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Pathways and timescales of Southern Ocean hydrothermal iron and manganese transport

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Scarcity of iron and manganese limits the efficiency of the biological carbon pump over large areas of the Southern Ocean. The importance of hydrothermal vents as a source of these micronutrients to the euphotic zone of the Southern Ocean is debated. Here we present full depth profiles of dissolved and total dissolvable trace metals in the remote eastern Pacific sector of the Southern Ocean (55–60° S, 89.1° W), providing evidence of enrichment of iron and manganese at depths of 2000–4000 m. These enhanced micronutrient concentrations were co-located with ³He enrichment, an indicator of hydrothermal fluid originating from ocean ridges. Modelled water trajectories revealed the understudied South East Pacific Rise and the Pacific Antarctic Ridge as likely source regions. Additionally, the trajectories demonstrate pathways for these Southern Ocean hydrothermal ridgederived trace metals to reach the Southern Ocean surface mixed layer within two decades, potentially supporting a regular supply of micronutrients to fuel Southern Ocean primary production.

Despite being the largest high-nutrient low-chlorophyll region^{1,[2](#page-7-0)}, the Southern Ocean (SO) is a globally important organic carbon sink. The interplay between circulation and remineralisation dynamics set the rate of carbon uptake by the biological carbon pump³ with 10% of global biological carbon export occurring in the region⁴. Iron (Fe) is an essential but often scarce micronutrient that can limit primary production and the efficiency of the biological carbon pump in the $SO^{5,6}$ $SO^{5,6}$ $SO^{5,6}$. Other elements can potentially be (co-)limiting alongside Fe in the $SO⁷$, with recent evidence for manganese (Mn) (co-)limitation in particular 2^{38-10} .

As non-conservative elements with residence times on the order of decades in the deep ocean¹¹, the distributions of dissolved Fe and Mn (dFe, dMn; $<$ 0.2 μ m) in seawater are spatially coupled to sources. The main external sources of Fe and Mn to the ocean are margin sediments, atmospheric dust, and hydrothermal venting $6,12$. Input from margin sediments enhances primary production downstream of islands in the $SO^{13,14}$ $SO^{13,14}$ $SO^{13,14}$, and sustains ecosystems in Antarctic shelf regions^{12,15}, where melting glacial ice can be an additional source of Fe and Mn^{16,17}. Marine aerosol Fe concentrations can vary by >3 orders of magnitude, causing sporadic and seasonal changes to the widespread deposition of Fe to the surface ocean¹⁸. However, according to both observations and models, the remote southern hemisphere oceanic gyres and polar regions have some of the lowest Fe aerosol deposition fluxes in the world (<0.01 g Fe m⁻² yr^{-1)[19](#page-8-0)}.

Both Fe and Mn are concentrated in hydrothermal fluids, often enriched by a factor of $>10^6$ relative to background seawater²⁰. Early studies suggested that Fe and Mn entering the ocean from hydrothermal vents were precipitated or scavenged and sedimented close to vent sites $12,21$, with precipitation reported to occur before the plume reaches neutral buoyancy²². However, it is now known that a small fraction of the dFe supplied from hydrothermal vents is sufficiently stabilised against precipitation to be transported in the ocean interior $^{23-25}$. Hydrothermal venting is estimated to supply 4 ± 1 Gmoles dFe yr⁻¹ to the wider deep ocean away from the proximal vent sites $23,26$, representing a continuous Fe input in contrast to short-term variations observed in other Fe sources, such as atmospheric deposition²⁶. Similarly, observed increases in deep ocean dMn concentrations have been attributed to hydrothermal activity¹⁶, and incorporating hydrothermal dMn inputs increases the accuracy of modelled oceanic dMn $distributions¹²$. Our understanding of the hydrothermal plume processes responsible for the physico-chemical stabilisation of Fe is developing, though not yet comprehensive. Within hydrothermal plumes, dFe has been shown to form inorganic nanoparticles^{27,28}, larger inorganic colloids²⁹, and

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organic complexes^{30,31}. The co-location of carbon with Fe in plume particles suggests that organic carbon may alter the chemical behaviour of Fe oxyhydroxides²⁹ and create localised regions of Fe(II) enrichment³¹.

Due to the inhibition of vertical mixing in the ocean interior by density stratification, hydrothermal plumes tend to travel predominantly along isopyncals^{32,33}. Westerly winds drive the Antarctic Circumpolar Current to create a northward surface flow (Ekman Transport), causing isopycnals to shoal in the $SO³⁴$. Consequently, upwelling deep waters in the SO may provide a pathway for hydrothermal trace metals to be mixed into the SO euphotic zone^{21,35}. Inputs of hydrothermal dFe from shallow (<500 m) and deep (>2000 m) vents have been linked to regional phytoplankton blooms within $36,37$ and outside of the SO³⁸. The potential for hydrothermal dFe to sustain primary production in the SO has been evaluated in models $39,40$ $39,40$, but consensus on the significance of this source of Fe and Mn has yet to be reached.

