

Tracking the spatial extent of redox variability in the mid-Proterozoic ocean

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23 ABSTRACT

24 Emerging geochemical evidence suggests considerable redox heterogeneity in the mid-
25 Proterozoic ocean. However, quantitative estimates of the extent of different modes of anoxia remain
26 poorly constrained. Due to their complementary redox-related behaviour, uranium and molybdenum
27 isotopes can be combined to reconstruct ancient marine redox landscapes, but this approach has not
28 been applied to the mid-Proterozoic. Here, we present new $\delta^{238}\text{U}$ and $\delta^{98}\text{Mo}$ data for marine rocks
29 from the ~ 1.4 Ga Xiamaling Formation, North China Craton, together with independent redox
30 indicators (Fe speciation and redox-sensitive trace metals). We find that most samples deposited
31 under oxic or dysoxic conditions retain low U and Mo contents, with $\delta^{238}\text{U}$ and $\delta^{98}\text{Mo}$ values
32 indistinguishable from continental crust, demonstrating a dominant detrital signal. By contrast,
33 euxinic samples with authigenic enrichments in U and Mo record the highest authigenic $\delta^{238}\text{U}$ and
34 $\delta^{98}\text{Mo}$ values, consistent with efficient reduction of U and Mo. Samples deposited under ferruginous
35 conditions exhibit a wider range of intermediate $\delta^{238}\text{U}$ and $\delta^{98}\text{Mo}$ values that generally fall between
36 the (dys)oxic and euxinic end-members. Using a coupled U-Mo isotope mass balance model, we infer
37 limited euxinia ($<0.5\%$ of the global seafloor area) but extensive low-productivity (dys)oxic and
38 ferruginous settings in ~ 1.4 Ga oceans. This redox landscape would have provided potentially
39 habitable environments for eukaryotic evolution in the mid-Proterozoic.

40

41 INTRODUCTION

42 Reconstructing the oxygenation history of Earth's surface environment is crucial to understand
43 the trajectory of Earth's habitability. The mid-Proterozoic (1.8–0.8 Ga) was a critical interval for
44 early eukaryote evolution (Knoll and Nowak, 2017), and while emerging evidence suggests that

45 biological innovation at this time occurred under heterogeneous ocean redox conditions (Sperling et
46 al., 2014; Zhang et al., 2016; Luo et al., 2021; Song et al., 2023), the global extent of different modes
47 of anoxia remains poorly constrained. As such, a quantitative assessment of the global redox
48 landscape may provide critical insight into the spatial extent of habitable conditions, thus ultimately
49 enabling improved consideration of potential controls on early eukaryote evolution.

50 Uranium and molybdenum isotopes ($\delta^{238}\text{U}$ and $\delta^{98}\text{Mo}$) are useful tools for reconstructing global
51 redox conditions because of their redox sensitivity and long oceanic residence times (Andersen et al.,
52 2017; Kendall et al., 2017). The largest U isotope fractionations occur during U reduction in anoxic
53 environments, with heavy U isotopes preferentially sequestered in sediments (Weyer et al., 2008).
54 The largest Mo isotope fractionations occur during adsorption to Mn-Fe (oxyhydr)oxides under oxic
55 conditions (Kendall et al., 2017). By contrast, minimal isotopic difference is expected between Mo
56 in sediments and coeval seawater during near-quantitative drawdown under strongly euxinic
57 conditions (Nägler et al., 2011). Since rapid Mo burial specifically requires relatively high dissolved
58 sulfide concentrations, but U burial only requires anoxia, a particularly robust reconstruction of the
59 extent of different ocean redox conditions can be achieved when considering $\delta^{238}\text{U}$ and $\delta^{98}\text{Mo}$
60 together (Andersen et al., 2020; Kendall et al., 2020). Although there are several studies using either
61 $\delta^{238}\text{U}$ or $\delta^{98}\text{Mo}$ to investigate ocean redox variability in the mid-Proterozoic (Arnold et al., 2004;
62 Yang et al., 2017; Gilleaudeau et al., 2019; Luo et al., 2021), there have been no studies of U-Mo
63 isotope co-variation during this period.

