Release from biogenic particles, benthic fluxes, and deep water circulation control Cr and δ53Cr distributions in the ocean interior

David J. Janssen a,*, Jörg Rickli a,b, April N. Abbott c,1, Michael J. Ellwood d, Benjamin S. Twining e, Daniel C. Ohnemus f, Philipp Nasemann a, Delphine Gilliard a,g, Samuel L. Jaccard a,g

a University of Bern, Institute of Geological Sciences & Oeschger Center for Climate Change Research, Botanistrasse 1-3, 3012 Bern, Switzerland
b Institute of Geochemistry and Petrology, ETH Zürich, Zürich, Switzerland
c Department of Earth & Environmental Sciences, Macquarie University, Sydney, NSW, Australia
d Research School of Earth Sciences, Australian National University, Canberra, Australia
e Bigelow Laboratory for Ocean Sciences, East Boothbay, ME, USA
f University of Georgia, Skidaway Inst. Oceanography, Dept. of Marine Sciences, Savannah, GA, USA
g University of Lausanne, Institute of Earth Sciences, Lausanne, Switzerland

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A B S T R A C T
Chromium (Cr) has shown promise as a paleoceanographic proxy due to the redox-driven control of dissolved Cr concentrations ([Cr]) and stable isotope composition (δ53Cr). To improve the mechanistic understanding of Cr cycling in the modern ocean and strengthen its potential proxy applications, we present new data from regeneration incubations, bottom and sediment pore waters, and a compilation of intermediate and deep water data. While Cr removal and biological export from the surface ocean is associated with organic carbon export, the deep water release of dissolved Cr from sinking particles is not directly dependent on organic carbon respiration, as indicated by differing trends between Cr, oxygen utilization and the regeneration of organic-associated macronutrients (e.g. N, P). Pore water and bottom water data demonstrate that benthic Cr fluxes are locally important and may be significant globally. The pore water dissolved Cr flux at our CaCO3-rich site is likely driven by the re-release of Cr scavenged from the water column by sinking particles, with minor contributions from lithogenic phases. We argue this is consistent with the highest open ocean [Cr] to date being found in the water column below oxygen minimum zones, likely reflecting the release of scavenged Cr in deep waters or surface sediments. Chromium released from suspended particles and surface sediments follows the global δ53Cr–[Cr] array, supporting the proposed role of biological export and regeneration in shaping global Cr and δ53Cr distributions. Global intermediate and deep water [Cr], δ53Cr and Cr:macronutrient relationships are thus shaped by a synergy of circulation patterns, water mass mixing, a deep Cr regeneration cycle, and benthic Cr sources. A biogenic control on global Cr distributions indicates that sedimentary Cr records may reflect biogenic as well as O2-dependent processes, while more research is needed to assess sediment Cr record fidelity based on an active diagenetic cycle.

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1. Introduction

Oceanic distributions of dissolved Cr show nutrient-type behavior, generally resembling primary macro- and micronutrients, though with muted surface depletions and enrichments at depth (e.g. Campbell and Yeats, 1981; Jeandel and Minster, 1987). Thermodynamic calculations predict that oxidized Cr(VI), a soluble oxoanion with low particle reactivity (Semeniuk et al., 2016), should account for all dissolved Cr in the oxygenated ocean (Elder-
field, 1970). However, the reduced form, Cr(III), is regularly found at low levels in oxygenated seawater (e.g. Cranston, 1983; Jeandel and Minster, 1987; Achterberg and van den Berg, 1997; Janssen et al., 2020), and may constitute a major fraction of dissolved Cr in oxygen minimum zones (OMZs; e.g. Murray et al., 1983; Rue et al., 1997). Chromium(III) is relatively insoluble and readily adsorbs to mineral and organic surfaces (Cranston and Murray, 1978; Mayer et al., 1984). Therefore redox transformations play a central role in Cr biogeochemical cycling. Iron(II) is an important Cr(VI) reductant in natural aquatic systems (Pettine et al., 1998), and Mn oxides (MnOx) are a primary Cr(III) oxidant (van der Weijden and Reith, 1982; Milletto et al., 2021), especially in Mn-rich environments such as sediments (Oze et al., 2007).

The main dissolved Cr removal processes in the ocean, both involving Cr reduction followed by particle scavenging, are in OMZs and through biologically-mediated export (e.g. Murray et al., 1983; Scheiderich et al., 2015). Chromium removal in OMZs has been reported in the eastern tropical North Pacific (ETNP) (Murray et al., 1983; Rue et al., 1997; Moos et al., 2020) and the eastern tropical South Pacific (ETSP) (Nasemann et al., 2020). Correlations between Cr and particulate organic carbon in sinking particles (Connelly et al., 2006), Cr(III) adsorption onto phytoplankton (Semeniuk et al., 2016), and agreements between dissolved Cr deficits and productivity-based inferred removal (Janssen et al., 2020) support a Cr sink associated with biological export. However, the subtle gradients in [Cr] depth profiles suggest that release from biogenic particles is relatively minor (e.g. Scheiderich et al., 2015; Goring-Harford et al., 2018; Moos and Boyle, 2019, Rickli et al., 2019), and circulation plays an important role in shaping distributions (e.g. Scheiderich et al., 2015; Goring-Harford et al., 2018; Rickli et al., 2019). Elevated bottom water [Cr] suggests surface to deep [Cr] gradients may be influenced by a flux from marine sediments (Murray et al., 1983; Jeandel and Minster, 1987), consistent with shallow pore water [Cr] enrichments on the California shelf, assumed to reflect release from organic matter (Shaw et al., 1990).

