B; dZn (nmol kg⁻¹) = 0.064 Si(OH)₄ (μ mol kg⁻¹) + 0.690; r² = 0.93; n = 120), a well-483 484 documented feature in the Southern Ocean and most of the global ocean (De Souza et al., 2018 and references therein). The linear correlation suggests a similar remineralisation length scale for Zn and 485 Si(OH)₄. Calculated b values for biogenic silica (bSi; 0.22 ± 0.53 ; Boyd et al., 2017) span the range of our 486 487 b values for P(0.41) and pZn(0.33) preventing meaningful comparison. However, considering the observed decoupling between vertical profiles of dZn and PO₄, and close coupling of dZn and Si(OH)₄, we would 488 489 expect the b value for bSi to be more similar to that of pZn than P. Ultimately, diatoms in the high latitude 490 Southern Ocean play a significant role in Zn cycling as a result of their unique uptake stoichiometries. 491 However, it is unlikely that this mechanistic link alone can explain the global trends observed for Zn-PO₄ 492 and Zn-Si(OH)₄.

493 4.4. Upwelling and lateral transport of Zn signatures

In addition to a taxonomic role, the physical water mass circulation pattern in the Southern Ocean is a

- 495 critical component of Zn and major nutrient cycling owing to the transport of biogeochemical signatures,
- 496 set in the high latitude AAZ, throughout much of the global low latitude ocean (Vance et al., 2017). To
- 497 elaborate, the net result of biological uptake by diatoms in the AAZ are surface waters that are depleted in

498 dZn and Si(OH)₄ with respect to PO₄. Physical circulation dictates that the resulting dZn and Si(OH)₄ 499 deficient surface waters are transported northward across the APF. Furthermore, a portion of the northward 500 flowing surface water subducts in the PFZ and SAZ, forming SAMW with inherently low dZn and Si(OH)4 501 concentrations relative to PO₄. Considering that SAMW is the main communication channel to the global 502 low latitude thermocline (Sarmiento et al., 2004), the northward transport of SAMW provides a likely 503 mechanism for the observed dZn and Si(OH)₄ depleted waters of the low latitude upper ocean. One of the 504 key components of the Vance et al. (2017) hypothesis is the export of diatom cells below the winter mixed 505 layer thus setting the deep Southern Ocean Zn-P and Zn-Si(OH)₄ relationships. Our pZn:P data from the 506 AAZ shows high ratios below the winter mixed layer, indicating export of diatom cells to the subsurface, 507 where remineralisation and vigorous mixing may resolve the deep water $dZn-Si(OH)_4$ correlation despite 508 being located in separate components of the diatom cell (Vance et al., 2017).

509 4.5. Sources of pZn_{lith}

The low pZn_{lith} (~0.001 nmol kg⁻¹) throughout the SML of the AAZ reflects the lack of an atmospheric pZn 510 source for this remote open-ocean region, consistent with previous observations of low dissolved 511 512 aluminium (dAl), a tracer of atmospheric dust supply, along this transect (Barrett et al., 2018). A localised pZn_{lith} signal was however evident above the mid-ocean ridge at 50°S where pZn_{lith} was 0.01 nmol kg⁻¹, the 513 514 highest estimate along the transect. The pZn_{lith} maximum occurred in the deepest sample (3500 m), roughly 515 500 m above the seafloor, and therefore resuspension of seafloor sediment seems an unlikely source. A 516 more probable source was hydrothermal fluids expelled from the mid-ocean ridge considering that highest 517 concentrations of pAl (6 nmol kg⁻¹) and particulate iron (pFe; 2 nmol kg⁻¹), two hydrothermal indicator elements, were also measured in this sample. Once ejected, hydrothermally derived pZn may either sink as 518 519 Zn-rich particles to the seafloor (German et al., 1991) or dissociate to dZn followed by lateral advection 520 (Roshan et al., 2016). The latter mechanism therefore has the potential to influence deep dZn distributions 521 although this was not evident in our data where the $dZn:Si(OH)_4$ ratio (Si(OH)_4 is not a hydrothermally 522 derived element), in the sample was 0.069 mmol mol⁻¹, consistent with deep (< 2000 m) dZn:Si(OH)₄ for 523 the whole transect $(0.066 \pm 0.003 \text{ mmol mol}^{-1})$.