64 Here, we present new U and Mo isotope data for drill core samples from the ~1.4 Ga Xiamaling
65 Formation, North China Craton. In combination with new Re concentration data, which enables
66 specific identification of dysoxic conditions (Crusius et al., 1996; Song et al., 2023; Li et al., 2025),

67 and existing Fe speciation, $\delta^{98}\text{Mo}$ and U-Mo concentration data, we first explore $\delta^{238}\text{U}$ and $\delta^{98}\text{Mo}$
68 systematics in the context of the local redox state. We then utilize an isotope mass balance model to
69 reconstruct the spatial extent of different redox conditions in the ~ 1.4 Ga ocean.

70

71 **GEOLOGIC SETTING**

72 The ~ 1.4 Ga Xiamaling Formation (Zhang et al., 2015) study site is located in the Xiahuayuan
73 region of Hebei Province, north China (Fig. 1A), where the formation can be divided into six units
74 (Wang et al., 2017; see Supplemental Material for detailed geologic background). Here, we focus on
75 fresh drill core material from the upper four units (Fig. 1B), which are dominantly composed of low
76 thermal maturity mudstones and black shales, representing deep-water deposition (Wang et al., 2017).

77

78 **RESULTS**

79 In total, 50 samples were analyzed for $\delta^{238}\text{U}$ compositions, while an additional 15 samples were
80 analysed to augment existing $\delta^{98}\text{Mo}$ data (Zhang et al., 2019; see Supplemental Material for methods
81 and data). Bulk $\delta^{238}\text{U}$ values show relatively constant values through unit 4 ($-0.22 \pm 0.04\text{‰}$, 2SD;
82 Fig. 2). An excursion to lower values (as low as -0.41‰) and then to higher values (up to 0.08‰)
83 occurs in unit 3, followed by a progressive increase to values approaching 0.2‰ at the top of unit 2
84 (Fig. 2). Samples in unit 1 show more scatter, but with an initial drop to lower values, followed by a
85 general increase up-section (Fig. 2). Authigenic $\delta^{238}\text{U}$ ($\delta^{238}\text{U}_{\text{auth}}$) compositions (see Supplemental
86 Material for detrital corrections; note that samples with low U and Mo contents were excluded from
87 authigenic correction) range from -0.44‰ to $+0.26\text{‰}$, and exhibit a similar trend to bulk $\delta^{238}\text{U}$ (Fig.
88 3). Our $\delta^{98}\text{Mo}$ data range from 0.15‰ to 1.26‰ , with negligible difference relative to authigenic

89 $\delta^{98}\text{Mo}$ ($\delta^{98}\text{Mo}_{\text{auth}}$) (Fig. 2; Table S1). In almost all cases, Re enrichment factors (Re_{EF} ; see
90 Supplemental Material for the enrichment factor (EF) calculation) are above 1 (Fig. 2).

91

92 **DISCUSSION**

93 **Local Redox conditions**

94 Previous detailed reconstructions have invoked a generally anoxic, but dynamic redox setting
95 for the Xiamaling Formation (Zhang et al., 2016; Wang et al., 2017; Song et al., 2023). Based on
96 mostly low but variable highly reactive Fe to total Fe ($\text{Fe}_{\text{HR}}/\text{Fe}_{\text{T}}$) ratios and $\text{U}_{\text{EF}}\text{-Mo}_{\text{EF}}$ values, coupled
97 with generally low pyritization (Fe_{py}) of the Fe_{HR} pool (Fig. 2), units 3 and 4 have been interpreted to
98 record orbital-scale variability in the spatial extent of a ferruginous oxygen minimum zone (OMZ),
99 which became more productive through unit 3 (Wang et al., 2017; Song et al., 2023). More expansive
100 deeper water anoxia developed in unit 2, with a progression from ferruginous to euxinic conditions,
101 while unit 1 documents continued euxinia, punctuated by more oxygenated conditions in the upper
102 half of the unit (Fig. 2; Wang et al., 2017).

