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Cadmium isotope variations in Neoproterozoic carbonates – A tracer of biologic production?

S.V. Hohl^{1,2*}, S.J.G. Galer³, A. Gamper⁴, H. Becker¹

Abstract

32

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Cadmium concentrations and stable isotopic compositions in seawater are important tools for studying the biogeochemical cycling of Cd in the modern oceans and as a proxy for micronutrient utilisation by phytoplankton. It is now well established that Cd isotopes become "heavier" as the primary production in the surface ocean increases, even though the mechanism driving the isotopic fractionation is still debated. Here, we use this property of Cd isotopes to examine changes that took place in the oceans during the emergence of multicellular life in the Neoproterozoic. Isotopic compositions and concentrations of Cd, N and C are reported in shallow-water carbonates of Ediacaran age from the Xiaofenghe section on the Yangtze Platform, South China. The Cd isotope data - reported as $\epsilon^{112/110}$ Cd - show positive excursions in the cap dolomites, while significantly lighter Cd is found in the overlying strata. After correction for salinity-controlled fractionation into inorganic calcite, calculated palaeo-seawater $\epsilon^{112/110}$ Cd_{sw} range from -2 to +1.5, overlapping values of modern surface seawater. Importantly, $\epsilon^{112/110}$ Cd_{sw} and δ^{13} C show a general positive correlation, as would be expected in bio-productive environments. However, the trend to lighter $\epsilon^{112/110}$ Cd up-section is not that explicitly expected for an "explosion of life" at the end of the Ediacaran. The upper Doushantuo also displays substantial fluctuations in REE abundances, δ^{15} N and δ^{13} C, which may be due to estuarine mixing. Our data suggest that the variations in $\epsilon^{112/110}$ Cd are a result of biologically-induced fractionation in at least some of the Ediacaran carbonates at Xiaofenghe. Further Cd isotope fractionation processes are clearly playing a role as well, such as precipitation of sulphides under anoxic pore-water conditions and fractionation into inorganic carbonates under variable salinity conditions. These effects have to be evaluated carefully when using Cd isotope systematics in ancient marine carbonates to look for palaeo-productivity signals.

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Introduction

Letter

The oceans underwent some of the most remarkable transformations in Earth's history in the Neoproterozoic, such as the appearance of animal life (Knoll, 2015), major excursions in sulphur and carbon isotope records (Kaufman and Knoll, 1995; Halverson and Hurtgen, 2007) and indications of a transition from an anoxic to a partially oxygenated deeper ocean in the aftermath of the Marinoan glaciation (Planavsky *et al.*, 2010). These changes can arguably be best studied in marine sediments from the Yangtze Platform, South China, that cover the critical time span from the Cryogenian to the Precambrian-Cambrian (PCC) boundary which saw the appearance of the first macroscopic fossils (*e.g.*, Yin *et al.*, 2007). For this reason, it may also be an ideal place to evaluate whether Cd isotopes in marine sediments can be used as a palaeo-productivity proxy. To test this, we analysed shallow-water carbonates from the Ediacaran Xiaofenghe section that was deposited on the S-SE-facing passive margin of the Yangtze Craton and hosts abundant fossil assemblages of multicellular life from the Doushantuo and Dengying Formations (Xiao *et al.*, 2012).