Chromium redox transformations are accompanied by isotopic fractionation. While fractionation patterns during Cr oxidation are variable (Zink et al., 2010; Milletto et al., 2021), reduction consistently enriches light isotopes in Cr(III) (e.g. Wanner and Sonnenthål, 2013). The redox control on [Cr] and δ53Cr makes δ53Cr a potentially powerful tracer of paleoredox conditions, especially in early earth studies (e.g. Frei et al., 2009). However, application and interpretation of δ53Cr records require an accurate mechanistic understanding of the process(es) that control Cr budgets and isotopic fractionation in terrestrial environments, the ocean, and during incorporation into the sediment record. The systematic relationship between seawater [Cr] and δ53Cr suggests that redox transformations likely control oceanic distributions of δ53Cr (Scheiderich et al., 2015; Goring-Harford et al., 2018; Moos and Boyle, 2019; Moos et al., 2020; Rickli et al., 2019; Janssen et al., 2020; Nasemann et al., 2020) implying an effective isotope enrichment factor (ε) of approximately ~0.7‰ associated with Cr reduction and removal. Recent δ53Cr and [Cr] data demonstrate fractionation associated with biologically driven Cr removal in the open ocean (Janssen et al., 2020). Similarly, Cr removal in shelf settings (Goring-Harford et al., 2018), as well as in OMZs (Moos et al., 2020; Nasemann et al., 2020; Huang et al., 2021) can result in isotopic fractionation. However, isotope fractionation factors for these removal processes remain poorly constrained.

To address uncertainties in the modern ocean Cr budget and the role of biogenic processes in seawater δ53Cr distributions, we present dissolved [Cr], δ53Cr and [Cr(III)] from shipboard incubations, new intermediate and deep water [Cr] and δ53Cr from eight research expeditions in the Southern, Pacific and Atlantic Oceans, and pore water [Cr] data from the Tasman Sea.

2. Oceanographic background - Southern Ocean intermediate and deep waters and their advection into the Pacific and Atlantic Oceans

The Southern Ocean is the formation site for much of the global deep and intermediate waters, including Antarctic Intermediate Water (AAIW) and Upper and Lower Circumpolar Deep Water (U/L-CDW). The extent to which these waters penetrate into lower latitudes in the Pacific and Atlantic Oceans varies depending on formation sites, bathymetry, and the prevalence of other water masses.

AAIW is reflected in salinity minima in depth profiles, although this erodes with northward transport, and it is found at neutral densities ~27 < γN < ~27.4 (Orsi et al., 1995; Talley et al., 2011; Bostock et al., 2013). AAIW flows northward to the tropical South Pacific (~10-15°S) where it mixes with Equatorial Pacific Intermediate Water (Bostock et al., 2013). Atlantic AAIW is diluted by mixing during northward transport, as evidenced by increased salinity, and reaches a maximum extent slightly beyond the equator (Suga and Talley, 1995). Tropical Atlantic AAIW is found at a depth of around 700-800 m discernible by low salinity (~34.20-34.40) and high O2 (≥190 μmol kg⁻¹) south of 20°S.

The deep waters of the Pacific Ocean comprise UCDW, LCDW and PDW, the latter of which is largely found at similar densities as UCDW (γN < 28.0; T > 1.5° C; S > 34.65 PSU) and is formed as the colder, more saline and more oxygen rich LCDW (γN > 28.0) is upwelled and mixed throughout Pacific (Orsi et al., 1995; Kawabe and Fujio, 2010, Talley et al., 2011). Densities occupied by UCDW transition to predominantly PDW in the tropical South to subtropical North Pacific. LCDW extends into the North Pacific along the western side of the basin, and the northernmost reaches are characterized by elevated [Si(OH)₄]₂⁻ (≥170 μmol kg⁻¹) (Kawabe and Fujio, 2010). CDW enters the South Atlantic below AAIW and can be dominant as far north as ~25°S, but its relative contribution weakens as the water mass continues northward, with CDW accounting for ≤50% of deep waters in the tropical Atlantic (Larqué et al., 1997).

3. Methods

Intermediate and deep water [Cr] and δ53Cr samples were collected on RV Investigator expeditions IN2018_V02 (Mar-2018) and IN2018_V04 (Sept-Oct 2018), ACE Legs 1 (Dec-2016 to Jan-2017) and 2 (Jan-2017 to Feb-2017), GEOTRACES section GP13 (Australian leg, May-June 2011), RV Meteor cruise M77/4 (Jan-Feb 2009), Line P cruise 12-13 (Aug-2012), and RV Discovery cruise DY110 (Oct-Nov 2019) (Fig. 1). Further analyses were performed on bottom and pore waters (IN2018V_04) and on samples from shipboard regeneration incubation experiments (IN2018V_02).

3.1. Incubation design and sampling

Incubation regeneration experiments were carried out at the Southern Ocean Time Series (SOTS) site, in the open subantarctic Southern Ocean south of Tasmania (IN2018_V02, ~4600 m water depth) in acid-cleaned 20 L LDPE cubitainers with Teflon-lined caps. Three experiments were conducted, with differences in incubation temperature and the particle collection depth (supplemental material). All experiments used particles from below the chlorophyll max (~60 m depth) and at or below the mixed layer depth (~90 m depth). Each experiment consisted of two treatments, each in triplicate. Cubitainers were filled with filtered (0.2 μm) surface (~5 m) seawater collected using a trace metal-clean tow fish, and aliquots were collected for initial (TN) nutrients, [Cr] and δ53Cr before particle inoculation. Filled cubitainers were stored in the dark at 1 °C until inoculating with particles (within 8 hours of filling).
Ambient suspended particle samples from the oxic water column were collected onto acid-cleaned 1 µm polycarbonate filters using in-situ pumps (McLane). Particles were resuspended from these filters, forming a particle-rich slurry that was added to the individual cubitainers to result in particulate concentrations approximately 2–4 fold above natural levels. Because only resuspended particles were added to the incubations, the high Cr blanks associated with many filter materials (e.g. Scheiderich et al., 2015) will not impact the incubations.

Particle-inoculated cubitainers were incubated in the dark at either 4 °C (ambient) or 1 °C (cold treatment) for five days, with three subsampling time points. For subsampling, the cubitainers were transported to a Class 100 clean van, gently mixed, and subsampled into acid-cleaned LDPE bottles (for trace metals) or polycarbonate tubes (for macronutrients). Subsamples for trace metals were transported to a laminar flow hood and filtered through acid-cleaned 0.4 µm polycarbonate filters using a Teflon vacuum filtration apparatus (Savillex). Loss of Cr to container walls has been shown to be minimal over short timeframes (e.g. Semeniuk et al., 2016) and was not considered here. Chemically-labile particulate metal and P concentrations were determined with magnetic-sector ICP-MS following solubilisation with a heated mixture of reductant (hydroxylamine hydrochloride) and weak acid (acetic acid) designed to dissolve biogenic material and Mn and Fe oxides (Berger et al., 2008). Analytical procedures followed those previously described in Rauschenberg and Twinning (2015).