524 The elevated pZn_{lith} found in subsurface water masses of the STZ, with a core of 0.009 nmol kg⁻¹ at 800 m,

525 may reflect the influence of ARC waters which were sampled during at our STZ station (Fig. 7). The ARC,

526 which extends between surface and > 1500 m (Lutjeharms and Ansorge, 2001), is supplied by the AC which

527 flows along the south east African continental margin before retroflecting south of Africa and flowing

- eastward, often in close association with the STF. The AC is characterised by massive lithogenic inputs of pAl and pFe, up to 673 and 224 nmol kg^{-1} , respectively, through interaction with the African continental
- shelf (Grand et al., 2015) and potentially entrains significant amounts of pZn (1.10 nmol kg⁻¹ of pZn_{lith})
- using the Zn/Al_{UCC} ratio in Eq. (2)) which are ultimately inherited by the ARC. The lithogenic signal is
- bowever diluted with distance from the shelf source (Barrett et al., 2018; Grand et al., 2015) and particles
- 533 may be remineralised or sink to subsurface waters during the timescales taken for the AC to retroflect and

- return as the ARC (on the order of weeks; Lutjeharms and Ansorge, 2001). Ultimately, the enriched pZn_{lith}
- 535 in subsurface waters at the northern stations may reflect the signature of ARC enriched in lithogenic
- 536 material derived from the AC, an observation consistent with sustained high Al in subsurface waters further
- 537 east (Grand et al., 2015). However, the remineralisation of the high pZn_{lith} in the STZ is not likely to
- 538 influence subsurface dZn significantly considering that pZn_{lith} is negligible compared to dZn (> 2%).

539 4.6. Inhibited mixing south of the Indian ridge

540 Highest concentrations of dZn were measured in deep waters south of the mid-ocean ridge (hereafter Indian ridge). South of the Indian ridge (St. 50°S and 56°S), dZn was 9.07 ± 0.06 (n = 2) and 8.27 ± 0.18 nmol 541 542 kg^{-1} (n = 2) below 3000 m (Fig. 2A), respectively while to the north of the Indian ridge (St. 48°S and 45°S), dZn was lower (6.84 \pm 0.23 nmol kg⁻¹; n = 4). While dZn data for bottom waters in the Southern Ocean is 543 544 extremely limited, our concentrations appear higher compared to concentrations measured below 3000 m $(7.25 \pm 0.58 \text{ nmol kg}^{-1}; n = 6)$ and nearer to the AABW formation region (Zhao et al., 2014). The higher 545 dZn in deep waters south of the Indian ridge was associated with the highest measured Si(OH)₄ (129 \pm 2 546 547 μ mol kg⁻¹) suggesting that the resuspension of opal-rich sediment may be contributing to elevated dZn and Si(OH)₄ as observed in the South Atlantic (Wyatt et al., 2014). While the deepest sampled depth at 50°S 548 549 was in close proximity to the seafloor, resuspension of opal-rich sediment is unlikely at 56°S where the 550 deepest sampled depth (3500 m) was well above the seafloor depth (~5500 m; Fig. 2A). Instead we propose 551 that inhibited mixing to the south of the Indian ridge may increase the residence time of 'pure' AABW 552 thereby allowing for the accumulation of dZn and Si(OH)₄ and likely other nutrients which show deep water 553 maxima. Pure AABW here refers to AABW in the absence of significant mixing/dilution with other water masses (> 75% AABW). Importantly, the long residence time of dZn (3000-11,000 years; Little et al. 554 555 (2014); Roshan et al. (2016)) and Si(OH)₄ (~10,000 years; Tréguer and De La Rocha (2013)) in the ocean, 556 longer than the timescales used to measure oceanic overturning circulation (centennial timescales; 557 Matsumoto (2007)), supports the idea of nutrient accumulation in bottom waters. Pure bottom waters in this 558 region have been shown to not extend north of the Indian ridge (Anilkumar et al., 2006) and that deep waters north of the Indian ridge are instead an altered variety of LCDW resulting from diapycnal mixing 559 560 between pure AABW and LCDW (Mantyla and Reid, 1995). In an effort to further validate these 561 observations, we performed an eOMP analysis which computes the contributions of pre-defined end-562 member water masses to each measured sample (see Section 2.7). Consistent with our hypothesis, results 563 of the eOMP analysis suggest that samples below 3000 m south of the Indian ridge are composed predominantly of AABW, up to 85% at the deepest sampled depth (Fig. 8). It therefore appears plausible 564 that inhibited mixing and increased residence times of deep waters south of the Indian ridge may contribute 565 to elevated dZn and Si(OH)₄ concentrations in this region. We do however acknowledge that more deep 566 567 water measurements are necessary to confirm this hypothesis.