Stable isotope fractionation of Cd is a new proxy for studying biogeochemical cycling of micronutrients in the present-day oceans (Lacan et al., 2006; Ripperger et al., 2007; Abouchami et al., 2011) and in sedimentary archives, such as Fe-Mn crusts (Schmitt, et al., 2009; Horner et al., 2010). Seawater profiles of Cd show a nutrient-like behaviour, with strong near-surface depletions and deep-water enrichments, mimicking profiles of macronutrient phosphate (Boyle et al., 1976). The surface depletions in Cd are associated with "heavier" Cd stable isotopic compositions, which are thought to be due to preferential uptake of the "lighter" isotopes during incorporation of Cd by phytoplankton (Lacan et al., 2006; Ripperger et al., 2007; Abouchami et al., 2011). At the moment, there is no general understanding of why cadmium, which is toxic to most life, should be taken up. Zinc is required by all organisms and in the case of phytoplankton is the cofactor in zinc carbonic anhydrase (Zn-CA), an enzyme involved in photosynthesis. Under zinc-poor conditions, Cd can substitute for Zn in Zn-CA and, furthermore, Price and Morel (1990) have described a cadmium-bearing carbonic anhydrase variant (Cd-CA) with the same functionality; but Cd-CA is only known from a few diatom species (Park et al., 2007) whose first appearance lies in the Mesozoic. Studying Cd systematics in Proterozoic sediments could therefore possibly provide insights into the lineage of the CA enzymes. It is equally possible that cadmium fulfills no biological role at all, in most cases, and that Cd uptake is driven by absorption onto organic matter or that Cd is pumped into the cell through the membrane, mistaken for Zn.

Similarly, it remains unclear what the precise mechanism is for causing the isotopic fractionation of Cd during uptake from seawater. Pure inorganic absorption (Wasylenki *et al.*, 2014), partitioning into calcite (Horner *et al.*, 2011) and biological utilisation are known to favour the light isotopes, which would all be consistent with the observation of heavy Cd in surface seawaters. But irrespective of these uncertainties concerning the reason for uptake and site of fractionation,



^{1.} Institut für Geologische Wissenschaften, Freie Universität Berlin, Germany

^{2.} State Key Laboratory for Mineral Deposits Research, Department of Earth Sciences, Nanjing University, Nanjing, China

^{*} Corresponding author (email: shohl@zedat.fu-berlin.de)

^{3.} Max-Planck-Institut für Chemie, Abteilung Klimageochemie, Mainz, Germany

^{4.} Museum für Naturkunde Berlin, Leibniz Institute for Evolution and Biodiversity Science, Berlin, Germany

Cd isotopes can be used as a convincing proxy for productivity in surface waters in the sense that the greater the depletion in Cd (and macronutrients like phosphate), the isotopically heavier the Cd becomes that is left behind in seawater.

So far the oldest sediments analysed for their Cd isotopic compositions are Permian in age (Georgiev *et al.*, 2015). Studying Cd isotopes in our Ediacaran-age carbonates could potentially document palaeo-seawater biogeochemical cycling of cadmium, providing insights into evolution of multicellular life at the PCC boundary. Furthermore, Cd isotopes may also be used to support interpretations from other bio-available isotopic tracers. In recent years, δ^{15} N has provided important insights into palaeo-ocean nutrient-cycling and redox conditions (Ader *et al.*, 2014). As the δ^{15} N proxy is often affected by post-depositional modification (Bebout and Fogel, 1992), combining with Cd isotopes is potentially useful in screening such datasets. Nevertheless, when analysing Cd isotopes in ancient sediments, other factors that may mask the true seawater signal must be carefully evaluated, such as fluid flow alteration and additional inorganic fractionation processes (*e.g.*, variable salinity, authigenic sulphide precipitation).

Results and Discussion

34

Stable isotope measurements of C and N were performed on carbonates and bulk rock using standard methods. Stable Cd isotope compositions on acetic acid leachates from alteration-free carbonates were determined using a double-spike method and TIMS (see Schmitt *et al.*, 2009; Abouchami *et al.*, 2011; detailed analytical protocols in Supplementary Information).

The $\delta^{13}C_{org}$ values obtained up the stratigraphic section remain relatively constant at -28 ‰ before decreasing to -34 ‰ in black shales of the middle Doushantuo, where TOC values increase to 1.1 %. These values are characteristic of enhanced planktonic productivity and remineralisation and resemble previously published data from the Doushantuo (*e.g.*, Ishikawa *et al.*, 2013). In the cap dolostones $\delta^{13}C_{org}$ and $\delta^{13}C_{carb}$ are decoupled ($\Delta^{13}C = 25 vs. 33$ in overlying strata, Fig. 1c) and this is mimicked in a shift in average bulk $\delta^{15}N$ values from +1.6 ‰ to +4.2 ‰ (Fig. 1f), possibly indicating a change in the nutrient regime to a NO₃⁻-dominated marine environment (Ader *et al.*, 2014). In the Doushantuo Formation, nitrate presumably became the main utilised nutrient for primary producers while N₂ partly remained unused in surface-waters, where incomplete denitrification shifts seawater to higher $\delta^{15}N$. This may possibly be attributable to increased oxygen availability, which is essential for organic matter (OM) remineralisation via nitrification-denitrification (Canfield *et al.*, 2010); however, smaller second-order variations in the $\delta^{15}N$ curve might simply be due to alteration.