3.2. Bottom water, pore water and sediment sampling

Pore water and bottom water samples for this study were collected from a multicore deployment near Station P33 in the Tasman Sea (Fig. 1, 3340 m water depth). Cores for pore water, bottom water, and sediment samples were processed in an N2 environment. A bottom water sample was extracted from the core tube approximately 10 cm above the sediment-water interface using an acid-cleaned rubber-free syringe and filtering through a 0.45 µm syringe filter into an acid-cleaned LDPE bottle. The bottom water was visibly clear and free from sediment resuspension, and the sediment surface appeared undisturbed.

Pore water samples were collected following procedures outlined in Abbott et al. (2015). In brief, 1 cm depth intervals were placed into acid-cleaned centrifuge tubes in a glove bag, sealed, then centrifuged at 4500 rpm for 20 minutes. The pore water (supernatant) was extracted from centrifuged samples and filtered as outlined for bottom water. The samples were acidified, stored and analyzed for [Cr] by isotope dilution as described in section 3.6. Pore water subsamples were diluted 40x and [Fe] and [Mn] were determined by high resolution inductively coupled plasma–mass spectrometry (HR-ICP-MS) (Element XR, ThermoScientific) at the Australian National University. Centrifuged sediments were stored frozen until freeze-dried, then analyzed for bulk composition (CaCO₃, organic matter, lithogenic material; supplemental material).

3.3. New and literature data from the Southern, Pacific and Atlantic Oceans

We present a compilation of new and literature [Cr] data in Southern-sourced intermediate and deep waters across the Southern, Pacific, and Atlantic Oceans (Fig. 1). Literature [Cr] data were not included if: (1) hydrographical data (temperature, salinity) are missing, (2) the data have not been peer-reviewed, or (3) later publications identified them as suspect or incorrect.

We add 12 samples from the Indian Sector of the Southern Ocean (ACE Leg 1) and one sample from the Drake Passage (ACE Leg 2) to [Cr] and δ⁵³⁷Cr available from the Pacific Sector of the Southern Ocean (Rickli et al., 2019) to further constrain the composition of southern-sourced end-members. The Drake Passage sample constrains the characteristics of intermediate waters entering the Atlantic Ocean.

We present data from five stations in the subantarctic Southern Ocean and the Tasman Sea (IN2018V_02 and IN2018V_04, 26 sam-

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**Fig. 1.** Map of sampling locations. Station locations are shown for new data (open circles) and literature data (filled circles). Split circles indicate locations with both new and literature data. Full data tables can be found in the supplemental material. The black star denotes the location of the multicore site, where bottom and pore waters were collected. The northernmost extent of AAW as a primary water mass component in the Pacific Ocean (Bostock et al., 2013), is shown as a dashed black line. The inset shows the subantarctic Southern Ocean, the Tasman Sea (located between Australia and New Zealand) and the subtropical southwestern Pacific, the sites for most of the new data. The flow paths of AAW are shown in the inset as black arrows, and M denotes regions of enhanced AAW mixing (Bostock et al., 2013). Dashed white and grey lines indicate representative fronts of the Southern Ocean (white – the Subtropical Front, light grey – the Subantarctic Front, dark grey – the Polar Front; Orsi et al., 1995). (For interpretation of the colors in the figure(s), the reader is referred to the web version of this article.)
precipitation, and four stations in the Tasman Sea and subtropical South Pacific (GEOTRACES section GP13, 12 samples) in order to characterize waters advecting northward into the Pacific Ocean. Four new samples from below one of the most intense OMZs in the global ocean (ETSP near Peru; M77/4) are added to existing ETSP (Nasemann et al., 2020), ETNP OMZ (Murray et al., 1983; Rue et al., 1997), and subtropical North Pacific (Moos and Boyle, 2019) data. Additionally, we present four new samples from a profile in the subarctic North Pacific (Line P 2012-13) to characterize the northern extent of deep water in the Pacific Ocean. Previous subarctic North Pacific [Cr] literature data in PDW and LCDW (Cranston, 1983; Mugo and Orians, 1993) show a clear offset from recent high-precision data and are not used (see Table S1).

To follow the northward advection of AAIW into the equatorial Atlantic, we present seven samples from five stations along a meridional transect (DY110). Additional literature data from the Atlantic are limited, with only one sample within AAIW (station 115, Goring-Harford et al., 2018) and no available CDW data.

3.4. Water column dissolved Cr sampling

IN2018_V02, IN2018_V04, and GP13 samples were collected using an autonomous trace metal clean rosette system and gravity filtered through acid-cleaned 0.2 µm filters (Pall Acropak) into acid cleaned LDPE bottles in a class 100 clean van. IN2018 samples were acidified to pH < 2 (2 mL L−1 sub-boiling distilled concentrated HCl) on land and stored for 3–26 months before analysis. GP13 samples were stored frozen until 2019, at which point they were thawed, acidified as for IN2018 samples, and stored at least 7 months before analysis. DY110 samples were collected with a conventional CTD-rosette, gravity filtered through acid-cleaned 0.2 µm filters (Pall Acropak) acidified to pH = 2 (1 mL L−1 concentrated HCl), and stored for at least 3 months before analysis. Sampling and sample handling are discussed in detail elsewhere for ACE (Rickli et al., 2019), M77/4 (Nasemann et al., 2020) and Line P (Janssen and Cullen, 2015).

3.5. Cr(III) samples

Dissolved chromium redox speciation ([Cr(III)]) samples were processed at sea using magnesium hydroxide (Mg(OH)2) co-precipitation, then analysed by isotope dilution ICP-MS at the University of Bern following the procedures of Janssen et al. (2020) (external [Cr(III)] 1σ = 10% RSD, Janssen et al., 2020).