103 Persistent enrichments in Re (Fig. 2), including for samples with low $\text{Fe}_{\text{HR}}/\text{Fe}_{\text{T}}$ ratios, further
104 suggest that deeper waters below the OMZ in units 3 and 4 were likely dysoxic (Crusius et al., 1996;
105 Song et al., 2023; Li et al., 2025), although intervals of fully oxygenated conditions (Zhang et al.,
106 2016; Wang et al., 2017) cannot be discounted (hence we refer to such samples as being (dys)oxic).

107 **Controls on U and Mo Isotope Compositions**

108 To explore $\delta^{98}\text{Mo}$ - $\delta^{238}\text{U}$ variability, we plot $\delta^{98}\text{Mo}$ and $\delta^{238}\text{U}$ profiles in the context of local
109 redox conditions (Fig. 2). The (dys)oxic samples have $\delta^{238}\text{U}$ values that are essentially
110 indistinguishable from average continental crust (-0.3‰), consistent with generally low U contents

111 and a dominant detrital contribution (Fig. 2; Fig. S1) (Andersen et al., 2017). Ferruginous samples
112 have $\delta^{238}\text{U}_{\text{auth}}$ values that span a relatively wide range (Fig. 2), consistent with a previous $\delta^{238}\text{U}$ study
113 of ferruginous lakes (Cole et al., 2020). In more detail, ferruginous samples with low TOC and U_{EF}
114 values (termed F2 samples) tend to have $\delta^{238}\text{U}$ similar to the detrital composition, whereas higher
115 $\delta^{238}\text{U}_{\text{auth}}$ values are observed for samples with higher TOC and U_{EF} (F1 samples) (Fig. 2). Euxinic
116 samples are generally enriched in authigenic U and record the highest $\delta^{238}\text{U}_{\text{auth}}$ values (Fig. 2; see
117 Supplemental Material for discussion of two euxinic samples with anomalously low $\delta^{238}\text{U}_{\text{auth}}$).

118 Integrating our new $\delta^{98}\text{Mo}$ analyses with previously published data (Zhang et al., 2019) shows
119 that (dys)oxic and F2 samples are dominantly characterized by low $\delta^{98}\text{Mo}$ values ($0.34 \pm 0.14\text{‰}$),
120 close to continental crust ($\sim 0.3\text{‰}$) (Voegelin et al., 2014), while the highest $\delta^{98}\text{Mo}$ values are
121 observed for euxinic and F1 sediments (Fig. 2). It is noteworthy that euxinic samples have $\delta^{98}\text{Mo}$
122 values spanning a wide range, possibly suggesting variable, and relatively low, water column sulfide
123 levels (Neubert et al, 2008). In addition, sulfidic pore waters may also have exerted an important
124 control on the observed $\delta^{98}\text{Mo}$ variability (Kendall et al., 2017).

125 While $\delta^{238}\text{U}$ systematics are commonly used to evaluate global ocean redox changes, the
126 potential impact of local redox conditions and organic carbon loading on fractionations recorded in
127 the sediments needs to be considered (Lau et al., 2022; Rutledge et al., 2024). Although bulk $\delta^{238}\text{U}$
128 values show a general positive correlation with TOC (when OMZ samples are excluded) and U
129 contents, $\delta^{238}\text{U}_{\text{auth}}$ values display no such correlation (Fig. 3), suggesting a negligible local
130 environmental control on $\delta^{238}\text{U}_{\text{auth}}$ fractionations. We also note that our $\delta^{238}\text{U}_{\text{auth}}$ values are generally
131 comparable to shale $\delta^{238}\text{U}_{\text{auth}}$ data (-0.27‰ to 0.16‰) from the ~ 1.36 Ga Velkerri Formation,

132 northern Australia (Yang et al., 2017), suggesting a consistent isotopic offset from global seawater
133 and relatively stable seawater $\delta^{238}\text{U}$.