Cadmium isotopic compositions in carbonates vary by more than four $\epsilon^{112/110}Cd$ units and show no definite correlation with Cd (Fig. 2a) nor K concentrations (see Fig. S-1b). Cadmium concentrations in detrital material are very



Letter



Figure 1 Isotope and concentration data from Xiaofenghe section. Samples are drawn in equidistance according to their sample numbers in Table 1 for better visibility. (a-c) Carbon isotope data. (d) Carbonate $\epsilon^{112/110}$ Cd (error bars = 2 σ), grey bar = discrimination line for modern seawater values (Ripperger et al., 2007). (e) Salinity-corrected $\epsilon^{112/110}$ Cd of seawater. (f) Bulk δ^{15} N, yellow envelope = 0.2 % uncertainty, grey bar = modern surface water. (g,h) Shale-normalised Y/Ho and Ce/Ce* in carbonates, grey bars represents discrimination of modern seawater values (Bau et al., 1995) and negative/positive Ce anomalies.

low, so the effect of detrital input on $\epsilon^{112/110}$ Cd is likely negligible in extent. Late fluid-flow overprint is typical for many Neoproterozoic carbonates and may also have altered the Cd isotope signatures. However, Hohl *et al.* (2015) have shown that, with the exception of the cap dolostones, carbonates at Xiaofenghe were not much affected by fluids (SEM pre-screening, no co-variation of δ^{18} O with 87 Sr/ 86 Sr or Mn/Sr); thus, their trace element budgets and isotopic compositions most likely reflect partitioning and incorporation from Ediacaran seawater. In the cap dolostones $\epsilon^{112/110}$ Cd > 0, while low Cd and N concentrations (Fig. S-1d) correlate with light $\delta^{13}C_{carb}$ (Fig. 1c) and high Mn/Sr ratios (see Fig. S-1a), commonly used as indicators of fluid overprinting (Brand and Veizer, 1980).

Table 1	Stable isotope data; N and C concentrations; assorted shale normalised REE ratios and Mn
enrichmen	ts relative to Cal-S from Hohl et al. (2015b). D1-D4 = Doushantuo Fm., DG = Dengying Fm.