To sustain SO primary production, upwelling timescales must be short enough to deliver hydrothermal Fe and Mn to the euphotic zone before removal processes, such as scavenging and precipitation, deplete these metals. Assuming that some degree of physico-chemical stabilisation of dFe in neutrally buoyant plumes may allow transportation to surface waters, model results indicate that hydrothermal Fe could support ≈15–30% of export production south of the Polar Front³⁹. However, estimates of dFe residence time in the East Pacific Rise (EPR) far-field plume at 15° S have been revised from quasi-conservative, based on comparisons with ³He measurements²³, to non-conservative with a residence time of 9-50 years, based on comparisons with 228 Ra measurements⁴¹. The 9-50 year residence time estimate is consistent with a global modelling study estimate of hydrothermal Fe residence time of 21–35 years³⁹. Similarly, deep ocean dMn residence times are estimated at $5-40$ years¹¹.

In comparison, transit time estimates for the shoaling of Circumpolar Deep Waters, originating in the Pacific $(\geq 30^{\circ} \text{ S})$, to the SO mixed layer range from 17 years to a few centuries⁴²⁻⁴⁵. Such uncertainty surrounding hydrothermal Fe input and stabilisation processes, alongside SO ventilation rates, is reflected in Fe modelling studies. A global steady-state inverse circulation model combined with a mechanistic Fe model was used to conclude that only 3–5% of hydrothermal dFe reaches surface waters globally⁴⁰. A subsequent study argued however that the inverse modelling approach reduces the magnitude of hydrothermal Fe inputs from ridge systems located within the SO relative to a spreading rate model approach³⁹. This ongoing debate is hindered by a lack of observational evidence of the key transport pathways. The Southeast Pacific sector of the SO has been indicated as a critical region for hydrothermal Fe and Mn input by models^{12,[39](#page-8-0)}, but is a 'data desert' for trace metal observations. In this study, we provide the first full-depth profiles of Fe and Mn from this SO region, which show a clear midwater hydrothermal trace metal signal. By modelling water pathways, we trace the hydrothermal signal back to specific ridge systems and evaluate whether the observed hydrothermal Fe and Mn could supply the SO mixed layer, and therefore fuel SO primary production.

Results and discussion

Observations of hydrothermal trace metals in the southeast Pacific sector of the Southern Ocean

Our study sampled a north–south transect along 89° W located in the Mornington Abyssal Plain in the southeast Pacific Ocean (Fig. 1A). As expected, SO surface waters were characterised by extremely low dFe (<0.2 µm, 0-30 m depth; 0.09 ± 0.04 nM, $n = 18$) and total dissolvable Fe (TdFe; unfiltered) concentrations (0–30 m depth; 0.19 ± 0.09 nM, $n = 13$; Fig. 1B, C) and excess macronutrient concentrations (nitrate $+$ nitrite 16.6–23.6 μ M, phosphate 1.0–1.6 μM; data not shown), consistent with previous upper ocean observations in this region⁴⁶ and wider SO biogeochemistry³⁵.

The most striking feature of our transect was Fe enrichment in deep waters (Fig. 1B, C), with dFe concentrations of 0.48–0.96 nM occurring between 2000 and 4000 m at the northern end of our transect Fig. 1A; Table S1). Elevated TdFe concentrations (>2.5 nM with a maximum concentration of 15.4 nM), associated with reduced dFe concentrations, were also observed in benthic nepheloid layers driven by particle accumulation and benthic sediment resuspension of particulate Fe and scavenging of dFe (Figs. 1C, [2D](#page-2-0)). For comparison, particulate Fe concentrations (up to 88 nM) were observed above the seafloor at open ocean stations in the North Atlantic 47 .

Fig. 1 | Study region and dissolved iron and total dissolvable iron transects. A Study area in the southeast Pacific and Southern Ocean. Red dots are the location of known active vent sites 82 . Lack of exploration means that knowledge of active vent site locations over the region is incomplete. The yellow arrow is the approximate flow of water at 2000-4000 m depth^{[83](#page-9-0)} and white arrow is the approximate flow of Southeast Pacific Deep Slope Water⁵⁴. The solid black line is the transect occupied by this study. The white cross is the sampling location of a previous ³He profile^{[49](#page-8-0)} collected in 1993 ~83 km northeast (bearing of 055°) of our northernmost station. Triangles indicate the start locations for forward-tracked trajectory modelling in this study: yellow—East Chile Rise, purple—West Chile Rise, white—Easter

Away from external sources, deep ocean dMn concentrations are typically very low (<0.2 nM) due to scavenging removal into the particulate phase and oxidation of Mn ions to insoluble Mn oxides¹⁶. However, like dFe,

Microplate, green—North East Pacific Rise, orange—South East Pacific Rise (SEPR) and blue—Pacific Antarctic Ridge (PAR). B, C Section plots of dFe (<0.2 μm) and TdFe (unfiltered) concentrations along the transect with neutral density overlain. Black dots mark sampling depths. Note that concentrations in (B) and (C) are on different scales and that TdFe concentrations were typically >2.5 nM near the seafloor. The maximum concentration measured was 15.39 ± 1.12 nM. OOI is the sampling location coinciding with the Ocean Observatories Initiative sustained observatory location, 54.08° S 89.67° W, [https://oceanobservatories.org/array/](https://oceanobservatories.org/array/global-southern-ocean-array) [global-southern-ocean-array](https://oceanobservatories.org/array/global-southern-ocean-array). Stations OOI, TN, and TS (C) are the locations of trace metal depth profiles displayed in Fig. [2.](#page-2-0)

