Timing and provenance of volcanic fluxes around the Permian-Triassic Boundary Mass Extinction in South China: U-Pb zircon geochronology, volcanic ash geochemistry and mercury isotopes

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Abstract

Anomalous mercury (Hg) contents recorded near the Permian-Triassic boundary (PTB) are often linked to Siberian Traps Large Igneous Province (STLIP) volcanism and the Permian-Triassic boundary mass extinction (PTBME). However, mounting evidence indicates that the relation between STLIP volcanism and Hg "anomalies" is not straightforward. This study focuses on the timing and provenance of volcanic fluxes around the PTBME in South China. We constrain carbon isotope (δ^{13} C) and Hg concentration and isotope records by utilizing high-precision U-Pb zircon ages from two expanded deep-water marine sections spanning the Late Permian to Early Triassic in the Nanpanjiang Basin. Results reveal two episodes of Hg enrichment. The oldest episode predates the onset of a large negative $\delta^{13^{\alpha}} \epsilon \xi_{\zeta \cup \rho \sigma \iota \circ \gamma}$, which is documented to be older than 252.07 \pm 0.130 Ma. The second episode occurred between 251.822 \pm 0.060 Ma and 251.589 \pm 0.062 Ma, coinciding with the nadir of the δ^{13} C excursion. Volcanic ash geochemistry and Hg isotope compositions suggest that mercury was sourced from subduction-related volcanic arc magmatism in the Tethys region, which peaked between 251.668 \pm 0.079 Ma and 251.589 \pm 0.052 Ma. These results are compatible with suggestions that regional arc volcanism contributed to the causes of the PTBME in South China and provide evidence that Hg anomalies close to the PTB are not a reliable stratigraphic marker for the PTB extinction event. This study demonstrates that the relations between volcanism, environmental perturbations and mass extinction during the Permian-Triassic transition are better resolved with the aid of high-precision U-Pb zircon ages.

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2	Timing and provenance of volcanic fluxes around the Permian-Triassic Boundary
3	Mass Extinction in South China: U-Pb zircon geochronology, volcanic ash
4	geochemistry and mercury isotopes
5 6	Oluwaseun Edward ¹ , André Navin Paul ² , Hugo Bucher ³ , Christian Vérard ² , Thierry Adatte ⁴ , Jeroen E. Sonke ⁵ , Urs Schaltegger ² , Torsten Vennemann ¹
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16	Key Points:
17	• Mercury concentrations show elevated volcanic fluxes to South China before and after
18	the Permian-Triassic boundary.
19	• The recorded mercury anomalies are attributed primarily to regional subduction-related
20	arc volcanism, not the Siberian Traps.
21	• Hg anomalies close to the Permian-Triassic boundary are not a reliable stratigraphic
22	marker for the mass extinction event.

23 Abstract

24 Anomalous mercury (Hg) contents recorded near the Permian-Triassic boundary (PTB) are often

- 25 linked to Siberian Traps Large Igneous Province (STLIP) volcanism and the Permian-Triassic
- 26 boundary mass extinction (PTBME). However, mounting evidence indicates that the relation
- between STLIP volcanism and Hg "anomalies" is not straightforward. This study focuses on the
- timing and provenance of volcanic fluxes around the PTBME in South China. We constrain
- carbon isotope (δ^{13} C) and Hg concentration and isotope records by utilizing high-precision U-Pb zircon ages from two expanded deep-water marine sections spanning the Late Permian to Early
- Triassic in the Nanpanjiang Basin. Results reveal two episodes of Hg enrichment. The oldest
- episode predates the onset of a large negative δ^{13} C excursion, which is documented to be older
- than 252.07 ± 0.130 Ma. The second episode occurred between 251.822 ± 0.060 Ma and 251.589
- ± 0.062 Ma, coinciding with the nadir of the δ^{13} C excursion. Volcanic ash geochemistry and Hg
- isotope compositions suggest that mercury was sourced from subduction-related volcanic arc
- magmatism in the Tethys region, which peaked between 251.668 ± 0.079 Ma and $251.589 \pm$
- 0.052 Ma. These results are compatible with suggestions that regional arc volcanism contributed
- to the causes of the PTBME in South China and provide evidence that Hg anomalies close to the
- 39 PTB are not a reliable stratigraphic marker for the PTB extinction event. This study demonstrates 40 that the relations between volcanism, environmental perturbations and mass extinction during the
- 40 Permian-Triassic transition are better resolved with the aid of high-precision U-Pb zircon ages.

42 Plain Language Summary

43 Unusually high mercury contents in sedimentary rock sequences and the mass extinction of

- 44 organisms during the transition from the Permian–Triassic Period are often linked to Siberian
- 45 Traps volcanism. However, results from several studies indicate that the relationship between
- this massive volcanism and mercury peaks in Permian- to Triassic-aged rocks in the South China
- 47 region is complex. This study combines the geochemical and isotopic records of carbon and
- 48 mercury from Late Permian to Early Triassic sedimentary rocks, with absolute ages determined
- 49 from interlayered volcanic ashes, to investigate the presence, timing, and source of volcanic
- 50 inputs to these rock successions in South China. Results show higher mercury concentrations in 51 two intervals, one before and after the Permian-Triassic boundary. Absolute age results indicate
- 51 two intervals, one before and after the Permian-Triassic boundary. Absolute age results indicate 52 that the mercury peak closest to the Permian-Triassic boundary occurred around 300,000 years
- after the mass extinction. We attribute the mercury peaks to more local volcanic activity than the
- far away Siberian Traps and suggest that this regional-scale volcanic activity began shortly
- before 252 million years ago (Ma) and peaked around 251.6 Ma.

56 **1 Introduction**

- 57 The Paleozoic Mesozoic transition (ca. 252 million years ago) was characterized by the 58 largest mass extinction event in the Phanerozoic – the Permian-Triassic boundary mass
- 59 extinction (PTBME) (Erwin, 1998; Stanley, 2016). This event paved the way for the faunal
- 60 transition from the Paleozoic evolutionary fauna to the modern fauna (Dal Corso et al., 2022).
- 61 Furthermore, major perturbations of the global carbon and mercury cycles (expressed as negative
- 62 carbon (C) isotope excursions and mercury (Hg) concentration spikes, respectively) are
- 63 documented for sedimentary successions straddling the Permian-Triassic boundary (PTB) in
- 64 several spatially disparate localities (e.g., Baud et al., 1989; 1996; Grasby et al., 2013; Korte &
- 65 Kozur, 2010; Sanei et al., 2012; J. Shen et al., 2019; S. Shen et al., 2013; Sial et al., 2020).
- 66 Causal mechanisms for the extinction are still debated. However, the most popular trigger

67 suggested by researchers is the temporally overlapping volcanic activity of the Siberian Large

Igneous Province (STLIP), based on a temporal overlap between U-Pb ages of STLIP rocks and

69 those of volcanic ash beds in the Meishan Global Stratotype Section and Point (GSSP) (Burgess

70 & Bowring, 2015; Burgess et al., 2017).