sample	Height [m]	Member	$\delta^{13}C_{carb}{}^1$	$\delta^{18}O_{carb}{}^1$	δ ¹³ Corg	$\Delta \delta^{13}$	ε ^{112/110} Cd ²	2SE	Cd [µg/g]
1	0.2	D1	-3.8	-6.7	-28	24.2	0.09	0.2	0.06
2	0.75	D1			-26.7		0.06	0.62	0.1
3	1.6	D1	-3.7	-7.7			0.46	0.23	0.03
4	2.3	D1	-3.8	-7.3	-26.7	22.9	0.25	0.19	0.07
5	3.85	D1	-3.2	-6.9			0.18	0.39	0.03
6	4.8	D1	-1.2	-12.3	-28.4	27.2			
7	6.65	D2	0.0	-6.8	-25.4	25.4			
8	12.1	D2			-28.1				
9	22	D2	0.6	-10.4	-29	29.6	1.07	0.35	0.03
10	36	D2					-1.69	0.09	0.2
11	37.5	D2			-28.3				
12	65	D2	6.5	-6.1	-29	35.5			
13	73.7	D2			-25.5				
14	75	D2	5.1	-4.8	-27.3	32.4			
15	75.6	D2	5.4	-4.4	-26.8	32.2	-2.04	0.25	0.04
16	76	D2	5.2	-4.5	-28.1	33.3	-1.11	0.25	0.05
17	76.5	D2							
18	77.5	D2	6.2	-4.1	-29.4	35.6			
19	77.5	D2			-29.4				
20	79.5	D2	6.2	-4.1	-27	33.2			
21	80.5	D2			-28.8				
22	81.5	D2	4.8	-5.0	-27.9	32.7			
23	84.5	D2	6.1	-1.6	-28.3	34.4	0.1	0.33	0.02
24	85	D2	5.4	-4.4	-28.8	34.2			
25	86	D2		•	-28.6	01 E			
26	87.2	D2	5.2	-3.9	-26.5	31.7			
27	88	D2			-30.1				
28	88.7	D2	5.0	4.0	-27.2	22.2			
29	89	D2	5.2	-4.0	-27	32.2			
30	91.2	D2 D2	5.5	-3.3	-27.7	33.2			
31	91.7 02.5	D2 D2	6.1	-2.9	-27.6	33.7			
32	92.5	D2 D2			27.2				
33	93	D2 D2			-27.3				
34	96.2	D2 D2			-28.0				
30	98	D2 D2			-27.9				
30	99 00.85	D2 D2	Εć	26	-27.9	24.0			
3/ 30	77.83 101 5	D2 D2	5.0 5.2	-2.0 3.4	-29.2 28.6	34.8 33.0			
30 20	101.3	D2 D2	5.5	-3.4	-20.0	33.9			
39 40	101.85	D2 D2	5.8 6.2	-2.4 1 2					
40 /1	102	D2 D2	0.5	-1.5					
41 42	102.4	D2 D2							
43	102.7	D2	5.6	-1.5	-32	37.6	-1.23	0.35	0.1

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$\delta^{15}N^3$	N [wt. %]	TOC [wt.%]	C/N	∑REE [µg/g]	Ce/Ce*4	Y _N /Ho _N	Pr _N /Yb _N	EF Mn _(Cal-S)
1.8	0.003	0.01 0.11	2.7	16	0.95	1.5	0.97	48
2.6	0.006			27	0.83	1.4	1.1	30
2.3	0.005	0.0001	0.02					
0.5	0.011							
0.8	0.004	0.06	15	2.1	0.71	1.4	0.97	1
4.9	0.066	0.02	0.2	13	1.2	1.5	1.05	496
2.4	0.008	0.25	32	4.9	0.84	1.4	0.85	7.8
4.2	0.011	0.12	11	2.5	1.11	1.3	0.41	8.9
				2.2	0.66	1.7	0.57	2.1
5.1	0.014	0.21	15	1.7	0.69	1.8	0.49	2.9
5.5	0.028	0.12	4.4	0.33				0.71
3.9	0.017			3.8	0.9	2.1	0.77	6.4
4.9	0.092			0.33	0.69	1.5	0.4	0.14
6.6	0.13	0.61	4.7	2.9	1.01	1.6	0.58	1.7
5.4	0.034	0.22	6.6	2.9	1.15	1.3	0.62	5.2
4.8	0.023							
5.9	0.046	0.43	9.3	3.4	1.19	1.3	0.63	5.2
5.0	0.032	0.42	13	1.4	0.75	1.6	0.49	4.8
5.2	0.044	0.41	9.4	1.7	0.69	2.6	0.37	1.7
4.9	0.049	0.38	7.7	13	0.86	1.4	1.31	3.8
2.5	0.092	0.01	0.1	3.2	0.64	2.1	0.5	1.6
4.6	0.031	0.45	14	3.6	0.76	1.9	0.65	2.2
4.8	0.034	0.48	14	1.9	0.72	2.3	0.36	2.5
4.5	0.067	0.92	14	6.4	0.89	1.6	0.81	2.1
5.3	0.044	0.71	16	6.9	0.76	1.7	0.89	2.5
4.8	0.037	0.36	10	7.3	0.97	1.7	0.7	2.2
4.6	0.048	0.71	15					
4.4	0.044			6.7	0.8	1.5	0.91	1.1
2.3	0.017	0.29	17					
5.4	0.087	0.77	8.9	1.3	0.69	2.58	0.49	2.3
3.9	0.020							
1.9	0.011	0.26	23	2.4	0.72	2.3	0.57	1.4
3.2	0.014	0.35	25					
4.7	0.04	0.35	8.7	29	0.94	1.2	0.92	3.2
2.8	0.011	0.15	13	16	0.82	1.6	1.52	3.1
4.8	0.06	0.32	5.3	10	0.92	1.4	1.36	2.9
3.9	0.019	0.65	34	3.5	0.95	1.4	0.82	3.6
2.4	0.017	0.28	16	6	0.8	1.6	1.22	4
4.3	0.045			9.4	0.92			2.5
4.8	0.094	0.95	10			1.6	0.8	
4.5	0.097	0.4	4.1				0.63	
4.4	0.085	1	12	5.8	1.19	1.5	0.73	4.0