568 5. Conclusion

569 Winter measurements of dZn and pZn are presented along the 30°E longitude in the Indian Sector of the 570 Southern Ocean. The results provided the unique opportunity to investigate Zn cycling over seasonal cycles 571 and to gain insights into processes driving the exchange between dissolved and particulate phases in a 572 region which is extremely data scarce. Vertical profiles of dZn showed surface depletions and progressive 573 enrichments with depth and contrasted profiles of pZn which showed enrichments near the surface followed 574 be a decline in concentration with depth. These characteristics reflect biological uptake in surface waters 575 and remineralisation in the subsurface. Seasonal variations in Zn distributions were most apparent at the 576 higher latitudes where winter dZn was higher, and pZn was lower, than spring and summer measurements 577 owing to reduced biological uptake. Additionally, deep winter mixing may preferentially resupply dZn 578 (relative to PO₄) to winter surface waters through access to dZn-enriched (from diatom remineralisation) 579 waters found below the spring/summer mixed layers. The composition of pZn was dominated by biogenic 580 pZn although lithogenic contributions were observed at depth over the Indian ridge, likely a hydrothermal 581 signal, and at mid-depths in the STZ, potentially from advection of Agulhas waters rich in lithogenic 582 material derived from the African continental margin. Calculated vertical attenuation factors for our 583 particulate data suggest Zn has a longer remineralisation length scale than P and provides a mechanism 584 explaining the observed differences in their respective dissolved phase vertical distributions, i.e., the deep 585 dZn maxima and intermediate PO₄ maxima result in a curvilinear relationship. There were however distinct differences in Zn cycling across the transect. In the AAZ for example, the high surface pZn:P ratios, 586 587 characteristic of Southern Ocean diatoms, was driven by increased dZn availability and potentially Fe-588 limitation. Furthermore, pZn:P in the AAZ did not vary significantly with depth, in contrast to the lower 589 latitude stations where pZn:P increased significantly with depth. The high pZn:P throughout the upper water 590 column of the AAZ indicates the export of diatom cells below the winter mixed layer, as hypothesised by 591 Vance et al. (2017), where rigorous mixing may resolve the strong linear $dZn-Si(OH)_4$ correlation in deep 592 waters despite these elements being located in separate components of the diatom cell. Lastly, inhibited 593 mixing leading to increased residence times of bottom waters south of the Indian ridge may partly explain 594 the elevated dZn and $Si(OH)_4$ in comparison to deep waters to the north of the Indian ridge.

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Table 1. Results from the ICP-MS analysis of GEOTRACES SAFE D2, GSC and GSP seawater reference materials, NASS-7 certified seawater reference material and our own WISOS internal reference seawater are compared with respective consensus values. Consensus values for SAFe D2 as of 2013 and GSC and GSP as of 2019. Certified values for NASS-7 as of 2018. Calibrated mean values for WISOS calculated by repeat analysis (n > 10) of a large volume surface seawater sample collected from 55°S; 28°E. Instrument

and method blanks as well as ICP-MS detection limits are also shown.

	dZn
	nmol kg ⁻¹
SAFe D2	
Consensus	7.43 ± 0.25
Measured $(n = 4)$	7.23 ± 0.25
GSC	
Consensus	1.40 ± 0.10
Measured $(n = 5)$	1.41 ± 0.10
GSP	
Consensus	0.03 ± 0.05
Measured $(n = 5)$	0.10 ± 0.02
NASS-7	
Certified	6.27 ± 1.22
Measured $(n = 5)$	6.59 ± 0.07
WISOS reference seawater	
TracEx Calibration (n = 10)	9.67 ± 0.23
Measured $(n > 30)$	9.63 ± 0.24
Blanks	
Instrument $(n = 5)$	0.07 ± 0.01
Method $(n = 5)$	0.09 ± 0.01
<i>Limit of Detection</i> $(n = 5)$	0.02