Fig. 2 | Depth profiles of iron, manganese and excess helium. A Depth profiles of dFe at stations OOI, TN and TS. B Depth profiles of dMn at stations OOI, TN and TS. C Depth profiles of dissolved trace metals (dFe and dMn) and TdFe at our northernmost station OOI (combined Fe data from four site visits) and nearby historic xs³He (see Fig. [1](#page-1-0)A; white cross). D All dFe and TdFe data from the transect and nearby historic xs³He plotted against neutral density. Error bars for dFe and TdFe are the range of duplicate analyses and for dMn the standard deviation of triplicate analysis.

basin-scale transport in vent plumes has been observed e.g. westward transport of dMn from the EPR (10–17° S) occurs in the low latitude Pacific Ocean²³. Along our transect we present, elevated dMn concentrations (up to 0.39 nM, Table S1) occurred coincidently with the enriched dFe signal (Fig. 2A–C), compared to a background dMn concentration (~0.2 nM) immediately above and below the mid-depth enrichment (Fig. 2B, C). The Fe and Mn enrichment we observed was centred on a neutral density surface
of $\text{Yn} = 28.0-28.1 \text{ kg m}^{-3}$ (Fig. 2D), within Circumpolar Deep of ϒn = 28.0–28.1 kg m[−]³ (Fig. 2D), within Circumpolar Deep Water (Fig. S1). In the southwest Pacific (170° W), Circumpolar Deep Water has background dMn concentrations of $~0.1-0.2~\text{nM}^{48}$ $~0.1-0.2~\text{nM}^{48}$ $~0.1-0.2~\text{nM}^{48}$ away from hydrothermal and sediment inputs. The Υ n = 28.0–28.1 kg m⁻³ density surface shoals by ~880 m towards the south of the transect as part of the wider Ekman driven upwelling occurring in the SO. The influence of this wind-driven upwelling is shown clearly in the maximum dFe concentrations, which tracked the isopycnals along the depth section (Fig. [1](#page-1-0)) and in the dMn profiles from stations OOI, TN and TS (Figs. [1B](#page-1-0), C and 2A, B; Table S1).

Mantle helium (He), which is enriched in primordial ³He, is also released into the ocean at mid-ocean ridges. A ratio of ³He/⁴He in excess of atmospheric values (hereafter xs³He) in deep ocean waters is a widely used tracer of hydrothermal inputs⁴⁹, though the basin-scale hydrothermal Fe supply from SO vents may not be well predicted by the spatial distribution of ³He inputs, likely because of inter-vent Fe:³He ratio variability³⁹. Nevertheless, at a single location, the coincident enrichment of ³He, Fe and Mn in deep ocean waters is indicative of hydrothermal input of these elements.

Helium isotope observations, previously sampled near station OOI (Fig. [1](#page-1-0)), revealed the presence of xs³He at the depths and neutral density of our observations of trace metal enrichment (Fig. 2C, D). Although the helium dataset was collected 26 years and 83 km away from our trace metal observations, the comparable distribution of dFe, TdFe and xs³He with neutral density (Fig. 2D) provides confidence that we observed a consistent oceanographic signal in both datasets. However, we cannot account for any temporal variability of end-member metal and helium concentrations during the 26-year gap between trace metal and helium sample collection⁵⁰. The Fe concentration maximum at the northern end of our study site was slightly deeper (215 m) than the mantle xs³He maximum. A similar offset in the distribution has been observed further north in the far-field plume of the EPR (20–26°S) and has been attributed to the reversible scavenging of dFe onto sinking particles, which deepens the metal concentration maximum⁵¹. The offset we observed may also be due to temporal variability of endmember metal and helium concentrations, the influence of multiple vent sites with varying Fe:³He ratios, and/or reversible scavenging combined with layering of fine particulate material with irregular element distribution within plumes 52 .

The balance between the release of dFe from remineralising organic matter and removal of dFe from solution via scavenging processes leads to an accumulation of dFe in Pacific Mode and Intermediate Wates^{[53](#page-8-0)}. We show that the signal we observed is decoupled from the mineralization of biogenic particles by plotting dFe against apparent oxygen utilisation (Fig. S2). In waters below the mixed layer, there is a positive linear relationship between

dFe and apparent oxygen utilisation indicating that remineralisation of sinking organic matter exerts an important control on dFe concentrations, consistent with previous investigations of Pacific mode and intermediate waters⁵³. However, the enrichment of dFe we observed in deeper waters clearly deviates from the linear dFe- apparent oxygen utilisation trend, indicating an additional deep water dFe source such as long-range transport of hydrothermal dFe^{24} .