The connection between volcanic activity and Hg concentration spikes in sedimentary 71 72 successions is founded on the understanding that volcanoes are the dominant natural source of Hg to the environment (Pyle & Mather, 2003). As such, LIP volcanism could result in the 73 emission and sequestration of high levels of Hg in the environment. Consequently, mercury 74 "anomalies" - relatively high Hg concentrations or Hg/TOC ratios in a portion of strata within a 75 given sedimentary succession – are used to trace the timing of Large Igneous Province (LIP) 76 magmatism in the geological record (Grasby et al., 2019; Percival et al., 2021; Yager et al., 77 78 2021). In the case of the δ^{13} C record, previous studies have interpreted the negative C isotope excursion associated with the PTB as the result of the intrusion of magma into surrounding 79 organic-rich country rocks (e.g., coal, shale) and petroleum-rich evaporites in the Tunguska 80 Basin, Siberia (Broadley et al., 2018; Payne & Kump, 2007; Svensen et al., 2009). Thus, Hg 81 anomalies and negative δ^{13} C excursions recorded close to the PTB are usually interpreted in the 82 context of STLIP volcanism and its associated deleterious environmental effects leading to 83

84 faunal extinction.

However, our understanding of the links between STLIP volcanism, carbon cycle 85 perturbations and mass extinction during the Permian-Triassic (P-T) transition are still 86 complicated by several factors. First, Hg and Hg/TOC content vary significantly near the PTB 87 between different localities and the relative timing of Hg anomalies with regards to the PTBME 88 horizon is inconsistent (e.g., J. Shen et al., 2019; Sial et al., 2020; Wang et al., 2018). In addition, 89 Hg anomalies can also be influenced by factors other than direct volcanic emissions. For 90 instance, Hg anomalies can arise from increased input of Hg remobilized from soils to marine 91 sediments due to increased erosion and continental weathering rates (Them II et al., 2019), as 92 93 well as enhanced sequestration by sulfides in euxinic depositional environments (J. Shen et al., 2020). In addition, sedimentary Hg records can be distorted by post-depositional weathering, 94 erosion, and thermal alteration (Charbonnier et al., 2020; Chen et al., 2022). Second, the 95 correlation of geochemical records between different localities is fraught with uncertainty 96 because the position of the PTB in many successions remains uncertain (e.g., Johnson et al., 97 2021; J. Shen et al., 2019; Sial et al., 2020). Also, sedimentary and volcanic successions 98 99 straddling the Permian and Triassic are often characterized by an unconformity at the PTB (Burgess & Bowring, 2015; Davydov, 2021; Yin et al., 2014), which for South China has been 100 101 estimated to represent a time gap of about 89 ± 38 kyr for the Permian part (Baresel et al., 102 2017a). Third, it has become apparent that several paleocontinents, including South China, experienced extensive regional intermediate to felsic volcanism during the P-T transition (Gao et 103 al., 2013; J. Shen et al., 2021; Vajda et al., 2020; Yin et al., 1992; H. Zhang et al., 2021), 104 105 increasing the potential sources of volcanic Hg input to PTB marine records at these localities. Fourth, absolute time calibration of the eruptive history of the STLIP remains coarse, limiting 106 precise temporal correlations between STLIP magmatism and mass extinction (Dal Corso et al., 107 2022), although, U-Pb geochronologic evidence suggests that the PTBME may have been limited 108 to only a short period of STLIP magmatism (Burgess et al., 2017). Consequently, considering the 109 complex biogeochemical cycling of Hg, the widespread incompleteness of the PTB rock record 110 (which limits the accuracy of chemostratigraphic and biostratigraphic correlations and the 111 placement of the PTB), and the occurrence of more proximal regional volcanic activity capable 112

of locally supplying Hg, links between Hg anomalies around the PTB and STLIP magmatism

cannot be unequivocally inferred without a thorough assessment of the provenance of the

recorded Hg anomalies and their timing relative to the PTBME using precise and accurate

116 geochronology.

Hg isotopes are widely used to trace the sources of Hg anomalies in ancient sedimentary 117 successions, as different transformations during the biogeochemical cycling of Hg can induce 118 mass-dependent (MDF) and/or mass-independent fractionation (MIF) of Hg isotopes (Blum et 119 al., 2014; Thibodeau & Bergquist, 2017). Previous studies have applied Hg isotopes in 120 investigating the source of Hg anomalies and its relationship to the PTBME in P-T successions 121 of South China and elsewhere (Grasby et al., 2017; J. Shen et al., 2021; J. Shen et al., 2019; 122 Wang et al., 2019a; Wang et al., 2018). However, these studies have provided mixed results, 123 with Hg MIF data from nearshore depositional settings reflecting a dominant terrestrial Hg 124 source and those from deeper-water depositional settings being congruent with atmospheric 125 volcanic Hg input (Grasby et al., 2017; Wang et al., 2018). Hence, it has been suggested that the 126 Hg isotope record for these successions mostly reflects their depositional setting (Yager et al., 127 2021) and that deep-water marine sections, being less vulnerable to terrestrial Hg inputs, may be 128 better suited for investigations of volcanic Hg input sources (Grasby et al., 2017; Thibodeau & 129 Bergquist, 2017; Wang et al., 2018). 130