134 Neither $\delta^{98}\text{Mo}$ or $\delta^{98}\text{Mo}_{\text{auth}}$ compositions show a systematic covariation with TOC or Mo
135 contents (Fig. 3), therefore local changes in productivity or sedimentation rate appear to have exerted
136 minimal impact on the observed $\delta^{98}\text{Mo}$ variability. Diamond et al. (2018) and Zhang et al. (2019)
137 reported two distinct $\delta^{98}\text{Mo}$ maxima for euxinic sediments of the Xiamaling Formation at two
138 different localities, which likely suggests heterogenous sulfide availability in the relatively low-
139 sulfate mid-Proterozoic ocean (Fakhraee et al., 2019). Nevertheless, a combined evaluation of $\delta^{238}\text{U}$ -
140 $\delta^{98}\text{Mo}$ can be used to estimate global ocean redox variability (Andersen et al. 2020; Kendall et al.,
141 2020).

142

143 **Reconstructing Global Redox Conditions**

144 Generally, fluctuating $\delta^{238}\text{U}$ and $\delta^{98}\text{Mo}$ values in the Xiamaling Formation are consistent with
145 local redox dynamics, and their ranges across different redox conditions are relatively constant
146 throughout the formation (Fig. 2), suggesting relatively stable oceanic U-Mo inventories and isotopic
147 compositions. To quantitatively evaluate the global oceanic redox distribution at this time, we employ
148 a stochastic isotope mass balance model, assuming that both U and Mo are dominantly sourced from
149 rivers and are buried under (dys)oxic, ferruginous and euxinic conditions. In this model, U-Mo burial
150 is a function of the areal proportion of each redox sink, and seawater $\delta^{238}\text{U}$ and $\delta^{98}\text{Mo}$ are calculated
151 from U-Mo burial and corresponding isotope fractionations for each redox sink, utilising specific
152 fractionation factors from previous studies (Table S2; see Supplemental Material for detailed
153 modelling approach). By running the model 10,000 times from a modern-day initialization through a

154 random selection of ferruginous and euxinic areal fractions, model outputs (i.e., U and Mo isotope
155 compositions for each redox sink) are produced that fit both the $\delta^{238}\text{U}$ and $\delta^{98}\text{Mo}$ data for the
156 Xiamaling Formation (Fig. 4). An advantage of our model is that specific isotope compositions for
157 ~ 1.4 Ga seawater are not required, as the model calculates coeval seawater isotope compositions in
158 association with randomly selected isotope compositions and fractionation factors for each sink.

159 The resultant area that intersects all coloured areas (area b in Fig. 4) suggests that, at most, $\sim 0.5\%$
160 of the global seafloor was overlain by euxinic waters at ~ 1.4 Ga, while the areal fraction of
161 ferruginous conditions was at least 20%, leaving the rest of the seafloor in a (dys)oxic state. Our
162 estimate of more expansive ferruginous conditions is consistent with ferruginous-dominated redox
163 models proposed for the mid-Proterozoic (Poulton et al., 2010; Planavsky et al., 2011; Poulton and
164 Canfield, 2011). By contrast, our estimate for the extent of euxinia at ~ 1.4 Ga is relatively small
165 compared to estimates for other intervals of the mid-Proterozoic ($< 10\%$), which were constrained by
166 either $\delta^{238}\text{U}$ or $\delta^{98}\text{Mo}$ (Gilleaudeau et al., 2019; Luo et al., 2021). This may reflect either temporal
167 variability in water column euxinia, or model limitations when considering one isotope system in
168 isolation.

169 Recent studies have suggested increased ocean oxygenation at ~ 1.4 Ga, relating to enhanced
170 nutrient-driven primary productivity (Cox et al., 2016). However, our results suggest that large
171 expanses of the ocean were more likely dysoxic than oxic, and as such, elevated productivity was
172 likely restricted to local settings experiencing increased nutrient availability, driven either by locally-
173 enhanced nutrient influxes or recycling under euxinic conditions (Song et al., 2023). The limited
174 extent of euxinia in ~ 1.4 Ga oceans would have diminished the specific ‘toxicity’ control of sulfide
175 on eukaryote evolution (Anbar and Knoll, 2002), which together with the expansive extent of at least

176 mildly oxygenated oceans (Heard et al., 2023), may have exerted a significant control on the evolution
177 of the biosphere in the mid-Proterozoic.