Identifying the source region of the observed hydrothermal signal The bathymetry of the southeast Pacific Ocean has several mid-ocean ridge systems (Fig. [1](#page-1-0)A), the most prominent being the EPR which is aligned meridionally at ~115° W at depths of around 2500 m. The southern extension of the EPR, the Pacific Antarctic Ridge (PAR), is aligned zonally (\sim 55–65° S). The Chile Rise (CR; \sim 40–45° S) bounds the North of the MorningtonAbyssal Plain (our study region) between the EPR and the coast

area of the OOI station (location indicated by pink line). Additionally, the grey line superimposed on the pink line highlights the depth range where dFe concentrations were >0.75 nM (Fig. [2](#page-2-0)). The cyan line marks the trajectory seeding depth which mimics a neutrally buoyant plume. Bathymetry along 54° S (to match station OOI) is shown. Note the colour bar scale is different for (C) compared to (A, B). D An eastwest transect of excess Helium (xs³He) at 54° S as an indicator of hydrothermal influence from the Jenkins et al. dataset⁴⁹.

of Chile. The OOI site is a little south of the confluence of the easterly flowing Antarctic Circumpolar Current and the eastern boundary current. The latter flows within ~1500 km of the coast of South America between 1500 and 3500 m, this Southeast Pacific Deep Slope Water⁵⁴ represents the main route for mid-depth flow in the South Pacific to enter the SO^{54-[56](#page-8-0)}. Volcanic activity at multiple sites along ridge systems (Fig. [1](#page-1-0)A), alongside complex interior circulation, results in widespread xs³He in intermediate and deep waters (\sim 1000–4000 m) in the South Pacific Ocean⁴⁵. A previous meridional transect along 88° W shows the southward extension of xs³He towards our study area (Fig. S3A), which could indicate plumes originating from ridges to the north e.g. the Sala Y Gomez Ridge and CR (Fig. [1A](#page-1-0)). Similarly, a zonal transect along 54° S shows an eastward extension of xs³He from the South EPR (SEPR) that intersects the OOI station (Fig. [3](#page-3-0)D; Fig. S3B). Because the xs³He observations do not allow us to discern a clear source region, we undertook trajectory tracking simulations to provide an independent estimate for the origin of the hydrothermal signal.

To trace trajectories from potential sources, a series of model release sites were first selected. Along the poorly explored southern ridge systems, bathymetry along with xs³He observations and previous literature^{[33](#page-8-0)} were used to select an arc of potential release sites to the north and west of station OOI. To simulate a neutrally buoyant hydrothermal plume, trajectories were initiated within a 0.5° grid around an assumed source extending 500 m upwards in the water column above the ridge at theWest Chile Rise (WCR), East Chile Rise (ECR), North EPR (NEPR), SEPR and PAR South (PARS) locations (Fig. [1;](#page-1-0) Table 1) and tracked forwards in time. A reverse simulation was also performed whereby trajectories were backtracked from within a 0.5° grid around the OOI site between 2000 and 3950 m depth, coincident with our hydrothermal Fe and Mn enrichment observations (Table 1; Fig. [2C](#page-2-0)), towards the ridge systems. The mid-ocean ridge system surrounding our study region was divided into five regions (Fig. S4; Table S4), including an additional region PAR North (PARN) which did not have a forwardtracking release site. We quantified the backwards trajectories that passed through the likely plume depths (i.e. extending up to 500 m of water column above a ridge^{[57](#page-8-0)-[59](#page-8-0)}) in these regions (Tables 1 and S2). For both forward and backwards tracking, trajectories were traced for 20 years by applying a Lagrangian simulator, called 'parcels' [\(https://oceanparcels.org/](https://oceanparcels.org/) 60), to the velocity components of a global ocean model (Nucleus for European Modelling of the Ocean; NEMO) 61 61 61 at an eddy-resolving (0.083°) horizontal resolution. Trajectories estimate the flow of water within the model ocean and do not represent concentrations of trace metals.

Trajectory estimates strongly supported a westerly source region with no influence from northerly ridge systems (Table 1). The SEPR region is identified as the most likely source for our observed signal. Forwardtracking simulations from the SEPR resulted in the greatest percentage of trajectories (9%) crossing within a 0.5° area around OOI at a depth coincident with the observed Fe and Mn enrichment compared to other release sites (Fig. [3](#page-3-0)C; blue line). Similarly, in the backward-tracking simulations, the greatest percentage of trajectories (21%) released from OOI passed within the SEPR source region (Table 1). In an oceanographic context, these percentages can be thought of as demonstrating the strength and coherency of transport pathways within the wider ocean circulation between the source regions and OOI i.e. the greater the percentage the more dominant the pathway. The SEPR also had the shortest median transit timescales to/from OOI (10–12 years). Further west, the backward-tracking simulations indicate that 8–20% of trajectories starting at OOI intersect with the PAR ridge systems with a median transit time of 13–14 years. However, the peak in the trajectory signal propagated from the PAR regions was deeper than the observed Fe and Mn peaks at OOI, especially for PARS (Fig. S3H). Our transit time estimates from the SEPR and PAR are at the lower end of estimated hydrothermal dFe residence times of 9-50 years^{[39](#page-8-0),41}.

The NEPR makes a minor contribution in both forward and backward trajectories over 20 years (0.3–1.3% of trajectories to/from OOI), with trajectories exhibiting longer median transit times to reach OOI (14–18 years), despite being much closer in proximity to OOI than the PARS site which made the second largest contribution (Fig. [1;](#page-1-0) Table 1). Similarly, trajectories from the CR did not intersect with the OOI site within 20 years.