¹ O and C isotopic data relative to VPDB; ² Cd relative to NIST SRM 3108; ³ N relative to Air;

⁴ calculated following equations given by Lawrence and Kamber, 2006.



Letter

Geochemical Perspectives Letters

Letter

sample	Height [m]	Member	$\delta^{13}C_{carb}{}^1$	$\delta^{18}O_{carb}^{1}$	δ ¹³ Corg	$\Delta \delta^{13}$	$\begin{array}{c} \epsilon^{112/110} \\ Cd^2 \end{array}$	2SE	Cd [µg/g]
44	103	D2			-33.8				
45	103	D2	6.8	-1.1	-28.1	34.9			
46	103	D2	5.1	-2.1	-27.2	32.3			
47	103.5	D3			-31.2				
48	103.5	D3	5.2	-3.4	-31.2	36.4			
49	104	D3	3.5	-2.8	-29.2	32.7			
50	104	D3			-29.2				
51	104.25	D3			-27.8		-1.14	0.12	0.4
52	104.5	D3	4.2	-2.2	-27.3	31.5			
53	104.5	D3	5.0	-4.9	-28.3	33.3			
54	105.5	D3			-25.8				
55	108.15	D3			-31.9				
56	108.7	D3	4.0	-1.0	-27.3	31.3			
57	109.5	D3	2.0	-6.3	-27.8	29.8	-1.33	1.21	0.03
58	109.5	D3			-27.8				
59	110	D3	3.6	-6.8	-28.6	32.2			
60	110.5	D3			-28.4				
61	111.1	D3	2.9	-6.8	-29.6	32.5			
62	112.2	D3	2.8	-8.6					
63	113.9	D3	6.6	-5.4					
64	116	D3	6.9	-4.6		~~~~			
65	118.5	D3	6.1	-6.4	-27.2	33.3			
66	123.8	D3	6.1	-2.3	-28.6	34.7	1 (0	0.47	0.02
67	125	D3	5.4	-3.2	-31	36.4	-1.62	0.47	0.02
68	130	D3	5.7	-4.9	20.1		-0.13	0.31	0.08
69	133.5	D3	2 7		-30.1	22.0			
70	135	D3	3./ E (-7.5	-29.2	32.9	1.00	0.41	0.01
/1 72	140	D3	5.6	-0.7	-21.8	55.4	-1.22	0.41	0.01
72	142 1427	D4 D4	12	0.0	20.2	27.0	0.44	0.05	0.04
73	142.4 156.5	D4 D4	-1.5	-0.9	-27.2 -28.1	∠1.9 27.0	-0.44	0.95	0.04
75	150.5	D4 D4	-1.1	-0.4	-20.1	27.0	-1.51	0.59	0.01
76	156	D4 D4			-20.5				
77	167	D4	04		-20.1	28.2	-3 41	0.1	0.18
78	172.2	D4	3.3	-18	-27.0	20.2	-0.11	0.1	0.10
79	174	D4	47	-3.0	-271	31.8			
80	175.5	D4	49	-17	-28	32.9	-1.21	0.17	0.11
81	179.8	D4	5.2	-4.5	-27	32.2	1.21	0.17	0.11
82	180	D4	5.4	-5.3	-25.6	31.0	-0.79	0.12	0.07
83	194	D4	1.8	0.0	-27.9	29.7	0.7 2	0.12	0.07
84	203	DG	0.9	-2.8			-2.55	0.14	0.13
85	209	DG					-1.08	0.24	0.08
86	213	DG	3.3	-4.0			-2.19	0.10	0.07
87	217	DG	3.2	-3.4			-1.55	0.17	0.07