3.6. Chromium concentrations and δ53Cr

Procedures for the determination of total dissolved Cr concentrations ([Cr]) and stable isotope composition (δ53Cr) are described in detail in Rickli et al. (2019), with minor modifications (see supplemental material; Janssen et al., 2020; Nasemann et al., 2020). Briefly, preliminary [Cr] for spiking purposes was determined on small sample aliquots by isotope dilution with Mg(OH)2 co-precipitation followed by cation exchange chromatography. The final reported [Cr] is from 53Cr determinations, except when 53Cr samples were not available (pure waters and incubation intermediate time points). Pore water samples (0.2 mL) were dried rather than co-precipitated before chromatography. For 53Cr, a 0.25-1 L sample aliquot was spiked with 53Cr-53Cr double spike, pre-concentrated by Mg(OH)2 co-precipitation, and purified by a 2 step column chromatography.

Samples were analyzed on a ThermoFisher Neptune Plus MC-ICP-MS at the University of Bern following Rickli et al. (2019). 53Cr procedural blanks were 0.3–0.6 ng Cr and insignificant relative to sample Cr. 53Cr data are reported relative to NIST SRM 979. Internal uncertainties were generally 0.02–0.03% (2 SEM); external reproducibility, based on replicate analyses of seawater samples, is ±0.033 (2 SD) for δ53Cr and <1% (1 RSD) for [Cr] (see Janssen et al., 2020). A Merck Cr(III) standard was run to monitor mass-spectrometric accuracy (δ53Cr = −0.426 ± 0.027‰, n = 9; literature: δ53Cr = −0.443 ± 0.022‰, Schoenberg et al., 2008). The procedural accuracy of our method has previously been validated by an inter-laboratory comparison (Rickli et al., 2019), and the precision of seawater analyses during this study was confirmed through regular analysis of separate aliquots of OSIL seawater (δ53Cr = 0.89 ± 0.03, n=11; www.osil.co.uk).

4. Results and discussion

4.1. Regeneration incubation experiments

Three incubation experiments to probe for broad, differential effects on the regeneration of trace elements were conducted at the SOTS site (subantarctic Southern Ocean, Fig. 1) within 10 days of each other, with varying experimental design (particle collection depth and incubation temperatures) and natural variability in the initial particulate regime and elemental concentrations. Only one incubation (Inc. 3, particles from 150 m depth) generated significant differences in [Cr] and δ53Cr by the final time point. We focus on this incubation here, with results from Inc. 1 and 2 provided in the supplemental material (Section S1, Figure S1, Tables S3-S4). Initial particulate P (pP), Mn (pMn) and Fe (pFe) concentrations from all incubations are shown in Table S1.

Time series sediment trap data from SOTS indicate that fluxes are dominated by biogenic particles (>90%, total fluxes at 1000-3800 m = 13–21 g m−2 yr−1, Wynn-Edwards et al., 2020), with high contributions from coccolithophores (Wilks et al., 2017), while lithogenic contributions are minor (<0.7 g m−2 yr−1, Trull et al., 2001). Supportively, flow cam data from our incubations document a predominance of small particles (mostly ~2 µm, nearly all <6 µm), with a high abundance of flagellates (including coccolithophores, data not shown). Furthermore, POC profiles (Figure S2) and sediment trap data (Wilks et al., 2017; Wynn-Edwards et al., 2020) identify general stability in bulk particulate composition with depth from 150 m, indicating that the particles used here as broadly representative of local export.

There was minimal to no net regeneration of organic-associated macronutrients as indicated by the lack of macronutrient variability (N and P) (Fig. 2, Table S2). Chromium concentrations, [Cr(III)] and δ53Cr remained stable over the first three days. However, [Cr] increased by an average of 0.46 ± 0.14 and 0.56 ± 0.18 nmol kg−1 (1 SD of the triplicates) by the final sampling for the ambient and cold treatments, respectively (Fig. 2), demonstrating that Cr release was independent from organic carbon respiration. Small increases in [Cr(III)] were also observed by the end of the incubation, though [Cr] increases were approximately seven times larger. Particulate Cr was probably primarily Cr(III) – Cr(III) dominates biogenic particle-adsorbed Cr, with Fe oxides playing an important role for Cr(III) scavenging (e.g. Semeniuk et al., 2016). Although carbonates may also host Cr (as Cr(VI)), the Cr content of CaCO3 is very low (Remmelzwaal et al., 2019), and exported CaCO3 is largely preserved with depth at this site (Wynn-Edwards et al., 2020), making carbonates unlikely to drive the observed dissolved [Cr] accumulation. Therefore, due to the small changes in [Cr(III)], particulate Cr was either oxidatively released or oxidized shortly after release.

While more observational constraints would be needed for a detailed mechanistic framework of the processes regulating Cr oxidation, we explore likely oxidants based on available data and geochemical constraints from the literature. The primary oceanic Cr(III) oxidants are MnOx (e.g. van der Weijden and Reith, 1982;
Milletto et al., 2021) and H$_2$O$_2$ (Pettine et al., 1991). Other redox-active species (e.g. N), while present in the incubations, do not have known roles for Cr oxidation and are unlikely oxidants based on reduction potentials (Rue et al., 1997) and oceanic data showing correlations with NO$_3^-$ reduction with Cr reduction rather than Cr oxidation (Rue et al., 1997; Moos et al., 2020; Nasemann et al., 2020).

Particles in our incubations were enriched in Mn beyond typical cellular quotas (Table S1, Twining and Baines, 2013), indicating the presence of MnOx. We also observe increasing pMn between the first and second time points, indicating further enrichment in MnOx, consistent with mid-depth scavenging of Mn (e.g. Ohnemus et al., 2019). The timing of the Cr release in Inc. 3 matches a decrease in pMn (Fig. 2), supporting Mn reduction coupled to Cr oxidation and release, though the increase in Cr(VI) stoichiometrically exceeds that suggested by pMn and the Mn(IV)-Cr(III) redox couple. Since we observe an active cycle of Mn, with first increasing and then decreasing pMn, repeated microbial Mn(II) oxidation followed by MnOx reduction with Cr(III) oxidation may explain the [Cr] accumulation above expected reaction stoichiometry. This has
been observed in recent seawater culture experiments (Milletto et al., 2021); however, our data do not unequivocally indicate MnOx-driven cycling, and other potential oxidants may be involved (e.g. H₂O₂, for which data are not available).