Our simulations indicate minimal exchange between waters north and south of 45° S in the Mornington Abyssal Plain, likely due to the South Pacific Current, which has diverging pathways above and below 40–43° [S55,62.](#page-8-0) This interpretation is supported by a model validation between the NEMO model used for the trajectory simulations and the Estimating the Circulation and Climate of the Ocean (ECCO) model state estimate^{63,64} (Fig. S5). Both showed that the dominant currents at OOI originate from the west with weaker currents north of 45° S. Overall, the trajectory simulations indicated that the observed OOI trace metal plume originates from a ridge to the west of OOI, which is consistent with our understanding of the fast-moving Antarctic Circumpolar Current and weak currents in the northern Mornington Abyssal Plain⁶⁵. Therefore, the strong xs³He signal to the north in the Mornington Abyssal Plain region (Fig. S3A) should not be interpreted as traversing southwards towards OOI. Our results indicate that the xs³He signal observed at OOI has most likely traversed eastwards from the SEPR and/or from along the PAR (Fig. [3D](#page-3-0)). The PARS/N trajectory simulations highlight a pathway for potential hydrothermal transport along the PAR, that tracks northwards up to the SEPR and then eastwards towards the OOI site (Fig. S4E, G). Thus, it is possible we observed the amalgamation of trace metal inputs from multiple vent sites along the PAR and SEPR at the OOI site and along the transect, rather than a discrete single vent signal.

The proportion of trajectories that passed within a 0.5° area around the OOI site is presented relative to the number of particles that were successfully released during each simulation (Tables S2 and S3). The source regions that trajectories were tracked backwards in time towards were much larger than the area around station OOI, as they span along sections of the ridges, allowing more trajectories to pass into the source regions in the backward-tracking simulations (Tables S2, S3 and S4).

Systematic surveying for hydrothermal activity is lacking for the majority of the southern EPR and PAR systems⁶⁶. However, it is expected that the ridge system is hydrothermally active as multiple lines of evidence suggest that hydrothermal venting is common. Metalliferous sediments (enriched in Fe and Mn compared to aluminium), formed by the precipitation of Fe and Mn from hydrothermal fluids, are found along these ridge systems⁶⁷. Additionally, the magmatic budget hypothesis⁶⁶ predicts that variability in magma supply is the primary control on the large-scale hydrothermal distribution pattern along spreading ridges and is supported by a linear relationship between spreading rate and frequency of vent fields. Extrapolation of this relationship has been used to estimate a total of ≈200–300 undiscovered vent fields along the SEPR and PAR with an average distance between vent fields along the SEPR of $25-30$ km^{[66](#page-9-0)}. Indeed, high-resolution optical and redox sensor measurements made along 1470 km of intermediate and fast-spreading mid-ocean ridge suggest that the frequency of vent sites is 3–6-fold higher than current observations (Fig. [1](#page-1-0)A), with a mean discharge spacing of $3-20 \text{ km}^{68}$.

Similarly, evidence from observational and modelling studies exists for the long-range transport of chemical signatures from these ridge systems. For example, sections of the PAR (south of 55° S) were identified as the likely source region of a ³He anomaly found on the neutral density surface Υ_n = 28.2 kg m⁻³, indicative of active venting³³. Modelling dFe hydrothermal vent input as a function of ridge spreading rate found good agreement with observed dFe anomalies in the abyssal SO, with some of the highest rates of hydrothermal Fe input in the Pacific sector along the PAR and SEPR^{[39](#page-8-0)}. Similarly, a global ocean modelling simulation of Mn predicted strong input from the SEPR which produced a dMn anomaly that extends to the location of our transect¹².

Could hydrothermal iron and manganese from southeast Pacific vents fuel Southern Ocean primary production?

The potential for hydrothermal Fe and Mn to fuel primary production depends on the balance between Fe and Mn residence times and the ventilation timescale of hydrothermal plume-influenced waters. A transit time of 99 ± 18 years for the shoaling of deep waters to the SO surface has been estimated using a global ³He mass balance model, in which ³He predominantly enters into the SO via deep South Eastern Pacific waters⁴⁵. Deep ocean re-exposure timescales of a few centuries have also been calculated using a low horizontal resolution (2°) global steady-state ocean circulation inverse model⁴². These ventilation timescales are likely incompatible with the notion that Fe and Mn from southeast Pacific vents could sustain SO productivity when compared with estimated seawater residence times for Fe and Mn. Indeed, coupling a mechanistic Fe model, where hydrothermal input is estimated from a fixed Fe:³He ratio, with an ocean inverse circulation model allows for scavenging processes to effectively trap hydrothermal Fe in the deep ocean 40 .

However, uncertainty around the magnitude of hydrothermal Fe input from SO ridge systems, and accounting for mesoscale processes when estimating ventilation timescales, may in fact allow for rapid transport of hydrothermal Fe and Mn to upper ocean waters. An inverse modelling approach reduces the magnitude of ³He input from SO ridge systems relative to a spreading rate model approach. A spreading rate model estimates ³He input as a function of ridge spreading rate, and likewise assigns an Fe:³He ratio to estimate a hydrothermal Fe flux^{[39](#page-8-0)}. Comparisons with observed dFe concentrations revealed that the inverse model approach does not replicate the magnitude of an observed SO hydrothermal dFe signal from the GEOTRACES GS01 section South of Tasmania³⁹. A consequence of the spatial redistribution of hydrothermal Fe input away from the SO vent sites by the inverse modelling approach is a reduction in the estimated amount of hydrothermal Fe reaching the upper 250 m of the ocean by 4–5 fold relative to the spreading rate model^{39,40}. It is also possible that previous studies may be over-estimating ventilation times due to their coarse resolution. Mesoscale (10–100 km) processes are important conduits for vertical mixing in the $SO^{43,44}$. Using a Lagrangian particle tracking model transit times of 17–90 years were estimated for Circumpolar Deep Waters

(originating from 30° S between 1000 and 4000 m) to upwell to the SO mixed layer^{43,44}. Importantly, transit time estimates decrease as model resolution becomes finer^{43,44} with the shorter 17-year timescale estimated using a 0.1° eddy-resolving model. This highlights that accounting for mesoscale eddies is important for estimates of timescales for SO upwelling⁴⁴.