The Upper Permian to Lower Triassic of South China is characterized by the widespread 131 occurrence of volcanic ash lavers (Gao et al., 2013; He et al., 2014; Yang et al., 2012; Yin et al., 132 1992), permitting precise and accurate U-Pb zircon geochronological calibrations of P-T 133 successions (e.g., Baresel et al., 2017a, 2017b; Burgess et al., 2014; Lehrmann et al., 2015; S. 134 Shen et al. 2019), as well as providing a means of evaluating the provenance of volcanic 135 products (e.g. zircon) to sedimentary successions in South China (e.g., Jiao et al., 2022; Zhao et 136 al., 2019). However, except for the Shangsi section (with published U-Pb zircon ages, S. Shen et 137 al., 2011), Hg anomalies recorded near the PTB from deep-water marine successions in South 138 139 China lack U-Pb zircon age calibration. Also, the published U-Pb zircon ages for Shangsi were not obtained with the most recent EARTHTIME spike (S. Shen et al., 2011; Yuan et al., 2019), 140 which makes a direct comparison with ages obtained using this spike problematic at high 141 temporal resolution. To accurately account for spike composition effects on the weighted mean 142 U-Pb age when comparing ages determined using different spikes, the tracer and analytical 143 uncertainty needs to be propagated (i.e., Y uncertainty, see section 2.5, not reported in Shen et al. 144 145 (2011)). This Y uncertainty propagation results in less-precise ages, inhibiting age comparison at high temporal resolution. In addition to enabling precise geochronology, volcanic ashes provide 146 an archive for the assessment of magmatic sources, as their geochemical compositions reflect 147 those of the source magma (e.g., He et al., 2014; Song et al., 2022; Yang et al., 2012). This study 148 presents, for the first time, a paired δ^{13} C and Hg (concentration and isotope) record calibrated by 149 precise and accurate U-Pb zircon ages from two sections (Laxian and Potuo) representing deep-150 water marine depositional environments in the Nanpanjiang Basin, South China. The aim of the 151 study is to assess the occurrence, timing and provenance of volcanic fluxes during the P-T 152 transition in South China. 153

154 1.1 Geological Context

The Potuo and Laxian sections (Fig. 1) are situated in the NE-SW trending Pingtang syncline, which in the present-day, is located in the northern part of the Nanpanjiang Basin, 157 South China (Bagherpour et al. 2020). The Changhsingian (Late Permian) sedimentary

- succession in Potuo is characterized by thin-bedded siliceous mudstones interbedded with
- volcanic ash layers: the Talung/Dalong Formation (Bagherpour et al., 2020; Baresel et al.
- 160 2017b). The Griesbachian (Induan, Early Triassic) consists of laminated black shales with 161 interbedded concretionary micritic limestones and occasional ash beds representing the
- interbedded concretionary micritic limestones and occasional ash beds representing the
 Daye/Ziyun Formation (Bagherpour et al., 2020). The Changhsingian to Griesbachian
- 162 sedimentary succession for Laxian is similar to that of Potuo except that there are no
- 164 concretionary micritic limestone beds interbedded with the Griesbachian shale units (Bagherpour
- 165 et al., 2020). The Late Permian to Early Triassic sedimentary sequence in these localities is
- 166 interpreted to have been deposited as a continuous section in a deep-water basin
- 167 paleoenvironment, hence without an unconformity between the Talung and Daye formations
- 168 (Bagherpour et al., 2020; Dai et al., 2019). As such, these sections are ideally suited for our
- 169 combined geochronologic and geochemical study, having been deposited in troughs within a
- horst-and-graben paleotopography (Bagherpour et al., 2020). In addition, the presence of
- volcanic ashes in these sections enables studies of latest Permian to earliest Triassic volcanic
 fluxes to the Nanpanjiang Basin of South China. Samples spanning the Upper Permian to Lower
- Triassic used for this study are the same as those analyzed by Bagherpour et al. (2020) and
- details on the geological setting, lithostratigraphy, biostratigraphy, as well as δ^{13} C record for
- 175 these sections are given in that study.

176 **2 Materials and Methods**

For this study, measurements of the Hg concentrations (n=70) and isotopic compositions (n=22), total organic carbon (TOC) contents (n=60), major and trace element concentrations for sedimentary rocks and interbedded volcanic ash beds (n=55) were made. These data are complemented by U-Pb zircon geochronology based on single zircon grains from five volcanic ash beds in the two sections (Table S1, supplementary information).

182 2.1 Stratigraphical correlation of the study sections

Stratigraphical correlation between the two sections studied – Laxian (25.78880°N, 183 107.29750°E) and Potuo (25.82638°N, 107.24861°E) follows Bagherpour et al. (2020). The PTB 184 is delineated based on the conformable lithostratigraphic boundary between the Talung and Dave 185 formations, which for deep-water sections of the Nanpanjiang Basin, has been shown to be 186 187 equivalent to the PTB as defined at the Meishan D GSSP based on U-Pb zircon ages (Burgess et al., 2014; Baresel et al., 2017b). The stratigraphical correlation of the Laxian and Potuo sections, 188 which is based on lithostratigraphy (Bagherpour et al. 2020) is congruent with the similarity of 189 the Hg trends for Potuo and Laxian, and is consistent with our new U-Pb zircon ages (see section 190 191 3).

192 2.2 Mercury concentration and isotopic composition

Mercury concentration was measured using a Zeeman R-915F high-frequency atomic absorption spectrometer at the University of Lausanne, Switzerland. Samples (sedimentary rocks, including volcanic ashes) were analyzed in triplicates to ensure analytical precision and the reference material – GSD-11, Chinese alluvium (Hg concentration of 72 ± 9 ppb; Zintwana et al., 2012) was used to monitor accuracy of the measurements (correlation coefficient = 0.99, standard residual deviation = 5 %).