$\delta^{15}N^3$	N [wt. %]	TOC [wt.%]	C/N	∑REE [µg/g]	Ce/Ce ^{*4}	Y _N /Ho _N	Pr _N /Yb _N	EF Mn _(Cal-S)
4.6	0.051	0.21	4.1	8.8	1.4	1.3	0.83	2.9
5.3	0.137	0.97	7.1	44	1.16	1.5	0.75	2.1
5.4	0.126	0.89	7.1	11	0.98	1.6	0.7	2.4
3.1	0.015	0.59	39	3.9	0.98	1.3	1.07	1.2
3.1	0.015	0.59	39	3.5	1.09	1.4	0.84	3.5
5.0	0.029	0.11	3.9	14	1.33	1.4	1.32	
5.0	0.029	0.11	3.9	2.6	0.87	1.5	0.94	1.6
3.4	0.015	0.17	11	2.7	1.09	1.5	1	3.4
4.1	0.012	0.15	12	3.4	1.23	1.5	0.73	5.3
2.7	0.018	0.23	13	2.9	0.76	2.1	0.31	2.3
3.0	0.009	0.43	48	2.9	0.71	1.6	0.75	1.5
4.9	0.039	0.21	5.5	5.7	1.08	1.5	1.2	9.5
5.6	0.126	0.56	4.4	15	1.03	1.4	0.97	2.4
4.6	0.013	0.13	10	8.6	1.06	1.7	1.39	34
4.6	0.013	0.1	0 <i>(</i>	11	0.86	1.5	1.29	4.4
4.8	0.01	0.1	9.6	14	0.94	1.7	2.59	116
5.5	0.01	0.09	8.7	16	0.86	1.6	3.24	257
5.2	0.015	0.14	9.3	11	0.97	1.5	2.23	31
3.7	0.008	0.05	F 0	8.3	0.79	1.5	2.03	74
2.7	0.007	0.05	7.3	3.5	0.69	1.7	1.83	
3.6	0.010	0.07	0.0	3.0	0.7	1.8	1.23	1(0
3.6	0.009	0.07	8.0	6.7	0.87	1.7	1.37	162
3.3	0.012	0.17	14.5	10	0.97	1.3	1.67	10
5.3	0.018	0.24	13.1	10	1.02	1.4	1.98	19
4.1	0.009	0.17	20	12	1.01	1.5	2.97	60
4.2	0.008	0.16	20	9.0	0.87	1.4	2.21	32
4.4	0.007	0.31	44	10	0.99	1.8	1.17	10
4.0 5.5	0.007	0.15	18	11	1.01	1.5	2.3	10
5.5 E 0	0.017	1 6 1	24	12	0.00	1.1	1.04	2.1
5.9	0.040	0.12	54	1.9	1.2	2. 4 1.4	0.38	0.1
1.2	0.023	0.13	9.5 9.1	7.1	1.2	1.4	0.42	2.5
4.5	0.008	0.07	5.5	1.4	1.1	1.0 2.1	0.54	0.6
0.4	0.000	0.04	5.5	0.00	0.7	2.1	0.19	0.0
4.1	0.01	0.02	18	91	0.7	1.7	0.23	2.7
4.1 5.1	0.01	0.10	10	14.5	0.64	1.5	1.4	13
33	0.022	0.09	33	14.5	0.64	1.3	1.4	2.6
17	0.005	0.03	29	13	0.75	1.5	2.7	1.0
1.7	0.005	0.14	29	13	0.75	1.4	2	3.5
0.2	0.008	0.09	12	10	0.7	1.5	4	0.0
0.2	0.000	0.07	14	0.43	0.76	17	0.15	19
				0.56	0.86	1.7	0.15	57
				0.51	0.88	1.0	0.18	27
				0.01	0.00	1.0	0.10	2.7

