Supplemental Material

Mercury isotopes track the cause of carbon perturbations in the Ediacaran ocean

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Note that the data sources are in the Supplemental Table Excel files.
Text S1: Geological setting of sections studied

The Doushantuo Formation at the Jiulongwan section

At the Jiulongwan section (30°48'54" N, 111°03'20" E) in the Nanhua Basin, the Doushantuo Formation was deposited below or near wave base, and it is commonly divided into four lithostratigraphic members with several transgressive-regressive cycles (Jiang et al., 2007; McFadden et al., 2008). Member I, which overlies Nantuo Formation glacial sediments, is an approximately 5 m thick cap dolostone that is associated with a transgression. Member II contains 70 m of organic-rich shale and dolostone beds that contain abundant pea-sized chert nodules. Member III is mainly composed of limestone, but it sometimes contains cherty bands in the lower part of the member. The upper part of members II and the whole member III mostly correspond with two regressive episodes. Member IV is mostly organic-rich shale that is widespread across the Yangtze Gorges area in the Nanhua Basin. The sharp lithostratigraphic contact between members III and IV represents a transgressive surface. The first negative carbon isotope excursion (NCIE-1) is preserved in member I and the lower part of member II (below 16 m). NCIE-2 was observed at the top of member II (mostly between 70 m and 80 m, (Jiang et al., 2007; McFadden et al., 2008)), which is not always recognized in other Ediacaran sections in South China. We did not observe NCIE-2 in this study due to the low sampling resolution. NCIE-3 occurs in the middle of member III and continues through to member IV (from ~ 100 m to the end of the formation), which is temporally correlated with the Shuram excursion from Oman (Jiang et al., 2007; McFadden et al., 2008). Two positive carbon isotope excursions, named PCIE-1 (16.1-73 m) and PCIE-2 (79-96 m), occur between these three NCIEs, as has been show in previous studies (Jiang et al., 2007; McFadden et al., 2008).

The Wonoka Formation in Adelaide Rift Complex

The Wonoka Formation of the Adelaide Rift Complex, South Australia, is a ~500–1500 m thick Ediacaran carbonate and siliciclastic succession (Calver, 2000; Haines, 1990; Husson et al., 2015). The Wonoka Formation overlies the siltstones of the Bunyeroo Formation and coarsens and shallows upward into the Pound Subgroup (e.g. Husson et al., 2015). Haines (1990) divided the Wonoka Formation into eleven mappable units. Units 1 (interbedded dolostone and shale) and 2 (sandstone turbidite beds and interlayered shale) were mostly deposited below storm wave base in a deep outer shelf setting. Up-section, units 3 to 7 are comprised of a shallowing-upwards shelf succession that, essentially, transitions from shale-dominated to limestone-dominated, indicating shallowing from an outer to middle shelf environment. Unit 8, which contains shale with minor limestone storm beds, indicates
a short-lived deepening of the paleoenvironment. Units 9 to 11 are peritidal facies and were deposited in a lagoonal setting. An obvious NCIE starts at the lowest part of this formation, with $\delta^{13}$C$_{\text{carb}}$ decreasing from -3.9‰ to -12‰, before increasing to +5‰ up-section. This NCIE, named the Wonoka excursion, is equivalent to the Shuram excursion (Calver, 2000; Husson et al., 2015). Detailed descriptions of the Wonoka Formation may be found in Haines (1990) and Husson et al. (2015).

**Text S2: Detailed Methodology**

**Processing of sediment samples:** After removing possible weathering surfaces, samples were ground and sieved using a 200 μm mesh size. After processing, these samples were stored in screw-capped polyethylene bottles.

**Total Hg concentrations:** Total Hg concentrations were determined using a DMA-80 automatic Hg analyzer (Milestone, Italy) at the State Key Laboratory of Environmental Geochemistry, Chinese Academy of Sciences (CAS). The DMA-80 analyzer was routinely calibrated, and data accuracy and precision were assessed by analyzing procedural blanks, certified standard reference materials (NIST SRM 2711 and BCR 482) and sample duplicates. Duplicate analysis of Hg concentrations, overall, shows a small relative standard deviation (mean = ±5.5%). Total Hg concentrations reported in this study are the average value of the duplicate tests.