Differences between the three incubations may also be related to metal oxides. Incubations from 150 m had much higher pMn than those from shallower depths (Inc. 1, 100 m & Inc. 2; Table S1), following depth trends in oceanic pMn (Ohnemus et al., 2019), and potentially explaining the lack of Cr release in shallower treatments. Particulate Fe in Inc. 1 was higher than in Inc. 3 (Table S1), indicating more Fe-rich mixed oxides, which may impact Cr oxidation and explain differences between these incubations. If particulate MnOx are important, the enriched particle concentrations in the incubations, possibly in combination with the formation of particle aggregates, would help to increase the probability of oxidative Cr release by MnOx relative to ambient seawater conditions. Consequently, if oxidative particulate Cr release is driven by reduction with MnOx, it may be facilitated by more Mn-rich oceanic particles at greater depth (Ohnemus et al., 2019) and in marine sediments, supporting a deeper regeneration cycle for Cr than organic matter.

The increase in [Cr] was accompanied by a statistically significant decrease in δ²⁵⁹Cr in four of six samples (Fig. 2, Table S2). δ²⁵⁹Cr decreases were not significant for the remaining two samples, consistent with the small increase in [Cr]. Inter- replicate variability is likely controlled by small initial differences in biological communities and particle compositions in each cubitainer, hence real natural variability, rather than analytical uncertainty, given differences in [Cr] between triplicates were well outside of analytical uncertainty (Fig. 2, Table S2). This release of low δ²⁵⁹Cr from particles provides evidence for predictions that Cr adsorbed onto biogenic particles is isotopically light (Scheiderich et al., 2015; Semeniuk et al., 2016; Janssen et al., 2020).

A strong linear relationship is observed between δ⁵³Cr and ln[Cr] in the regeneration incubation samples (r² = 0.89) with an implied enrichment factor (ε) = −0.66‰, matching that implied by the global δ⁵³Cr-[Cr] array (global ε ≈ −0.70‰, Fig. 2). This supports release from biogenic particles as an important process in driving global [Cr] and δ⁵³Cr distributions. While proposed earlier (Scheiderich et al., 2015), previous studies have highlighted that biogenic Cr accumulation was lacking strong support in oceanic depth profiles (Scheiderich et al., 2015; Goring-Harford et al., 2018).

### Table 1

<table>
<thead>
<tr>
<th>Potential sources</th>
<th>Sediment phase (% of bulk)</th>
<th>Cr per phase (ppm)</th>
<th>Cr contributed to bulk sediment (ppm)</th>
<th>Cr needed (ppm)</th>
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<td>CaCO₃</td>
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<td>1</td>
<td>0.3</td>
<td>18</td>
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<tr>
<td>Lithogenic material/Upper continental crust</td>
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<td>90</td>
<td>18</td>
<td>49</td>
</tr>
</tbody>
</table>

**Fig. 3.** Pore water [Cr] profile. The left panel shows pore water dissolved [Cr], [Mn] and [Fe] with depth in the sediment. Note that [Cr] and [Mn] are presented as nmol kg⁻¹, while Fe is presented as µmol kg⁻¹ with concentrations multiplied by ten to match axis scaling (i.e. 2 µmol kg⁻¹ Fe would correspond to the axis tick mark for 20 in this figure). Concentrations represent pore water concentrations in sample intervals of 1 cm. Dissolved bottom water [Cr] is shown from a sample collected approximately 10 cm above the sediment-water interface (shown as a dashed line). Uncertainty is smaller than the symbol size. Note that the y axis scale above the sediment-water interface is different from the scale below the interface. The right panel shows bulk sediment composition for the same core and sediment intervals as pore water data (CaCO₃ – grey, lithogenic material – brown, organic matter – green; see supplemental section 3.3 and Table S7). The x axis scale has a break between 0X and 76X.

Moos and Boyle, 2019; Riclli et al., 2019). This study provides the first direct evidence that Cr release from biogenic particles acts as a control on δ⁵³Cr across intermediate and deep waters, and ultimately also supports the inverse of this process – the adsorption of Cr onto biogenic particles in the surface ocean – as a control of δ⁵³Cr and [Cr] (Scheiderich et al., 2015; Semeniuk et al., 2016; Goring-Harford et al., 2018; Janssen et al., 2020).

4.2. Benthic Cr supply: the flux of Cr to bottom waters from pore waters

Pore waters were sampled from calcareous ooze sediments (~3350 m water depth) collected from a shallower feature punctuating the abyssal plain (~4000-5000 m) in the Tasman Basin (Fig. 1). Regional sedimentation rates are on the order of 1-2 cm kyr⁻¹ (Cochran and Osmond, 1976). Organic matter is around 1% at the sediment surface, though upper sediments remain oxic and organic carbon content decreases with depth (Table 1, Table S7). Surface sediment Mn/Al (~0.016) and Fe/Al (~0.53) are higher than upper continental crust values (0.009 and 0.48 respectively, Rudnick and Gao, 2003; Table S7), suggesting slight authigenic oxide enrichment.

Pore water dissolved [Cr] shows a shallow sub-surface maximum (0-1 cm below the sea floor) of 47.4 nmol kg⁻¹, approximately an order of magnitude greater than local bottom waters (4.81 nmol kg⁻¹, Fig. 3), and pore water [Cr] decreases with depth. Pore water [Mn] and [Fe] do not show µM level enrichments, reflectingoxic sediment conditions (Fig. 3). The lowest pore water [Cr] is observed in the deepest samples (6.7 nmol kg⁻¹ at 9-10 cm depth), which remain elevated relative to bottom water. This upper pore water [Cr] maximum is consistent with observations by Shaw et al. (1990), though in contrast to our data, Shaw et al. (1990) observed Cr removal in the uppermost oxic sediments, above near-
surface pore water [Cr] maxima. The absence of Cr removal in our uppermost oxic sediments may reflect the strong differences in bulk sediment composition between lithogenic-dominated sites (Shaw et al., 1990) and our carbonate-dominated site.