	pZn	Р
	mmol kg ⁻¹	mmol kg ⁻¹
PACS 3		
Certified	5.80 ± 0.23	29.38 ± 2.91
Measured	6.24 ± 0.72	29.41 ± 0.31
% recovery	108	100
MESS 4		
Certified	2.25 ± 0.09	33.58 ± 5.17
Measured	2.25 ± 0.59	35.58 ± 0.51
% recovery	100	106
BCR 414		
Certified	1.71 ± 0.04	428.84 ± 51.20
Measured	1.66 ± 0.21	409.34 ± 52.14
% recovery	97	95
Filter Blank	1.51 pM	0.08 nM
Limit of Detection	2.82 pmol kg ⁻¹	0.03 nmol kg ⁻¹

Table 2. Results for pZn and P from the ICP-MS analysis of PACS 3, MESS 4 and BCR 414 certified
 references materials. Filter blanks as well as ICP-MS limit of detection are also shown.

892 **Table 3.** Properties characterizing the source water types (SWTs^a) considered in this study. The square of 893 correlation coefficients (r^2) between the observed and estimated properties are also given.

	Potential temperature (°C)	Salinity	$\begin{array}{c} O_2{}^0 \\ (\mu mol \; kg^{\text{-}1})^b \end{array}$	NO3 ⁰ (μmol kg ⁻¹) ^b	PO4 ⁰ (µmol kg ⁻¹) ^b	Si(OH)4 ⁰ (µmol kg ⁻¹) ^b
UCDW	3.67	34.31	319.00	20.67	1.52	36.51
LCDW	0.40	34.68	345.17	19.99	1.50	115.18
NADW	2.93	34.91	323.28	12.48	0.84	13.21
AABW	-0.47	34.66	353.12	21.92	1.62	124.91
r^2	0.99	0.99	0.97	0.91	0.82	0.93

^a UCDW: Upper Circumpolar Deep Water; LCDW: Lower Circumpolar Deep Water; NADW: North Atlantic Deep Water; and AABW: Antarctic Bottom Water.

^b Oxygen (O_2^0) , nitrate (NO_3^0) , phosphate (PO_4^0) and silicic acid $(Si(OH)_4^0)$ represent preformed values; note that O_2 represent saturation.

895 Table 4. Vertical attenuation factors (b values) for phosphorus (P), particulate zinc (pZn) and particulate
896 cadmium (pCd). ND, no data. The P normalised (P norm) b values for pZn and pCd were obtained by
897 dividing by that of P.

		all data					< 1000 m					
Lat (°S)	Region	Р	pZn	P norm.	pCd	P norm.	Р	pZn	P norm.	pCd	P norm.	
41	STZ	0.42	0.24	0.57	0.43	1.01	0.40	0.28	0.69	0.40	1.01	
43	SAZ	0.56	0.18	0.33	0.52	0.93	0.55	0.23	0.42	0.50	0.92	
45	SAZ	0.39	0.24	0.61	0.37	0.96	0.34	0.18	0.53	0.31	0.89	
48	PFZ	0.50	0.29	0.57	0.55	1.09	0.49	0.27	0.55	0.52	1.07	
50	AAZ	0.45	0.41	0.90	0.50	1.10	0.42	0.39	0.93	0.46	1.08	
56	AAZ	0.49	0.37	0.76	0.58	1.18	0.51	0.39	0.76	0.56	1.10	
58	AAZ	0.27	0.27	0.97	0.32	1.16	0.23	0.22	0.97	0.27	1.18	
All		0.41	0.33	0.78	0.46	1.11	0.39	0.30	0.78	0.43	1.10	
Ellwood et al. (2020)	SAZ	2.04	0.82	0.40	2.41	1.18						
Boyd et al. (2017)	South Pacific	$\begin{array}{c} 0.88 \\ \pm \ 0.48 \end{array}$	$\begin{array}{c} 0.77 \\ \pm \ 0.34 \end{array}$	0.88	ND	ND						