**Isotopic composition of Hg:** Hg in sedimentary samples was preconcentrated into 10 mL of 40% mixed acid solution (v/v, HNO$_3$/HCl = 2:1) using a double-stage combustion method developed for Hg isotope analysis (Sun et al., 2013). Immediately after the preconcentration, the trapping bottle and impinger were rinsed three times with 10 mL of Milli-Q water, which was added into the trapping solution to yield an ultimate acid concentration of ~20%. The final trapping solutions were kept in a refrigerator at 2–4 °C until the Hg isotope analysis was performed. Hg isotope ratios were analyzed using cold vapor-multicollector inductively coupled plasma mass spectrometry (CV-MC-ICPMS, Nu Instruments, U.K.) at the State Key Laboratory of Environmental Geochemistry, CAS (Guiyang, China) (Fu et al., 2019). Isotopic ratios were corrected for mass bias by standard-sample-standard bracketing using the National Institute of Standards and Technology (NIST) Standard Reference Material (SRM) 3133. Hg isotopic compositions are reported as delta values ($\delta$) in permil (‰) relative to the mean ratios measured for the NIST SRM 3133 before and after each sample using equation (1) (Blum and Bergquist, 2007):
where $\Delta^{\text{x}}$ refers to the mass of each Hg isotope between 199 and 204. MIF is reported in capital delta values ($\Delta$), which is the difference between the measured isotopic compositions ($\delta^{\text{x}}Hg$) and the theoretically predicted values based on the kinetic MDF law:

$$
\Delta^{\text{x}}Hg (\%) = \left[ \frac{\delta^{\text{x}}Hg_{\text{sample}}}{\delta^{\text{199}}Hg_{\text{NIST SRM 3133}}} - 1 \right] \times 1,000 \quad (1)
$$

$\delta^{\text{x}}$ refers to the mass of Hg isotope $^{199}$Hg, $^{200}$Hg, $^{201}$Hg, and $^{204}$Hg. The corresponding $\beta$ values of these Hg isotopes are 0.252, 0.502, 0.752, and 1.493, respectively. The analytical uncertainties of Hg isotopic compositions were evaluated by repeated analysis of the isotopic compositions of NIST SRM 3177 ($n = 39$), Lichen CRM BCR 482 ($n = 10$) and NIST SRM 2711 (Montana soil, $n = 5$). The measured Hg isotopic compositions of NIST SRM 3177, Lichen CRM BCR 482, and NIST SRM 2711 are presented in Table S1 and agree with previously published results (Biswas et al., 2008; Estrade et al., 2010; Yin et al., 2016). The analytical uncertainty of Hg isotopic composition in this study is the larger 2σ value of either the NIST SRM 3177 or repeated analysis of the sediment sample over different analytical sessions.

**Carbon isotope of carbonate:** Carbon isotope data from all samples of the Doushantuo Formation were measured at the Third Institute of Oceanography, State Oceanic Administration, China. Briefly, approximately 10-30 mg of powdered sample was extracted by concentrated phosphoric acid in a vacuum vial at 70 °C for 200-300 s using a Kiel IV device. The extracted CO2 was directly introduced into a GasBench II isotope ratio mass spectrometer (IRMS). The isotopic composition of carbon in carbonate ($\delta^{13}C_{\text{Carb}}$) was defined as the standard per mil notation (‰) relative to V-PDB (Vienna Peedee Belemnite). The long analysis of NBS-19 and Chinese national standard GBW04416 indicates an error better than 0.2‰ (2σ). Carbon isotopic compositions of carbonate in samples from the Wonoka Formation were taken from Ward et al. (2019).