Twenty-two samples (Potuo = 11, Laxian = 11) with sufficiently high Hg concentrations 199 $(\geq 9 \text{ ppb})$ were selected for Hg isotopic analysis at the Observatoire Midi-Pyrénées, Toulouse, 200 France. Mercury was preconcentrated using a double-stage tube furnace – acid-trapping protocol 201 as detailed by Sun et al. (2013). Powdered samples were loaded in quartz tubes (pre-cleaned at 202 550 °C), capped at both ends with quartz wool and heated in a combustion furnace connected to 203 a flow of Hg-free oxygen and set to ramp up from room temperature to 900 °C within 6 hours. 204 Liberated gaseous Hg⁰ was then purged into a decomposition oven, held at 1000 °C, using a 205 continuous flow of oxygen. Subsequently, the Hg⁰ was collected by oxidation to Hg^{II} in a 206 trapping solution of 40 % (v/v) inverse agua regia (iAR, 2HNO₃/1HCl), which was then diluted 207 at the end of the 6 h pre-concentration to 20 % (v/v) iAR and stored at 4 °C in the dark before Hg 208 isotopic measurements (Sun et al., 2013). Two certified reference materials, NIST 1632D 209 (bituminous coal, n=2) and MESS3 (Arctic marine sediment, n=2) were processed along with the 210 samples. Hg isotope compositions were subsequently measured in duplicate over two analytical 211 sessions by cold vapor multi-collector inductively coupled plasma mass spectrometry (CV-MC-212 ICP-MS) following analytical procedures detailed by Sonke et al., 2010; Sun et al., 2013 and 213 Jiskra et al., 2021. Briefly, the Hg^{II} in the iAR trap solution was reduced to Hg⁰ vapor using 214 SnCl₂ solution (3 %, w/v, in 1 M HCl) and then analyzed for Hg stable isotopes using an online 215 CETAC HGX-200 cold vapor generator coupled to a Thermo-Scientific Neptune PLUSTM 216 equipped with a $10^{12} \Omega$ resistor on the ¹⁹⁸Hg isotope. The MC-ICP-MS instrumental mass bias 217 was corrected by sample-standard bracketing using the NIST 3133 standard at matching standard 218 and sample concentrations (0.71 ppb and 2.1 ppb). Long-term instrumental precision was 219 monitored by repeated analysis of the ETH-Fluka (n = 6) and UM-Almaden (n = 3) Hg standard 220 solutions at Hg concentrations corresponding to the samples (i.e., 0.71 ng/g and 2.1 ng/g). 221 Procedural blanks had an average Hg concentration of ~ 0.01 ng/g (n=3). Mass-dependent 222 fractionation (MDF) of Hg isotopes is reported in small delta notation (δ) as δ^{202} Hg in permil 223 (‰) relative to the bracketing NIST 3133 standard: 224 $\delta^{202} \text{Hg} = [((^{202} \text{Hg}/^{198} \text{Hg})_{\text{sample}}/(^{202} \text{Hg}/^{198} \text{Hg})_{\text{NIST3133}} - 1) \times 10^3]$ 225 Mass independent fractionation (MIF) values are denoted using capital delta (Δ) notation and are 226 defined as the difference between the measured values of δ^{199} Hg, δ^{200} Hg, δ^{201} Hg, δ^{204} Hg and 227 those predicted for MDF with respect to δ^{202} Hg using the kinetic MDF law as follows: 228 $\Delta^{xxx}Hg = \delta^{xxx}Hg - K_{xxx} \times \hat{\delta}^{202}Hg$ 229 (2)where xxx refers to Hg isotope masses 199, 200, 201, and 204, and K_{xxx} refers to the constants 230 that are used to calculate values for Δ^{xxx} Hg, which are: 0.2520, 0.5024, 0.7520 and 1.4930 for 231 δ^{199} Hg, δ^{200} Hg, δ^{201} Hg and δ^{204} Hg respectively (Blum & Bergquist, 2007). Hg isotopic 232 compositions are reported as the mean of duplicate measurements and analytical uncertainty of 233 isotopic analysis are reported conservatively, as either the 2σ (2 × standard deviation) of the 234 replicate sample measurements or that of the standard with the largest 2σ , whichever was larger. 235 ETH-Fluka and UM-Almaden standard solutions yielded mean values ($\pm 2\sigma$) of -1.48 ± 0.14 ‰, 236 0.09 ± 0.03 ‰, 0.04 ± 0.03 ‰, 0.04 ± 0.07 ‰; and -0.59 ± 0.09 ‰, -0.03 ± 0.10 ‰, 0.02 ± 0.06 237 % and -0.05 \pm 0.07 % for δ^{202} Hg, Δ^{199} Hg, Δ^{200} Hg, and Δ^{201} Hg respectively. NIST 1632D and 238 MESS-3 had mean values of -1.93 ± 0.32 ‰, -0.01 ± 0.05 ‰, -0.06 ± 0.08 ‰, -0.06 ± 0.01 ‰; 239 and -2.25 ± 0.02 ‰, 0 ± 0.04 ‰, 0.04 ± 0.03 ‰ and -0.14 ± 0.14 ‰ for δ^{202} Hg, Δ^{199} Hg, Δ^{200} Hg, 240 and Δ^{201} Hg respectively. These mean values are comparable with reported values for these 241 standard solutions and certified reference materials (Jiskra et al., 2019; Kwon et al., 2015; Sun et 242 243 al., 2013).

244 2.3 Organic carbon content

All total organic carbon (TOC) content data for Laxian (n=26) were acquired during the present study. For the Potuo section, 10 data points are from Bagherpour et al. (2020) and an additional 28 samples have been analyzed for this study. Organic matter content and preservation was assessed by Rock-Eval pyrolysis using a Rock EvalTM 6 with the analytical procedures described by Behar et al. (2001). This included measurements of total organic carbon (TOC) content, pyrolysis temperature (T_{max}), hydrogen index (HI) and oxygen index (OI). The standard IFP160000 was used for calibration of the samples and instrumental precision was about 0.1

- wt.% for TOC, 10 mg HC/g for HI and 10 mg CO_2/g for OI.
- 253 2.4 Major and trace element analysis

Major and trace element concentrations were analyzed by X-ray fluorescence (XRF)
 spectrometry on glass discs and pressed tablets, respectively, using a PANalytical PW2400 XRF
 spectrometer at the University of Lausanne, Switzerland. The standard reference materials: JLS 1, JDO-1 and BHVO were used for assessment of analytical accuracy. External reproducibility
 is between 0.5 % and 5 % depending on the element, and detection limit for major elements

is ca. 0.01 % and between 1 and 7 ppm for trace elements.