To investigate the potential fate of the observed dFe and dMn signals, we continued to forward-track trajectories that passed through our cruise transect at depths of the observed trace metal peak (2000–4000 m; Fig. [4](#page-6-0)). We thus estimated the proportion of trajectories which shoal to depths coincident with SO winter mixed layer depths, the key seasonal supply mechanism of micronutrients to SO surface waters³⁵. Eighty-three percent of the trajectories passing through the cruise transect between 2000 and 4000 m continued east through the Drake Passage and then flowed northward away from the polar front (∽60° S) into deep water masses (Fig. [4](#page-6-0)A; median depth 2087 m), and so are unlikely to supply hydrothermal Fe and Mn to the SO mixed layer. However, trajectories which pass through the Drake Passage and traverse pathways south of 60° S (17%) tended to shoal in the water column by the end of the 20-year simulation (median depth 1043 m). Of the 17% of trajectories that passed south of 60° S, 10% (i.e., 1.7% of total trajectories) reached depths shallower than 600 m, a proxy for the typical winter maximum mixed layer depth in deep mixing regions of the $SO⁶⁹$, while 31% (i.e. 5.3% of total) reached shallower than 1000 m, the maximum observed winter mixed layer depth⁶⁹.

Our simulated trajectories thus indicate pathways from the observed deep southeast Pacific sector of the SO to the SO mixed layer within 20 years. Furthermore, the number of trajectories south of 60° S reaching winter mixing depths is likely to continue to increase beyond the temporal limit of our simulations (20 years). As discussed above, resolving mesoscale features in high-resolution models, as in our study (0.083° horizontal resolution), results in SO ventilation timescale estimates (within 20 years) that allow for hydrothermal Fe and Mn from SO vents to reach SO mixed layer depths within estimated Fe and Mn residences times $(5-50 \text{ years})^{41}$ $(5-50 \text{ years})^{41}$ $(5-50 \text{ years})^{41}$. Moreover, the effects of submesoscale mixing and dispersion, which will influence the transport and shoaling rates of dissolved constituents, are not resolved in the physical model driving our trajectory calculations but could further shorten ventilation timescales⁷⁰. Trajectory modelling offers an alternate approach to identifying hydrothermal source regions and ventilation pathways and timescales, which may be more tractable for localised studies. It remains challenging to isolate what drives the differences between Fe supply from different modelling approaches $39,40$ and ventilation timescales from high-resolution studies $\left(\langle 20 \rangle \right)^{43,44}$, compared to coarse resolution or global inventory studies $({\sim}100$ years)^{[42,45](#page-8-0)} discussed above, as several factors change in tandem. The significance of transport pathways to the mixed layer must ultimately depend on the residence times of hydrothermal Fe and Mn being comparable or longer than the ventilation timescales we predict, as well as the consistency and stability of hydrothermal Fe and Mn inputs, both of which remain poorly constrained.

Conclusion

We identify potential pathways for Fe and Mn from SO hydrothermal vents to reach the SO mixed layer. Enhanced Fe and Mn concentrations were observed between 2000 and 4000 m depths within the southeast Pacific sector of the SO. The observed trace metal enrichment most likely originated from the SEPR (south of 30° S). An origin further westward, such as from the PAR, was also possible. We further identified pathways for rapid (<20 years) transport of water parcels enriched in hydrothermal Fe and Mn, from vents located within the understudied Pacific sector of the SO, to SO winter mixed layer depths. Consensus on the role of deep ocean hydrothermal trace metal inputs in fuelling upper ocean primary production has not yet been reached, with modelling studies disagreeing on the importance of SO vent systems for supplying hydrothermal Fe supply to the wider SO euphotic zone^{39,40}. However, we conclude that due to the vigorous action of the Antarctic Circumpolar Current, Pacific sector SO hydrothermal inputs provide the potential for sustained input of Fe and Mn to the euphotic zone south of the Antarctic Polar Front and far downstream of the deep ocean vent sites, potentially fuelling productivity in remote areas of the SO. Given the existence of ridge systems throughout the SO, it is likely that other similar pathways exist that rapidly transport Fe and Mn from vent sites to the SO euphotic zone.

Fig. 4 | Trajectory locations at the end of the 20 years simulation. Indication of trajectory locations at the end of the 20-year simulation from the source regions. A Trajectory depth and location at 20 years and B the depths presented as above (red) and below (blue) the absolute maximum winter mixed layer depth in the SO (1000 m). C Trajectory density at 20 years. OOI is shown by the red marker. Note that only trajectories that passed within the cruise transect (OOI to 60° S) and the observed hydrothermal signal depth range (2000–4000 m) are shown.