178

179 CONCLUSIONS

180 Uranium and Mo isotope systematics in the ~1.4 Ga Xiamaling Formation provide constraints
181 on global ocean redox evolution in the mid-Proterozoic. Our $\delta^{238}\text{U}$ and $\delta^{98}\text{Mo}$ data exhibit distinct
182 ranges related to different redox conditions, with (dys)oxic samples having $\delta^{238}\text{U}$ - $\delta^{98}\text{Mo}$ values
183 indistinguishable from the detrital input. By contrast, the highest $\delta^{238}\text{U}_{\text{auth}}$ - $\delta^{98}\text{Mo}_{\text{auth}}$ values are
184 preferentially recorded in euxinic samples. Ferruginous sediments with low TOC tend to have low
185 $\delta^{238}\text{U}$ - $\delta^{98}\text{Mo}$ values, while higher-TOC ferruginous samples tend to have intermediate to high $\delta^{238}\text{U}$ -
186 $\delta^{98}\text{Mo}$. Coupled U-Mo modelling results suggest that eutrophic euxinic settings covered ~0.5% of the
187 global seafloor at most, while less-productive ferruginous and (dys)oxic settings were much more
188 extensive at ~1.4 Ga. This redox partitioning may account for the moderate diversification of
189 eukaryotes observed in the Xiamaling Formation. Similar studies linking mid-Proterozoic isotopic
190 and fossil records will help to determine whether the Xiamaling Formation documents a relatively
191 static mid-Proterozoic redox environment, or one stage in a temporal progression towards increased,
192 stable ecological niche space that promoted enhanced eukaryotic diversification.

193

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321 FIGURE CAPTIONS

322 Figure 1. A: Geological map of the North China Craton, after Zhao et al. (2005) and Wang et al.
323 (2017). Red square shows the location of the study area. B: Simplified stratigraphy of the Xiamaling
324 Formation, with age constraints from Zhang et al. (2015).

325

326 Figure 2. Stratigraphic geochemical profiles for the Xiamaling Formation. Total organic carbon
327 (TOC), Fe speciation, U and Mo concentration data are from Wang et al. (2017). Re data are from
328 this study (closed circles) and Song et al. (2023) (open circles). The bulk $\delta^{98}\text{Mo}$ profile comprises
329 data from Diamond et al. (2018) (red open circles), Zhang et al. (2019) (dark open circles for core
330 samples and grey open circles for outcrop rocks), and this study (closed circles). All $\delta^{238}\text{U}$ data are
331 from this study. Blue shading on the $\delta^{98}\text{Mo}_{\text{auth}}$ and $\delta^{238}\text{U}_{\text{auth}}$ profiles represent isotope ranges for
332 (dys)oxic and F2 samples, while grey shading indicates the euxinic isotope range. EF–enrichment
333 factor. (U)CC–(upper) continental crust. F1–high-TOC ferruginous. F2–low-TOC ferruginous.

334

335 Figure 3. Cross-plots of $\delta^{238}\text{U}$ and $\delta^{98}\text{Mo}$ *versus* their respective elemental and TOC contents. $\delta^{238}\text{U}_{\text{det}}$
336 and $\delta^{98}\text{Mo}_{\text{det}}$ represent average continental crust values of -0.3‰ (Andersen et al., 2017) and 0.3‰
337 (Voegelin et al., 2014), respectively. Note that the regression line on the $\delta^{238}\text{U}$ *versus* TOC plot
338 excludes the OMZ data (an R^2 of 0.12 is obtained when the OMZ data are included).

339

340 Figure 4. Simplified U-Mo isotope mass balance model outputs. Most likely isotope compositions
341 are mapped as a function of the relative areal fraction in different redox sinks. Area (a) represents the
342 solution space that satisfies the $\delta^{238}\text{U}$ data for the three sinks, while area (b) represents the solution

343 space that additionally satisfies the $\delta^{98}\text{Mo}$ data. Thus, area (b) represents the inferred spatial extent of
344 ferruginous and euxinic conditions, as satisfied by both isotope systems.

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347 ¹Supplemental Material. Detailed descriptions of the geological background, analytical methods,
348 detrital corrections, further discussion, modelling approach and geochemical data. Please visit
349 <https://doi.org/10.1130/XXXX> to access the supplemental material, and contact
350 editing@geosociety.org with any questions.

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