**Content of total organic carbon (TOC), total sulfur (TS), major and trace elements:** To measure TOC content, 2N HCl was firstly used to leach the carbonate fraction. After centrifuging, the residual samples were washed three times using 18 Ω purified water. After washing, the residual samples were analyzed for total carbon, which represents TOC fraction. TOC and TS concentrations in sediment samples were
measured using a LECO C-S element analytical instrument at Guangdong Province Research Center for Geoanalysis. Major elements were analyzed using X-ray fluorescence (XRF) at the ALS Chemex facility in Guangzhou, China. Trace elements were measured using ICP-MS at the State Key Laboratory of Ore Deposit Geochemistry, IGCAS (Liang et al., 2000). The analytical uncertainty for trace elements was better than ±5 % based on the long-term results of two standard reference materials (carbonate, COQ-1; shales, SBC-1).

**Text S3: Hg isotope of the Ediacaran sediment background in the Nanhua basin**

In the Doushantuo Formation, most chert and dolostone samples contain very minor detrital contamination (Al₂O₃ <0.7%) and lower TOC (mostly <1%) and TS (mostly <0.1%). These samples exhibit relatively low total Hg concentrations (as low as 6.9 ppb) and homogenous $\delta^{202}\text{Hg}$ values (-1.40±0.17‰, N=12) and $\Delta^{199}\text{Hg}$ values (0.15±0.06‰, N=12) (**Table S4**). These values are very close to the average Hg isotope composition of (1) limestones ($\delta^{202}\text{Hg}$= -1.43±0.10‰, and $\Delta^{199}\text{Hg}$ = -0.07±0.04‰) such as those reported by Wang et al. (2015) and Sun et al. (2016), (2) early Cambrian phosphorites ($\delta^{202}\text{Hg}$= -1.39±0.26‰, $\Delta^{199}\text{Hg}$ = 0.18±0.05‰) with low Al₂O₃, TOC and TS content (Yin et al., 2017) and (3) pre-anthropogenic marine sediments that were dominated by background atmospheric Hg deposition ($\delta^{202}\text{Hg}$ =-1.42‰ and $\Delta^{199}\text{Hg}$ =0.12‰, estimated by Zheng et al. (2018)). However, these $\delta^{202}\text{Hg}$ values are lower than those of pre-anthropogenic Mediterranean background (-0.76±0.16‰) and sapropel samples (-0.91±0.15‰) (Gehrke et al., 2009), as well as sediments from the central Portuguese Margin at 2084 m water depth (-0.59±0.04 ‰) (Mil-Homens et al., 2013). These differences could be the result of there being a larger amount of detrital contents (Al₂O₃ > 9%) in those samples than in ours. Under marine conditions, organic matter (OM) drawdown plays an important role in the sequestration of dissolved Hg from seawater into the marine sediments (Grasby et al., 2019). One study suggested that seawater Hg isotope signals can be inherited by both TOC-poor and -rich sediments, following a small difference in $\delta^{202}\text{Hg}$ value (0.15‰) between background sediments and sapropel samples (Gehrke et al., 2009).

Subsequently, an experimental study indicated an enrichment of light Hg isotope in thiol phase ($\Delta^{202}\text{Hg}_{\text{Thiol-seawater}}$ = -0.5 to -0.6‰) (Wiederhold et al., 2010). If this constant isotope fractionation is considered, it is more likely that homogenous negative $\delta^{202}\text{Hg}$ values near -1.4‰ and positive $\Delta^{199}\text{Hg}$ (up to 0.25‰) in our chert and carbonate samples reflect the Ediacaran sediment background, at least in the Nanhua basin.
Figure S1: The positive relationship between total Hg concentrations and TOC (A) and TS contents (B). The sample with the highest total Hg concentrations (1443 ppm) is not included.

Figure S2: The negative relationship between $\Delta^{199}\text{Hg}$ values and Rb/Sr ratios for all samples from the Doushantuo Formation (A) and negative relationship between $\Delta^{199}\text{Hg}$ values and total Hg concentrations for samples from the PCIE-1 interval.
**Figure S3**: The negative relationship between $\delta^{202}\text{Hg}$ and $\Delta^{199}\text{Hg}$ values for samples from the PCIE-1 and -2 intervals. Two samples at 16.8m and 86.6m are not included.

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