A near-surface pore water [Cr] maximum suggests pore waters act as a diffusive source of dissolved Cr to the ocean. Bottom waters composed of CDW are enriched in [Cr] and are isotopically distinct ([Cr] = 4.81 nmol kg⁻¹, δ⁵³Cr = 0.76 ± 0.03‰) from CDW at nearby stations not in contact with sediments ([Cr] = 3.95 nmol kg⁻¹, δ⁵³Cr = 0.84 ± 0.03‰, 2SD; n = 6 samples ≥3000 m at stations TS8 and PS2, Table S9). Given the gradient of [Cr] from the pore water to the overlying bottom water, the diffusive flux of dissolved Cr can be estimated using Equation (1)

\[
\text{Flux}_{\text{Cr}} = D_i \frac{\Delta C}{\Delta z}
\]

where \(D_i\) is the Cr diffusion coefficient corrected for temperature and tortuosity (supplemental material, see also Abbott et al., 2015), giving an estimated benthic flux of ~3 nmol Cr cm⁻² yr⁻¹.

To contextualize this, if global oxic sediments were characterized by a benthic source of this magnitude, the global benthic Cr flux would be comparable to or larger than riverine inputs, currently believed to be the dominant Cr source (Bonnand et al., 2013; supplemental material), consistent with box model estimates (Jeandel and Minster, 1987). While quantitative estimates of global benthic Cr fluxes await greater data availability across diverse sediment types, our data identify that benthic sources are at least locally important and support previous studies that: (1) have argued for benthic Cr fluxes based on globally distributed elevated deep water [Cr] (Cranston, 1983; Murray et al., 1983; Jeandel and Minster, 1987), and (2) have invoked elevated pore water [Cr] to explain Cr enrichments in sedimented planktonic foraminifera compared to water column samples (Rennelzwael et al., 2019).

We use bulk sediment composition to examine potential sources of Cr to pore waters by applying mass balance calculations based on the average composition of the upper 4 cm, reported Cr concentrations of these phases, and regional sedimentation rates (Table 1). These calculations suggest that neither lithogenic, carbonate-hosted, nor surface phytoplankton Cr can account for the Cr flux out of these carbonate-rich and detrital-poor sediments (Table 1). Therefore, we suggest a combination of mechanisms causing Cr enrichments in particles delivered to sediments can explain the observed pore water data. First, respiration may result in relative Cr enrichments in particles reaching the seafloor because the release of Cr from particles appears decoupled from organic matter respiration (see sections 4.1, 4.3). At the same time, scavenging of particle-reactive Cr(III) may increase the Cr content of biogenic and non-biogenic particles. Regardless of which process(es) are involved, the Cr released from these sediments must largely be derived from the water column based on mass balance and is incorporated into the sediments as particle-adsorbed Cr that has been scavenged onto particle surfaces in the upper ocean (Connelly et al., 2006; Semeniuk et al., 2016; Janssen et al., 2020), and at intermediate depths (see section 4.3), while detrital Cr may be more inert. The similar [Cr] and [Mn] profiles in upper pore waters (Fig. 3) may reflect related geochemical controls (e.g. oxidative release of Cr(III) coupled to Mn reduction).

By mass balance calculations using observed bottom and deep water [Cr] and δ⁵³Cr (bottom water: [Cr] = 4.81 nmol kg⁻¹, δ⁵³Cr = 0.76 ± 0.03‰; deep water: [Cr] = 3.95 nmol kg⁻¹, δ⁵³Cr = 0.84 ± 0.03‰, 2SD; supplemental material), we estimate pore water δ⁵³Cr at 0.34 ± 0.25‰ (2 SEM propagated error), which is heavier than silicate earth (~0.124 ± 0.101‰, Schoenberg et al., 2008). While the closeness of these ranges indicates that some lithogenic component may contribute to the pore water flux, these data and the likely refractory nature of lithogenic Cr (e.g. Bauer et al., 2019), argue against significant lithogenic Cr contributions to the pore water flux. In contrast, pore water δ⁵³Cr is lighter than dissolved δ⁵³Cr in the upper 1000 m ([δ⁵³Cr ≈ 1.1–0.9‰] by a similar amount as the enrichment factor predicted by the global δ⁵³Cr–[Cr] array (ε ≈ −0.7‰). In other words, removal of dissolved Cr to particles in the upper 1000 m with δ⁵³Crparticulate-dissolved following the global array would result in δ⁵³Crparticulate similar to the calculated δ⁵³Cr_pore water.

Although the exact origin of pore water Cr is hard to discern, our mass balance calculations highlight the importance of internal processes acting to redistribute Cr within the water column, transferring dissolved Cr from upper waters to deep waters. Therefore, the benthic flux may act more as an attenuation of Cr removal associated with particle export than an entirely new Cr source. Pore water data from diverse sediment types are needed to determine whether a truly new contribution from lithogenic material may be regionally important in more lithogenic-rich sediments.

4.3. Accumulation of Cr in intermediate and deep water masses

To investigate the oceanic implications of the mechanistic insights from our incubation and pore water findings, a compilation of globally-distributed [Cr] data is presented along with macronutrients and apparent oxygen utilization (AOU, a quantification of the O₂ used for organic matter respiration) following the northward advection of Southern Ocean water masses (Fig. 4). Intermediate and deep waters are split into three water mass ranges based on hydrographic properties: AAIW, UCDW/PDW and LCDW (Figure S3, Tables S9–S10).

Deeper waters (UCDW and LCDW) show relatively uniform [Cr], AOU and macronutrients at their southern origin, and concentrations increase with northward transport (Fig. 4). However, key differences emerge between tracer primarily reflecting organic matter respiration (AOU, PO₄), phytoplankton frustule-associated Si(OH)₄, and [Cr]. AOU and PO₄ consistently show latitudinal maxima in the upper deep water mass (UCDP/PDW) relative to LCDW (Fig. 4). However, Si(OH)₄ shows maxima in LCDW, and [Cr] in LCDW is comparable to or higher than in UCDW. These deeper maxima, illustrating an apparent deeper regeneration cycle, may reflect globally important benthic sources as shown for Si (Tréguer and de la Rocha, 2013), consistent with our calculated pore water fluxes and earlier global Cr models (Jeandel and Minster, 1987). Chromium–Si(OH)₄ distributions show a stronger correlation (\(r^2 = 0.58, n = 65\)) than Cr–AOU (\(r^2 = 0.44, n = 71\)) and Cr–PO₄ (\(r^2 = 0.52, n = 67\)) (Fig. 5), supporting Cr release from biogenic material as being mechanistically independent from organic matter respiration.