260 2.5 U-Pb zircon geochronology

In total, nine volcanic ash layers from the Talung (Late Permian) and Daye (Early 261 Triassic) formations at Potuo (POT66T, POT67T, and POT68T) and Laxian (LAX8T, LAX9T, 262 LAX10T, LAX11T, LAX 13T and LAX14T) were processed for U-Pb zircon dating. Of these, 263 seven ash layers yielded sufficient zircon crystals for CA ID-TIMS U-Pb analyses. Zircon U-Pb 264 isotopic compositions were determined by chemical abrasion, isotope dilution, thermal ionization 265 mass spectrometry (CA-ID-TIMS) at the University of Geneva, Switzerland, following the 266 267 procedure described by Widmann et al. (2019). Zircons were extracted from ash beds by conventional methods (crushing, milling, sieving, magnetic and density separation), including 268 hand-picking of high aspect ratio grains free of visible inclusions. The zircon grains were 269 thermally treated at 900 °C for 48 h to stimulate self-annealing prior to partial dissolution in 270 concentrated hydrofluoric acid (HF_{conc}), to remove structurally damaged domains that may have 271 suffered lead (Pb) loss (see Widmann et al. (2019) for the detailed procedure). The partially 272 273 dissolved grains were then extracted and washed in 6N HCl in 3 ml Savillex beakers overnight (min. 12 h), at 80 °C. Further cleaning steps involve cycling of 7N HNO₃ and ultra-sonic bathing 274 prior to loading into 200 μ l capsules for dissolution in 2 to 3 drops of HF_{conc} for 48 h at 210 °C in 275 pressurized ParrTM vessels. A ²⁰²Pb-²⁰⁵Pb-²³³U-²³⁵U tracer solution: ET2535 (EARTHTIME 276 2535, Condon et al., 2015) was added prior to dissolution, and Pb and U were isolated using ion 277 exchange resin chromatography. Uranium and Pb isotopic compositions were measured on an 278 279 IsotopX Phoenix TIMS at the University of Geneva. Pb was measured using a dynamic peak jumping routine on a Dalv ion counting system, while uranium was measured as UO₂ in static 280 mode using $10^{12} \Omega$ resistor Faraday amplifiers for samples POT66T, POT67T, POT68T, and 281 using ATONA Faraday amplifiers (with a 30 s integration time) for samples LAX8T and 282 LAX10T. The measured isotopic ratios were corrected for interferences of ²³⁸U¹⁸O¹⁶O on 283 235 U¹⁶O₂ using a 18 O/¹⁶O composition of 0.00205 based on repeat measurements of the U500 284 standard. Mass fractionation of U was corrected using a double isotope tracer with a ²³⁵U/²³³U of 285 0.99506 ± 0.005 . The Pb blank isotopic composition is ${}^{206}\text{Pb}/{}^{204}\text{Pb} = 17.43 \pm 0.71 (1\sigma)$, 286

287 ${}^{207}\text{Pb}/{}^{204}\text{Pb} = 14.73 \pm 0.38 \text{ (1}\sigma\text{)} \text{ and } {}^{208}\text{Pb}/{}^{204}\text{Pb} = 35.58 \pm 1.04 \text{ (1}\sigma\text{)}, \text{ based on total procedural blank measurements.}$

U-Pb dates of zircons were calculated using data reduction software Tripoli and ETRedux 289 290 (Bowring et al., 2011; McLean et al., 2011)), and all uncertainties are reported at the 95 % confidence interval. The interpreted weighted mean age is reported in the format: "weighted 291 mean" $\pm /X/Y/Z$, where X corresponds to analytical uncertainty, Y to analytical + tracer 292 uncertainty and Z to analytical, tracer and decay constant uncertainty (Schoene et al., 2006). All 293 zircon 206 Pb/ 238 U dates were corrected for 230 Th- 238 U disequilibrium using a Th/U_{magma} of 3.5 ± 294 1.0. We discuss dates at the precision level of analytical uncertainty (X), since most U-Pb data in 295 the relevant literature were obtained using the same isotope tracer (ET2535) and mode of 296 analysis (ID-TIMS), effectively eliminating inter-lab uncertainty. Repeat analyses of the ET100 297 solution $(^{206}\text{Pb}/^{238}\text{U} \text{ age: } 100.173 \pm 0.003 \text{ Ma; Schaltegger et al., } 2021)$ yielded a value of 298 100.1678 ± 0.0046 Ma (mean square weighted deviation (MSWD) = 3.2, n = 32/40). One batch 299 comprising 8 ET100 samples was rejected due to an anomalous young average age (this batch is 300 internally consistent). Similarly young, anomalous ET100 ages observed in Schaltegger et al. 301 (2021) were explained by fractionation effects during the sample preparation process. Inclusion 302 of this rejected batch would result in an average ET100 age of 100.1639 ± 0.0039 Ma (MSWD = 303 3.8, n = 39/40). Excess scatter is indicated by the elevated MSWD value and is commonly 304 observed for the synthetic ET100 solution. This is potentially derived from instrument instability 305 306 and from the sample preparation process (Schaltegger et al., 2021).

307 **3 Results**

Results of all geochemical analyses are given in the supplementary information and presented in figures 2 to 5.

310 3.1 U-Pb zircon geochronology

311 A total of 98 zircons from 7 volcanic ash beds were analyzed. Zircon U and Pb data are presented in the supplementary information and interpreted U-Pb dates are illustrated in Fig. 2. 312 For each volcanic ash bed, the U-Pb weighted mean date is interpreted on the following basis: i) 313 not rejecting any analyses that are concordant, ii) the youngest cluster of interpreted concordant 314 zircons consists of > 3 analyses and iii) assuming that all Pb-loss is effectively removed by the 315 chemical abrasion procedure. This data reduction strategy is in line with previous U-Pb ID-TIMS 316 317 studies concerning volcanic ash beds straddling the Permian-Triassic Boundary and the Early Triassic (e.g., Augland et al., 2019; Baresel et al., 2017b; Burgess et al., 2014; Widmann et al., 318 2020), making all of these previous ID-TIMS ages directly comparable. 319

Only in one sample (POT 66T) do we reject one younger, concordant analysis, as it is 320 younger than the statistically significant, slightly older age plateau of 12 analyses (Fig. 2). We 321 assume that this deviation is due to unresolved, residual lead loss not mitigated by the chemical 322 abrasion procedure. Analyses that are older than the youngest, statistically valid, date plateau are 323 considered detrital or due to inherited or antecrystic cores. The principal guideline for the 324 youngest weighted mean age determination is a statistically valid MSWD for the chosen age 325 cluster. Applying this strategy, volcanic ash beds LAX8T, LAX10T, POT66T, POT67T and 326 POT68T have statistically significant 206 Pb/ 238 U weighted mean ages of 252.07 ± 0.13/0.15/0.31 327 Ma (Mean Square Weighted Deviation (MSWD) = 1.3, n = 5), $251.822 \pm 0.060/0.089/0.28$ Ma 328 $(MSWD = 1.9, n = 6), 251.589 \pm 0.052/0.083/0.28 \text{ Ma} (MSWD = 1.67, n = 12), 251.668 \pm 0.052/0.083/$ 329 0.079/0.10/0.29 Ma (MSWD = 1.99, n = 9) and $251.64 \pm 0.13/0.15/0.31$ Ma (MSWD = 1.5, n = 330

12), respectively (Fig. 2). Ash beds LAX14T and LAX11T (Table S1) did not satisfy the data
 reduction criteria outlined above and are thus not interpreted.