Geochem. Persp. Let. (2017) 3, 32-44 | doi: 10.7185/geochemlet.1704



39



Figure 2 (a) Mn vs. $\varepsilon^{112/110}$ Cd. Cap dolomites (open triangles) have high Mn concentrations and exhibit positive $\varepsilon^{112/110}$ Cd. (b) N vs. Cd isotopic compositions reveal negative trend. (c) Carbon vs. Cd isotopic compositions show slight positive correlation when extreme values (red squares) and cap dolomites are excluded. (d) Cd concentrations vs. $\varepsilon^{112/110}$ Cd values scatter along modern Southern ocean fractionation line (Abouchami *et al.*, 2011), open circles represent coeval calculated $\varepsilon^{112/110}$ Cd_{sw}; note that the seawater Cd concentrations cannot be estimated from those measured in the carbonates.

As has been discussed above, there is still no consensus as to why cadmium is taken up by phytoplankton in the surface ocean, nor at which step (or steps) in this process the isotope fractionation takes place. Nevertheless, it is clear that $\epsilon^{112/110}$ Cd is a good indicator of biological productivity in the modern surface ocean: Cd uptake into OM leads to a depletion of light Cd in the photic zone. But can Cd isotopes in a marine sedimentary archive be used as a palaeo-productivity proxy? Under closed-system conditions (e.g., a restricted basin) Ediacaran carbonates precipitating in biologically productive environments would be expected to have higher (heavier) $\epsilon^{112/110}$ Cd and $\delta^{13}C_{\rm carb}$. The Xiaofenghe carbonates do exhibit a positive correlation between $\epsilon^{112/110}$ Cd and $\delta^{13}C_{\rm carb}$ (Fig. 2c), consistent with increasing bio-productivity. However, this correlation only exists once light $\delta^{13}C_{\rm carb}$ values from the basal Doushantuo (oxidation of a ^{12}C -enriched hydrocarbon source) and one sample with extremely heavy $\delta^{13}C_{\rm carb}$ of +9 (presumably due to evaporation) are excluded.

The $\epsilon^{112/110}$ Cd and δ^{15} N curves in Figure 1 show an anti-correlation in some parts of the profile - for example, higher $\epsilon^{112/110}$ Cd in the cap dolomites with δ^{15} N having low but positive values (Fig. 2b). This trend would be consistent with

40



Letter

strong N₂ fixation by diazotrophs in the euphotic zone (Sachs and Repeta, 1999). Cyanobacteria blooms of this sort are often documented from redox-stratified basins (Struck *et al.*, 2004), and recent work by Georgiev *et al.* (2015) has highlighted the redox control on Cd distribution in late Permian marine sediments. Cd is a not a redox-sensitive element *per se*, but Cd bound to OM will become enriched in anoxic sediments whenever oxidation of OM is inhibited.

Cadmium may also be co-precipitated with sulphides under euxinic conditions (Framson and Leckie, 1978), such as prevail in the deeper parts of the Black Sea today, where Cd is removed almost quantitatively from the water column and enriched in bottom sediments (Lewis and Landing, 1992). The effect of authigenic sulphides on Cd isotope fractionation is still not well understood. Cadmium isotope data on hydrothermal sulphide ores (Schmitt *et al.*, 2009; Wen *et al.*, 2016) and quantum chemical calculations (Yang *et al.*, 2015) suggest that the light isotopes of Cd would be preferentially bound into the sulphide phase. On the other hand, Rolison *et al.* (2015) did not find the high $\varepsilon^{112/110}$ Cd in the deeper parts of the Black Sea water column that one would expect. All in all this means that sulphide-bearing anoxic sediments should most likely have higher Cd concentrations and more negative $\varepsilon^{112/110}$ Cd than those of oxic surface waters.