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901 Figure 1. A) Station locations during the 2017 Winter Cruise along the GEOTRACES GIpr07 transect (30°E longitude). Positions 902 of the frontal systems intersected, and corresponding oceanic zones, are shown as well as the main surface currents. Acronyms in 903 alphabetical order are as follows; AAZ: Antarctic Zone; AC: Agulhas Current; ACC: Antarctic Circumpolar Current; APF: 904 Antarctic Polar Front; ARC: Agulhas Return Current; MIZ: Marginal Ice Zone; PFZ: Polar Frontal Zone; SACCF: Southern 905 Antarctic Circumpolar Current Front; SAF: Subantarctic Front; SAZ: Subantarctic Zone; SBdy: Southern Boundary; STF: 906 Subtropical Front; STZ: Subtropical Zone. B) Scatter plot of potential temperature (°C) against practical salinity used to 907 characterize the water mass regime. Potential density ($\sigma\theta$; kg m-3) isolines also shown. Acronyms in alphabetical order are as 908 follows; AABW: Antarctic Bottom Water; AAIW: Antarctic Intermediate Water; AASW: Antarctic Surface Water; LCDW: Lower 909 Circumpolar Deep Water; NADW: North Atlantic Deep Water; SAMW: Subantarctic Mode Water; SASW: Subantarctic Surface 910 Water; SICW: South Indian Central Water; STSW: Subtropical Surface Water; UCDW: Upper Circumpolar Deep Water. Figures 911 constructed using Ocean Data View (ODV; Schlitzer, 2020).

- 912 **Figure 2.** Section plots of A) dZn (nmol kg⁻¹) overlain with potential temperature (°C), B) pZn overlain with potential density (kg
- 913 m^{-3} , and C) pZn_{lith} (nmol kg⁻¹) overlain with the percentage (%) that pZn_{lith} contributes to total pZn. Each figure is separated into 914 0-500 m (upper panel) and 500 m to seafloor (bottom panel). Frontal positions and corresponding zones are shown. Figure 915 constructed using Ocean Data View (ODV; Schlitzer, 2020).

Figure 3. Surface water (~25 m) comparisons of A) dZn and B) pZn from the Indian, Pacific and Atlantic sectors of the Southern
Ocean during winter, spring and summer. Data references as follows: winter Indian Sector (this study), winter Pacific Sector
(Ellwood, 2008), winter Atlantic Sector (Cloete et al., 2019), spring Indian Sector (Wang et al., 2018), spring Pacific Sector (Coale
et al., 2005), spring Atlantic Sector (Löscher, 1999), summer Indian Sector (Barrett et al., 2018; Janssen et al., 2020), summer
Pacific Sector (Butler et al., 2013; Coale et al., 2005; Ellwood et al., 2020; Sieber et al., 2020), and summer Atlantic Sector (Cloete
et al., 2019; Croot et al., 2011; Sieber et al., 2020).

- 922 **Figure 4.** Surface water (~25 m) dZn, pZn, pZn:P, dFe and Si(OH)4 across the transect. Note: where necessary units were converted 923 to plot on a single vertical axis for direct comparison. The biogeochemical zones crossed during the transect are shown on the top
- 924 horizontal axis.

Figure 5. pZn (grey bars) and P (black bars) normalised to their respective maxima (bottom axis) for the upper 500 m at A) 41°S,
B) 43°S, C) 45°S, D) 48°S, E) 50°S, F) 56°S and G) 58°S. Red circles indicate the absolute pZn:P ratios at each depth (top axis).
The dashed blue line represents the MLD at each station.

Figure 6. Scatter plots of A) dZn vs PO4 and B) dZn vs Si(OH)4 for the full dataset. Sample points are coloured based on neutral density. Figure constructed using Ocean Data View (Schlitzer, 2020).

Figure 7. Map showing the presence of the Agulhas Current system, determined using Absolute Dynamic Topography (in meters)
 during our occupation of St. 41°S (10 July 2017). The contour defining the Agulhas Current system is the 0.75 m contour. Image
 courtesy of Marcel du Plessis.

Figure 8. Results of the extended optimum multiparameter analysis (eOMP) optimised for resolving the contribution of Antarctic
 Bottom Water (AABW) to each measured sample. Figure constructed using Ocean Data View (ODV; Schlitzer, 2020).

