333 3.2 TOC contents and Hg concentrations

TOC content is stratigraphically variable and generally low for Laxian relative to Potuo 334 with a range between < 0.1 - 0.3 wt.% and 0.1 - 2.9 wt.% respectively (Fig. 3). TOC generally 335 shows moderate correlation for Laxian ($r^2 = 0.36$) and Potuo ($r^2 = 0.50$) (Fig. 4). Only 3 samples 336 for Laxian have TOC content above 0.2 wt.%, a suggested threshold for robust Hg/TOC 337 normalization (Grasby et al., 2016). In contrast, only 6 out of 38 samples for Potuo have TOC 338 values < 0.2 wt.%. Therefore, Hg data for Laxian are not normalized to TOC. OI values for 339 Laxian are high, and have a range between 69 and 1386 mg CO₂/g TOC. OI values are between 3 340 and 334 mg CO₂/g TOC for Potuo, except for volcanic ash samples with values between 64 and 341 967 mg CO₂/g TOC. HI values are between 19 and 283 mg HC/g TOC and between 17 and 229 342 343 mg HC/g TOC for Potuo and Laxian respectively.

Hg concentrations of samples from the Talung Fm. in Potuo are between 2 ppb and 18 344 ppb and are generally lower than for Laxian, which have a range between 9 ppb and 64 ppb. The 345 main feature of the latest Permian Hg record for Potuo is a minor Hg/TOC excursion (referred to 346 as E1 here) with a peak Hg/TOC value of 70 ppb/wt.%. In the Daye Fm., a significant Hg 347 concentration increase (here labelled as E2) is recorded in the lower part of the successions at 348 both sections. This Hg excursion is simultaneously expressed by the Hg and Hg/TOC records for 349 Potuo (Fig. 3). Peak Hg concentrations within E2 Hg anomaly (Laxian = 251 ppb, Potuo = 203350 ppb) are higher than the mean so far reported (62 ppb) for sedimentary rocks (Grasby et al., 351 352 2019).

Volcanic ashes from the Talung Fm. at Potuo have Hg concentrations that are at least one order of magnitude higher than for interbedded siliceous mudstones. This difference does not hold in the Daye Fm., where Hg concentration values range between 68 ppb and 165 ppb for volcanic ashes and between 84 ppb and 203 for interbedded rocks within the E2 Hg anomaly interval (Fig. 3b).

358 3.3 Mercury isotopes

Mass dependent fractionation of Hg isotopes (δ^{202} Hg) shows near-zero to negative values for both sections throughout the studied interval (range: +0.23 ± 0.32 ‰ to -1.75 ± 0.32 ‰; mean = -1.13 ± 1.02 ‰), except for 2 volcanic ash samples from Potuo with more negative values (Fig. 5, POT66T: -2.49 ± 0.32 ‰ and POT67T: -2.29 ± 0.32 ‰).

Hg isotope MIF (Δ^{199} Hg and Δ^{200} Hg) values for both sections are near-zero throughout 363 the studied interval (i.e., $0.1 \ \% > z > -0.1 \ \%$, where z = sample Hg MIF value), except for 364 sample POT 59, which has a Δ^{199} Hg value of 0.12 ± 0.11 ‰. Mean ($\pm 2\sigma$) Δ^{199} Hg values are 365 0.02 ± 0.11 ‰ and -0.01 ± 0.11 ‰ for Potuo and Laxian respectively. For Potuo, the mean 366 Δ^{200} Hg value is -0.01 ± 0.10 ‰ and for Laxian, it is 0.00 ± 0.05 ‰. Hence, the Pingtang syncline 367 sections record no measurable mass independent fractionation of Hg isotopes, in contrast to Hg 368 isotope records for deep-water marine sections outside the Nanpanjiang Basin in South China 369 (Fig. 3c). 370

371 3.4 Major and Trace Elements

Al₂O₃ concentrations for both sections show similar patterns, having lower values in the 372 Talung Fm. (with a range of 2 to 4 wt.%, except for 1 Potuo sample) relative to the Dave Fm. For 373 Laxian, Al₂O₃ values in the Daye Fm. are $3 \times$ higher (range: 15 - 22 wt.%) than those of the 374 Talung Fm. Fe_2O_3 concentrations show a similar trend to Al₂O₃, with lower values in the Talung 375 376 Fm. (1 to 5 wt.%) relative to the Daye Fm. for both localities. However, unlike Al₂O₃, values in the Daye Fm. for both localities are identical (between 2 and 9 wt.%). Al and Fe (proxied by 377 Al₂O_{3 and} Fe₂O₃) show little to no correlation with Hg for both Potuo (Al: $r^2 = 0.02$, Fe: $r^2 = 0.23$) 378 and Laxian (A1: $r^2 = 0.27$, Fe: $r^2 = 0.16$). Mo and U are redox-sensitive trace elements used to 379 track redox variations in sedimentary environments (Algeo & Maynard, 2004; Hardisty et al. 380 2018; Tribovillard et al., 2006). Redox-sensitive trace element concentrations are usually 381 normalized to Al to account for variations that may be unrelated to changes in redox conditions, 382 such as changes in sediment input or authigenic mineral formation (e.g., Algeo & Maynard, 383 2004; Grasby et al., 2013; Rolison et al., 2017). For Laxian, 73 % of samples analyzed have Mo 384 contents lower than the lower limit of detection (LLD) of the XRF spectrometer (i.e., 1 ppm). 385 Mo/Al values show no stratigraphic trend for either studied locality (Fig. 5e) and have no 386 correlation with Hg contents (Potuo: $r^2 = 0.04$, Laxian: $r^2 = 0.06$; Fig. 4D). Laxian samples with 387 measurable Mo (n = 7) have Mo/Al values between 0.1 and 0.6 ppm/wt.%. Mo/Al values for 388 389 Potuo are slightly higher and range between 0.1 and 1.1 ppm/wt.%, except for sample POT 80 with a value of 5.7 ppm/wt.%. Potuo U/Al values range between 0.2 and 2.3 ppm/wt.%, except 390 for sample POT 63 (U/Al = 4.9 ppm/wt.%). For Laxian, U/Al ranges between 0.2 and 0.8 391 ppm/wt.%. As with Mo/Al, no correlation is observed between Hg and U/Al (Potuo: $r^2 = 0.10$, 392 Laxian: $r^2 = 0.03$; Fig. 4b). 393 Volcanic ash samples have high LOI (loss on ignition) values between 9 to 14 wt.%, 394 consistent with results from contemporaneous ash layers in South China (He et al., 2014). The 395 ash samples have LOI-corrected mean $(\pm 1\sigma)$ concentrations of 58 ± 5 wt.%, 25 ± 2 wt.% and 5 396 \pm 3 wt.% for SiO₂, Al₂O₃ and total alkali (Na₂O+K₂O) respectively (Table S2, supplementary 397 398 information). To mitigate the chemical effects of secondary alteration of the volcanic ashes, only immobile elements (such as Th, Nb, Ta, Zr, Hf, Ti, Y), known to be unaffected by post-399 depositional alteration (Portnyagin et al., 2020), are utilized for subsequent chemical 400 classification and tectonic interpretation (Fig. 6). The ashes mainly plot within the fields of 401 rhyodacite/dacite, andesite and trachyandesite on the Nb/Y vs. Zr/Ti diagram (Winchester and 402