Methods

Trace metal and macronutrient methods

Sampling was conducted during December 2019 and January 2020 on board the RRS Discovery along a transect in the eastern Pacific sector of the SO (Fig. [1](#page-1-0)). All trace metal samples were collected following GEOTRACES protocols 71 .

Briefly, dFe (0.2 μm filtered) and TdFe (unfiltered) were analysed using flow injection with chemiluminescence detection, after spiking with hydrogen peroxide⁷², in a clean laboratory at the University of Plymouth. The mean limit of detection was 0.012 ± 0.009 nM and the limit of quantification was 0.038 ± 0.026 nM ($n = 54$). Additional seawater samples collected in >1 L volumes, and acidified for >6 months, were used as in-house quality control materials as a measure of precision and were analysed every ten samples. Four in-house quality control materials were analysed; #1 0.17 ± 0.01 nM $(n = 45)$, #2 0.22 ± 0.04 nM $(n = 54)$, #3 0.26 ± 0.03 nM $(n = 124)$ and #4 0.19 \pm 0.03 nM $(n = 70)$. GEOTRACES GSP and SAFe D1 and S consensus values were analysed to determine accuracy, consensus values are $(GSP\ 0.16 \pm 0.05 \text{ nM}, \text{D1} \ 0.65 \pm 0.04 \text{ nM}, \text{S} \ 0.10 \pm 0.01 \text{ nM}),$ which compared well to values determined (GSP 0.20 ± 0.020 nM, D1 0.66 ± 0.066 nM, S 0.10 ± 0.01 nM). The authors have previously published bottom-up and top-down combined analytical uncertainty estimates (u_c) 5–10% $(k = 1)$) for this technique^{73,74}.

A subset of samples from the northern, middle, and southern stations were chosen for further trace metal analysis. dMn (0.2 μm filtered) concentration was determined using a standard addition method with off-line pre-concentration and subsequent high-resolution ICP-MS²⁹ at the National Oceanography Centre, Southampton, UK. The mean limit of detection was 0.011 ± 0.004 nM ($n = 3$). Consensus values for SAFe S $(0.812 \pm 0.062 \text{ nM})$ and D2 $(0.360 \pm 0.051 \text{ nM})$ reference material compared well with our measured values $(0.769 \pm 0.059 \text{ nM}$ and 0.372 ± 0.030 nM, respectively).

Inorganic nutrient analysis

Seawater samples were analysed for nitrate (determined as nitrate $+$ nitrite; $NO_3^- + NO_2^-$), silicate (SiO₄⁴⁻), nitrite (NO₂⁻) and phosphate (PO₄³⁻). Samples were drawn from Niskin bottles into rinsed 15 mL centrifuge tubes. Colorimetric analysis was undertaken within 24 h of the samples being taken and performed using SEAL QuAAtro 39 segmented flow autoanalyser⁷⁵. Two sets of certified reference materials (CRM Lots CJ and CB, KANSO, Japan) were determined at a start and the end of each sample run to ensure accuracy. Depending on an analysis run, the detection limits for each parameter ranged as follows: $0.02-0.2 \mu M$ for nitrate + nitrite, 0.02–0.15 μM for silicate, 0.01–0.033 μM for nitrite, and 0.002–0.019 μMfor phosphate.

Temperature and salinity calibration

Trace metal clean samples were taken with a titanium frame Conductivity Temperature Depth (CTD) fitted with 10 L Niskin bottles and an SBE 9 plus underway unit. Conductivity (and calculated salinity) data were calibrated against samples taken from each cast and run on an Autosal 8400B salinometer. The salinity was accurate to within ±0.004. There was no SBE35 thermometer fitted on the titanium frame but comparison with co-located calibrated stainless steel casts showed good agreement and data is accurate to better than ±0.005 °C

Excess helium calculations

³He is a primordial substance trapped in the mantle during planetary formation. Ratios of ³He/⁴He in excess of atmospheric values indicate a contribution from the mantle, therefore ³He can be used to identify ocean waters that have been influenced by hydrothermal activity. Excess ³He is the approximate measure of non-atmospheric ³He over saturation, we followed the approach outlined in ref. [23](#page-8-0) (Eq. 1):

³He_{xs} =
$$
(\delta^3 \text{He} - \delta^{3*} \text{He})/100 \times \text{C}[\text{He}] \times 1.384 \times 10^{-6}
$$
 (1)

where δ^3 He = 100 × (Rx/Ra – 1) × 100%, Rx and Ra are the ³He/⁴He ratios of the sample and air (1.384 × 10⁻⁶), respectively. δ^{*3} He is the helium isotope ratio anomaly in solubility equilibrium with the atmosphere.

Trajectory modelling

To identify the potential source of the observed hydrothermal Fe and Mn signal we focus on regions along the CR, EPR (NEPR and SEPR) and PAR (PARN and PARS), with regions of suspected hydrothermal activity shown in Fig. [1A](#page-1-0) (triangles).

Trajectories were used to identify the region(s) that were most likely to be the source of the deep water hydrothermal iron signal. Trajectories were tracked forwards from possible vent sites which may act as a source, and backwards from the OOI sustained observatory site, where enrichments in midwater micronutrient concentrations were observed. The velocity components used to calculate trajectories do not represent submesoscale mixing or dispersion processes 60 , which may affect the pathway of dissolved constituents in the ocean.