The highest deep water [Cr] to date is found below the ETSP OMZ, where accumulated Cr represents up to ~45% of total deep water [Cr] based on Southern Ocean end members. While suboxic sediments are a net sink term (Moos et al., 2020; Nasemann et al., 2020), these deep water [Cr] enrichments may reflect proximity to a benthic source from deeper oxic sediments (section 4.2, Fig. 1). Deep ETNP samples are also enriched relative to the subtropical South Pacific and subarctic North Pacific (Fig. 4), suggesting a connection to the intense OMZs overlaying these deep [Cr] enrichments (Murray et al., 1983; Rue et al., 1997; Moos et al., 2020). Mechanistically, enhanced water column [Cr] removal and export of particulate Cr to depth combined with elevated pMn present below these OMZs (e.g. Murray et al., 1983) would facilitate the oxidative release of Cr from particles in the water column or oxic sediments. This Cr-specific enrichment process would cause deep waters below intense OMZs to deviate from Cr–macronutrient correlations, which is confirmed by the resulting strengthening of the Cr correlation with Si (\(r^2 = 0.73\)) when removing samples below the ETSP
Fig. 4. Chromium and macronutrient concentrations and AOU along intermediate and deep water circulation. Modification of deep water \([\text{Cr}]\) is shown, along with parameters associated with the regeneration of organic material \([\text{PO}_4]\), AOU and dissolution of plankton skeletal/frustule phases \([\text{Si(OH)}_4]\). Panels on the left show AAIW (circles – Southern and Pacific Ocean data, diamonds – Atlantic Ocean data) while panels on the right show Southern and Pacific Ocean UCDW, PDW and LCDW, with UCDW and PDW in circles and LCDW in squares. Open circles denote samples with Antarctic Surface Water or Winter Water contributions. Colors match sampling locations shown in Figs. 1 and S3, and full data and sources are shown in Tables S8-S10. Direct comparisons between \([\text{Cr}]\) and AOU, \text{PO}_4 and \text{Si(OH)}_4 are shown in Fig. 5. Note the different x axis ranges for left and right panels.
OMZ (correlations with PO$_4$ ($r^2 = 0.56$) and AOU ($r^2 = 0.41$) remain similar; Fig. 5).

Intermediate waters are associated with high carbon respiration with northward advection (Fig. 4). Southern waters with properties similar to newly-formed AAIW show the highest [Cr], and concentrations are generally stable or decrease with northward transport, indicating [Cr] is decoupled from organic matter respiration. A general northward decrease in [Cr] is found for Atlantic AAIW, accompanied by increasing salinity ($r^2 = 0.54$). This trend likely reflects mixing of AAIW with Cr-poor but more saline thermocline waters and NADW (Fig. 6, Rickli et al., 2019). However, the weakness of the correlation likely suggests more complicated mixing of multiple water masses, variable Cr scavenging in intermediate waters, or both.

4.3.1. Implications of water mass [Cr] accumulation trends on Cr–macronutrient coupling in the global ocean

Widely observed Cr–macronutrient correlations suggest shared biogeochemical controls (e.g. Campbell and Yeats, 1981; Cranston, 1983; Jeandel and Minster, 1987; Rickli et al., 2019; Nasemann et al., 2020). Recent studies of other metal–macronutrient associations have highlighted the importance of three dimensional processes, rather than vertical-controlled simplifications, with pre-formed relationships, scavenging, and variable metal:macronutrient uptake ratios across taxa and nutrient regimes (e.g. Quay et al., 2015; Vance et al., 2017; Ohnemus et al., 2019) driving globally-correlated distributions.

While correlations between [Cr] and PO$_4$ exist on different spatial scales (e.g. Cranston, 1983; Rickli et al., 2019), biogenic
Cr fluxes are low relative to other nutrient-type metals and macronutrients as apparent in smaller surface [Cr] depletions, lower surface-to-deep [Cr] gradients, reduced [Cr] accumulation in global deep water, and [Cr] accumulation in subsurface waters that is independent from organic matter respiration (sections 4.1–4.3, Fig. 5). Therefore observed associations between [Cr] and PO4 probably reflect mixing between distinct oceanic end-members (see Rickli et al., 2019) rather than coupled vertically-driven processes (e.g. particle sinking and regeneration). Consequently, end-member pre-formed Cr:macronutrient ratios and global circulation would largely shape Cr–macronutrient relationships in the ocean interior and their variability, with end-members impacted by biological uptake at their formation sites.

A prominent benthic source of dissolved Cr provides a mechanistic explanation for [Cr] and Si(OH)4 coupling by vertical processes in addition to water mass mixing. However, deep water data demonstrate that [Cr] and Si(OH)4 can decouple (Figs. 4–5), and local differences in Cr–Si(OH)4 slopes (Jeandel and Minster, 1987) suggest the relationship between Cr and Si(OH)4 may reflect a combination of advection and mixing of waters with different pre-formed Cr:Si(OH)4 ratios, Cr and Si(OH)4 drawdown by phytoplankton in surface waters, and the impact of sediment type and location on benthic dissolved Cr and Si(OH)4 fluxes (e.g. elevated [Cr] below OMZs, Figs. 4–5). Based on these variable controls, and recognizing that circulation likely plays a central role, interpretations of Cr–macronutrient trends within depth profiles should be made with caution.

5. An updated global ocean Cr biogeochemical cycle

The foundations for understanding oceanic δ53Cr distributions were first presented by Scheiderich et al. (2015), who demonstrated a tight coupling between [Cr] and δ53Cr. Building on the [Cr] literature, Scheiderich et al. (2015) hypothesized that δ53Cr distributions reflected the known controls on [Cr]: reduction and removal in OMZs and Cr export with biogenic particle fluxes along with regeneration from biogenic material. Subsequent δ53Cr research has attempted to assess these mechanisms and their associated fractionations to clarify their roles in the global Cr and δ53Cr cycle. We combine these initial hypotheses and recent advancements with our new data to improve our mechanistic understanding of Cr cycling in the global ocean and highlight remaining uncertainties. This set of mechanistic controls defines how internal oceanic processes regulate δ53Cr, forming the global relationship (Fig. 7 panel A), and will help to guide paleoceanographic applications of marine δ53Cr.