Floyd, 1977), separate from STLIP volcanic rocks, except for 1 sample: POT 14T (Fig. 6a).
Incompatible trace elements normalized against primitive mantle values (Sun and McDonough,
1989) show that the volcanic ashes are characterized by pronounced negative Nb, Ta and Ti
anomalies (Fig. 6d), similar to the trace element pattern for volcanic ashes from other localities
in South China (He et al., 2014; Yang et al., 2012).

408 4 Discussion

409 4.1 Sedimentary hosts of Hg and interpretation of the Hg anomalies

410 4.1.1 Hosts of Hg

Hg enrichment in sediments may reflect enhanced Hg sequestration related to increased
abundance of its sedimentary host phase(s) or enhanced Hg input fluxes during sediment
deposition (Grasby et al., 2019; J. Shen et al., 2020). Due to the high affinity of Hg for organic

414 matter (OM), Hg concentration is usually normalized to TOC content to account for any OM

increases that could have preferentially enhanced Hg sequestration in sediments (e.g., Sanei et al., 2012; Grasby et al., 2017, 2019). However, in samples with low OM content (< 0.2 wt.%

al., 2012; Grasby et al., 2017, 2019). However, in samples with low OM content (< 0.2 wt.%
 TOC), the Hg/TOC ratio is deemed unreliable as the Hg/TOC values become exaggerated,

producing false Hg/TOC peaks (Grasby et al., 2019; but see Yager et al., 2021). In addition to

TOC variations, Hg sequestration in sedimentary rocks can be influenced by other sedimentary

host phases such as clay minerals, iron oxides and sulfides (Charbonnier et al., 2017; J. Shen et al. 2020)

421 al., 2020).

Based on cross plots of Hg vs. TOC, Al, Fe, Mo/Al and U/Al values for both sections, Hg is best correlated to TOC (r^2 : Potuo = 0.50 and Laxian = 0.36, Fig. 4c). This correlation suggests that Hg sequestration in the studied sites was partially controlled by organic matter availability and may explain the more efficient sequestration of Hg in the black shales of the Daye Formation relative to the siliceous mudstones of the Talung Formation. Nevertheless, plots of Hg/Al and Hg/Fe ratios for both localities, including Hg/TOC for Potuo (Fig. 3; Fig 5) show similar trends

of peak Hg contents in the lower part of the Griesbachian record, suggesting that the Hg anomaly

429 in this interval cannot be explained by TOC variability or clay mineral inputs alone.

430 Furthermore, lithological changes are unlikely to control the Hg concentration spike as increases

in Hg concentration do not coincide with the change in lithology in either of the two sections.

432 Moreover, relatively high and low Hg concentrations alike are measured for the Talung and

433 Daye formations where Hg and Hg/TOC anomalies are recorded (Fig. 3; Fig. 8).

434 4.1.2 Evaluation of Hg preservation

As OM is the dominant host of Hg in the studied successions, the potential impact of 435 post-depositional OM degradation on the Hg record warrants evaluation. Rock-Eval TOC data 436 cross plots (HI, OI, T_{max}) (Fig. 7) are routinely used to evaluate the type and maturity of 437 preserved OM in sedimentary successions (e.g., Charbonnier et al., 2020; Espitalié et al., 1985; 438 Fantasia et al., 2018). The low HI and OI values of many of the samples with enough OM for 439 Rock-Eval data interpretation (i.e., TOC > 0.2 wt.%, Fig. 7a) may indicate diagenetic alteration 440 of marine OM (altered type II) and/or high input of terrestrial OM (type III) (Charbonnier et al., 441 2020; Fantasia et al., 2018). The range of T_{max} values (400 – 525 °C) suggests that some of the 442 OM in the studied successions are thermally mature and have undergone post-depositional 443 oxidation (Fig. 7b; Espitalié et al., 1985). The thermal maturity of preserved OM in some strata 444 of the studied successions suggests that original Hg contents may have been diagenetically 445 modified (Charbonnier et al., 2020). Consequently, the Hg content measured for strata 446 447 characterized by thermally mature OM may be a minimum estimate of the original Hg content of these rocks. 448

Despite the probable partial loss of the original Hg content for some samples, several 449 points argue in favor of the reliability of the general Hg trend documented in the present study. 450 The Changhsingian to Griesbachian Hg trend for both localities is similar despite the vast 451 difference in OM content (Fig. 3). The lack (or loss) of OM is more prevalent in the Laxian 452 succession, yet the background Hg content in this succession is $3 \times$ that of Potuo. Also, volcanic 453 ashes mostly have similar Hg content in the Talung and Daye Fm. In contrast, interbedded rocks 454 differ strongly in Hg content between these rock formations (Fig. 3). The much higher Hg 455 content of volcanic ashes despite their higher susceptibility to weathering (e.g., Jiao et al. 2022) 456 and as such, Hg loss, suggests that: i) Hg is reasonably well preserved in these strata, and ii) that 457 the disparity in Hg content between volcanic ashes and interbedded rocks is primary. 458