We estimated 20-year trajectories using 'parcels'(version 2.2.0; [https://](https://oceanparcels.org/) oceanparcels.org/[60](#page-8-0)). Parcels were applied to the NEMO eddy-resolving general circulation model (ORCA0083-N06⁶¹). Model output at 1/12[°] horizontal resolution from 1992 to 2011 was used as a 20-year climatology of the hydrodynamic flow field. NEMO was coupled to a sea-ice model (LIM[276\)](#page-9-0) and was forced with historical atmospheric reanalysis (Drakkar Forcing Set 5.2 $\frac{77}{2}$. The simulations were carried out using 3D Runge-Kutta fourth order (RK4) timestep integration as the advection scheme with a timestep of 1 day to estimate the trajectories⁷⁸. The trajectory position was exported every 5 days.

Applying 'parcels' to velocity components on a C grid whilst using an RK4 advection scheme, with a Lagrangian timestep of 1 day, can lead to trajectories 'overshooting' the ocean grid cells into bathymetry. To reduce the occurrences of particles travelling out of bounds we added a criterion that returned such trajectories to 10 m above the seafloor⁷⁹. This criterion was applied to 4.7% of the total timesteps in the backwardtracked trajectories and for an average of 22.9% of timesteps for the forward-tracked trajectories, due to the trajectory start locations being in close proximity to seafloor topography, i.e. vent sites along ocean ridges.

Forward-tracking trajectories

Six potential source regions were identified (Fig. [1](#page-1-0)A, triangles) and referred to as the ECR, WCR, Easter Microplate (EM), NEPR, SEPR and PAR (Table S2). Trajectories were initialised over a 0.5° grid around each potential source location at 100 m intervals between 2000 and 4000 m. Two thousand trajectories were initiated per depth. In regions of bathymetry shallower than 4000 m, fewer trajectorieswere tracked. All calculations were normalised to the number of successfully tracked trajectories (Table S2) that were within a plume.We define the plume by the deepest depth of successful trajectories initialisation within each 0.5° release area, to mimic the deepest possible vent depth, up to 500 m shallower in the water column. Trajectories that crossed within a 0.5° area around the OOI site were isolated to estimate the dilution percentage of trajectories from the source regions (trajectory start locations) to OOI (Table S3). The EM had no trajectories passing near the OOI site and was not included in the analysis (Table S2).

Backward-tracking Trajectories

To complement the forward tracking we initiated trajectories throughout the water column at OOI to coincide with the depths at which the dFe and dMn signal was observed. Trajectories were initialised over a 0.5° grid around the OOI sustained observatory site every 50 m between 2000 and 4000 m. One thousand trajectories were released per depth and tracked backwards in time for 20 years (2011–1992; Table S2).

We defined five potential hydrothermal signal source regions along the PAR, EPR and CR to further evaluate where the iron signal was most likely to originate from—ECR and WCR as CR, NEPR, SEPR, PARN, and PARS (Table S4). Trajectories were only classed as originating from that source region if they entered at depths within a range of 500 m above the deepest possible vent height within each release area to mimic a plume $80,81$. For PARN, which had no specific vent release site for the forward-tracking simulation, a vent depth of 2500 m was chosen by examining the depths of the topography within the PARN area as a histogram to identify the ridge depths. The fraction of trajectories passing into each source region is presented as the dilution percentage (Table [1\)](#page-4-0).

Data availability

Lagrangian trajectories from the tracking using parcels are available at Zenodo [\(https://doi.org/10.5281/zenodo.8153763](https://doi.org/10.5281/zenodo.8153763)). The NEMO model velocity components used toforce parcels can befound on JASMIN [\(https://](https://www.ceda.ac.uk/services/jasmin/) www.ceda.ac.uk/services/jasmin/). The micronutrient data have been submitted to the British Oceanographic Data Centre [\(https://doi.org/10.5285/](https://doi.org/10.5285/0c7760cc-98fb-5861-e063-6c86abc0a998) [0c7760cc-98fb-5861-e063-6c86abc0a998](https://doi.org/10.5285/0c7760cc-98fb-5861-e063-6c86abc0a998)). The macronutrient data are available from the British Oceanographic Data Centre ([Series Document](https://www.bodc.ac.uk/data/documents/series/2121989/)— [BODC Document 2121989](https://www.bodc.ac.uk/data/documents/series/2121989/)).

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Author contributions

A.J.B. and C.A.B. wrote the initial draft and all co-authors contributed to draft revisions. A.J.B., C.A.B., N.J.W., K.P., H.J.V., C.M.M., I.T., A.M., S.J.U. and A.P.M. participated in the field campaign. A.J.B. and N.J.W. conducted trace metal (micronutrient) analysis. K.P. performed macronutrient analysis. H.J.V. processed conductivity-temperature-depth sensor package datasets. C.A.B. performed Lagrangian modelling experiments and analysis. S.O. carried out the water mass analysis. A.P.M., C.M.M., S.J.U. and A.M. wrote and planned the original research questions and funding proposal. A.P.M. led the research programme. The views expressed in this paper are those of the authors alone, and not the organisations for which they work.

Competing interests

The authors declare no competing interests.

Additional information

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