In the Xiaofenghe carbonates, Mn enrichments correlate with high $\epsilon^{112/110}$ Cd (Fig. 2a). As Mn in oxic waters is usually incorporated into Mn oxyhydroxides, such enrichments are likely to reflect reduction of Mn to the 2+state at the anoxic pore-water sediment interface and subsequent incorporation into CaCO₃ (Thomson *et al.*, 1986). The cap dolomites exhibit Mn enrichments, more positive $\epsilon^{112/110}$ Cd and low Cd concentrations. Furthermore, sedimentary sulphides ranging from 10 to 40 µm in size are abundant in thin section (see Fig. S-2). The fact that Ce anomalies are negative (Ce/Ce* < 0.9) in shallow Yangtze Platform cap dolomites (Fig. 1h) suggests that early Ediacaran surface-waters were already oxic; nonetheless, pore-waters may still have been anoxic, leading to the formation of sulphides may raise $\epsilon^{112/110}$ Cd in associated carbonates, and might have affected the Xiaofenghe cap dolomites.

Experiments on precipitation of inorganic calcite from seawater by Horner *et al.* (2011) show that the seawater/calcite fractionation factor for Cd is insensitive to temperature, Mg concentration and precipitation rate. By contrast, precipitation of calcites from freshwater does not lead to any Cd isotopic fractionation, presumably as a result of ion blocking on mineral surface sites. Since these experiments by Horner *et al.* (2011) used only seawater and freshwater end-members, we will assume to first order that Cd isotope fractionation into calcite depends linearly on salinity. Using Y/Ho ratios as an indicator of palaeo-salinity we are then able to calculate $\varepsilon^{112/110}$ Cd for seawater in equilibrium with the Xiaofenghe carbonates (see Fig. S-3 for more details). The resulting $\varepsilon^{112/110}$ Cd_{sw} range between -2 and +1.5 (Fig. 1e), which is still slightly lower than the lower bound of modern surface seawater $\varepsilon^{112/110}$ Cd but scatter around the Rayleigh fractionation line defined by modern Southern Ocean seawaters (Abouchami *et al.*, 2011; Fig. 2d). If the Cd



isotopic variations observed are controlled by changes in seawater salinity at the Xiaofenghe depositional site, estuarine mixing of seawater and river water, as described in Hohl *et al.* (2015), may account for the observed variation. The correlation between Y/Ho and $\varepsilon^{112/110}$ Cd (see Fig. 1g and Fig. S-1c) may therefore be due to an increased riverine input, flattening the REE patterns, decreasing the total salinity and thus reducing the isotopic fractionation into calcite. Any further increase in riverine input would most likely drown out any potential biological signal in the stable isotope proxies, such as $\varepsilon^{112/110}$ Cd, which would then resemble typical crustal values.

Implications

Stable Cd isotopic compositions of Ediacaran-age carbonates from the Xiaofenghe section on the Yangtze Platform show significant variations of up to four $\epsilon^{112/110}$ Cd units. There are several possible reasons for these variations: part of the signal may be biological in origin while abiological (inorganic) processes almost certainly play a role as well. Factors controlling the Cd isotopic compositions may include fluid flow alteration, the precipitation of authigenic sulphides under anoxic pore water conditions, both processes possibly modifying $\epsilon^{112/110}$ Cd in the cap dolomites; additionally there are changes in Cd isotope fractionation into calcite as a function of salinity, as suggested here for the upper Doushantuo. Encouragingly, our salinity-corrected palaeo-seawater $\epsilon^{112/110}$ Cd_{sw} signals overlap those of modern surface waters. A positive correlation between $\delta^{13}C_{carb}$ and $\epsilon^{112/110}$ Cd is observed, which suggests that the Cd isotope fractionation might, in part, have a biological origin, meaning that phytoplankton were present as early as the Ediacaran. Future research on marine sediments will need to address the abiotic fractionation processes above, and correct for them, in order to arrive at any biologically-derived Cd isotope signal present.

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Additional Information

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43

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