Here, we have demonstrated that Cr release from biogenic particles, either in the water column oxic sediments, can explain the [Cr]-rich side of the array (Fig. 7B) and C) consistent with previous work on biological uptake shaping the [Cr]-depleted side of the δ53Cr–[Cr] array (Fig. 7B). Goring-Harford et al., 2018; Janssen et al., 2020). In OMZs, dissolved Cr can be scavenged in the water column (Moos et al., 2020) and at the sediment surface (Moos et al., 2020; Nasemann et al., 2020), a process that largely follows the global array (Fig. 7D). Mixing will generally act to homogenize process- and source-induced variability, with the specific effect depending on the signature of mixing water masses (Fig. 7E, see also section S4, Figure S4 and Rickli et al., 2019). Hydrothermal circulation may also impact [Cr] and δ53Cr in the modern ocean and paleoceanographic interpretations (Holmden et al., 2016), but remains largely unconstrained at present (Fig. 7 panel F).

Coastal environments are more likely to deviate from the global δ53Cr–[Cr] array due to localized influences, including shelf sources (Goring-Harford et al., 2018) (Fig. 7 panel C), more quantitative removal in shelf OMZs (Nasemann et al., 2020, Fig. 7 panel D); dilution from meltwater (Scheiderich et al., 2015, Fig. 7 panel E), and local riverine inputs (Fig. 7 panel F). Consequently, reconstructions of seawater δ53Cr from coastally-sourced marine sediment records, including continental margins, should be approached with caution as these records may not reflect global ocean conditions for δ53Cr and/or [Cr].

Taken together, the available body of oceanic data indicates that Cr and δ53Cr distributions are controlled by biological adsorption and scavenging onto sinking particles acting across broad spatial scales, with local enhancement in OMZs, and regeneration from particles in the water column and/or oxic sediments. These signals are then transported and mixed through ocean circulation. Consequently, δ53Cr records in marine-origin sediments should reflect the combination of these processes, with both export productivity and OMZ reduction contributing to Cr accumulation in sediments and with temporal δ53Cr records reflecting modification of these processes over time and with changes in global climate, rather than exclusively reflecting changes in O2 availability. Additional research is needed to understand early sediment diagenesis, as indicated by elevated pore water [Cr], and how this may alter sediment Cr and δ53Cr signals.

6. Conclusions

Our incubation data provide the first direct evidence of Cr release from biogenic particles and demonstrate that this release, which may be driven by Cr oxidation, is mechanistically independent from the regeneration of major elements in organic matter (e.g. C, N, P). Based on Cr geochemistry and our data, a likely Cr oxidant is MnOx, though we cannot conclusively exclude other potential oxidants. Our pore water data indicate large benthic Cr fluxes can be locally important, and are possibly a globally-important component of the ocean’s Cr budget. This benthic flux likely reflects the release of Cr scavenged onto particles in the water column, rather than ‘new Cr’ from the dissolution of lithogenic material. δ53Cr data from both bottom waters and incubations identify that release from biogenic particles follows the global δ53Cr–[Cr] array, demonstrating that Cr release, either in deep waters or surface sediments, can explain the high [Cr] end-member of the global δ53Cr–[Cr] array. Furthermore, the data confirm that biogenic export from the surface ocean and release at depth is important in shaping distributions of both [Cr] and δ53Cr throughout the global ocean. A global compilation provides clear evidence of [Cr] accumulation in deep waters, supporting a deep regeneration cycle for Cr that is decoupled from organic matter respiration, includes a potential benthic source, and is enhanced below tropical
Pacific OMZs due to release of OMZ-scavenged Cr in underlyingoxic waters and sediments. Intermediate waters show stable ordecreasing [Cr] with northward advection, reflecting minimal releasefrom particles and a potential net water column Cr sink via scavenging.Because both [Cr] and δ53Cr are impacted by biological andphysical processes, and pore water data indicate active cyclingin near-surface sediments, the role of these processes with respect toCr paleoredox proxy viability should be incorporated into futureδ53Cr O2-reconstruction applications.

CRediT authorship contribution statement

DJJ led the design of the study along with SLJ, with assistancefrom MJE, who organized trace metal sampling operations for ACELeg 1 & IN2018V_O2 and all operations for IN2018V_04, BST &DCO, who designed and conducted the shipboard incubations, andANA, who planned coring operations and conducted sediment andpore water sampling for cruise IN2018V_04. DJJ conducted seawatersampling, processed, and analyzed samples from ACE Leg 1, IN2018V_02, IN2018V_04, and Line P cruise 2012-13. SLJ conductedsampling for ACE Leg 2. DJJ processed and analyzed samplesfrom DY110 and ACE Leg 2. MJE assisted in seawater samplingfor ACE Leg 1, IN2018V_02 and IN2018V_04. BT and DO analyzedparticulate samples from IN2018V_04 regeneration experiments.PN processed and analyzed samples from M77/4. DJJ and DGprocessed and analyzed samples from GP13. MJE analyzed POC andpore water [Fe] and [Mn] samples. DJJ led interpretations and writinand all authors contributed to the manuscript.

Declaration of competing interest

The authors declare that they have no known competing financialinterests or personal relationships that could have appeared toinfluence the work reported in this paper.

Data availability

Full cruise CTD and macronutrient data for cruises IN2018V_02and IN2018V_04 are available from CSIRO ([https://www.cmar.csiro.au/data/trawler/]). Full macronutrient and CTD data from the ACE transect are available from the ACE collection on Zenodo ([https://zenodo.org/communities/spi-ace]). Full macronutrient and CTD data
from the M77/4 cruise are available from the Pangaea repository (https://www.pangaea.de/). Full Line P macronutrient and CTD data are available from the Line P website (http://www.waterproperties.ca/linep/data.php). Shipboard incubation data are available in Tables S1-S4. Pore water data are available in Table S5 and sediment data are available in Table S7. New and literature data used for the isopycnal compilation are available in the supplemental material (Tables S9 and S10). All new Cr data in this manuscript are also available in an open access dataset in the Zenodo repository (https://doi.org/10.5281/zenodo.5408613).

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