Furthermore, both sections record the E2 Hg anomaly, and with a similar excursion 459 magnitude (Fig. 3, Fig. 8). This similarity in the Hg trends and hence, their spatial reproducibility 460 suggests that the primary Hg concentration trends are preserved. In addition, there is no 461 correlation between HI and TOC (Fig. 7c) and little correlation ($r^2 = 0.30$) between HI and Hg 462 values (Fig. 7d), suggesting that OM maturity does not influence Hg trends to any large degree in 463 these successions. Also, the E2 Hg anomaly coincides with the climax of the negative δ^{13} C 464 excursion at the P-T transition (Fig. 3, Fig. 8), suggesting that the Hg trend is controlled by 465 environmental perturbations (as indicated by the δ^{13} C excursion), and not diagenetic alteration. 466 Finally, Charbonnier et al. (2020) noted that despite the oxidative weathering of OM observed 467 for weathered rock samples, there were no significant changes in Hg/TOC ratios, suggesting that 468 the Hg/TOC ratio is less susceptible to the effects of post-depositional OM degradation. In the 469 present study, both Hg anomalies reported are present in the Hg/TOC record and more 470 significantly, the main excursion (E2) is present in both the Hg and Hg/TOC record, suggesting 471 that these Hg trends are primary. 472

473 4.1.3 Hg isotopes

Hg isotopes are used to trace the source(s) and depositional pathway(s) of Hg to natural 474 environments, as Hg isotope mass-dependent fractionation (δ^{202} Hg) and mass-independent 475 fractionation (MIF, Δ^{199} Hg and Δ^{200} Hg) compositions vary across Earth surface reservoirs and 476 transport mechanisms (Bergquist & Blum, 2007; Blum et al., 2014; Fu et al., 2021). Here, we 477 focus on Hg isotope MIF, as this occurs via fewer processes compared to MDF (Blum et al., 478 2014). Also, δ^{202} Hg values of direct volcanic emissions overlap with those of terrestrial runoff 479 and atmospheric Hg^{II} deposition (Yager et al., 2021), making them less diagnostic than Hg 480 isotope MIF values. 481

Hg isotope MIF values are considered to be resistant to diagenetic alteration (e.g., Grasby 482 et al. 2017; Thibodeau et al., 2016). This view was recently strengthened by the experimental 483 study of Chen et al. (2022), who documented that high-temperature or high-pressure alteration 484 of rocks does not result in alteration of Δ^{199} Hg and Δ^{200} Hg values. Δ^{199} Hg is commonly used to 485 interpret the sources and pathways of Hg deposition (Thibodeau & Bergquist, 2017; Yager et al., 486 2021) and recently, Δ^{200} Hg has been proposed as a complementary tracer of Hg sources to land 487 and oceans (Jiskra et al., 2021). This proposition is because even number-Hg isotope MIF (e.g., 488 Δ^{200} Hg) only occurs via upper atmospheric oxidation-reduction pathways and thus, Hg 489 transformations near Earth's surface yield no measurable even number-Hg isotope MIF (Chen et 490 al., 2012; Fu et al., 2021). Hg from terrestrial biomass usually has negative Δ^{199} Hg values while 491 oceanic reservoirs (e.g. marine sediments and seawater) are characterized by near-zero to 492 positive Δ^{199} Hg values (Blum et al., 2014; Yin et al., 2022). Furthermore, Hg released to the 493 atmosphere may undergo MIF via photochemical reactions during its cycling, acquiring positive 494 or negative MIF values (Blum et al, 2014) before long-term burial in marine sediments 495 (Thibodeau et al., 2016). Hg derived from direct volcanic emission is thought to have no 496 measurable MIF (i.e., Δ^{199} Hg, Δ^{200} Hg = ~ 0 ‰, Thibodeau & Bergquist, 2017; Zambardi et al., 2009), consistent with a recent estimate of the Δ^{199} Hg value of the primitive mantle (0.00 ± 0.10 497 498 ‰, Moynier et al., 2021). Therefore, near-zero Δ^{199} Hg values recorded for rock samples in the 499 geologic record have been interpreted as reflecting unaltered volcanic Hg input to the 500 depositional environment (e.g., Font et al., 2021; Thibodeau et al., 2016; Yager et al., 2021; H. 501 Zhang et al., 2021). Alternatively, near-zero Δ^{199} Hg values have also been interpreted as 502

reflecting Hg contributions from a combination of terrestrial and oceanic Hg reservoirs (e.g., J.
Shen et al., 2019; 2021; Wang et al., 2019a).

The near-zero and invariant Δ^{199} Hg and Δ^{200} Hg values throughout our studied interval 505 (Fig. 8) are consistent with direct volcanic Hg input from a volcanic center close to the 506 Nanpanjiang Basin, or a mixture of terrestrial and marine Hg sources. Several lines of evidence, 507 however, argue in favor of the former as an explanation for the Hg MIF record of the studied 508 successions. First, the lack of correlation between Hg and Al suggests that Hg sequestration was 509 unrelated to terrestrial input to the Pingtang syncline. As terrestrial input is characterized by 510 negative Hg isotope MIF values (Thibodeau & Bergquist, 2017; Yager et al., 2021; Yin et al., 511 2022), enhanced clastic input during the Griesbachian (as evidenced by dominantly shale 512 lithology and increase in Al content, Fig. 5) is expected to have resulted in more negative Δ^{199} Hg 513 values. However, Δ^{199} Hg values immediately after the PTB are near-zero to slightly positive for 514 both sites(Fig. 3), suggesting that atmospheric volcanic Hg input was the dominant source of Hg 515 to the Nanpanjiang Basin. Second, the Nanpanjiang Basin was situated close to a volcanic center 516 during the P-T transition. This proximity is evidenced by the occurrence of thicker volcanic ash 517 beds in coeval marine successions in southwest South China (i.e., in the Nanpanjiang Basin) 518 519 relative to the rest of South China (He et al., 2014; Zhao et al., 2019), the occurrence of coeval acidic-intermediate volcanic and intrusive rocks in southwestern South China and northern 520 Vietnam, as well as copper concentration and isotope excursions linked to felsic "super-521 522 eruptions" in South China during the Changhsingian (H. Zhang et al., 2021 and references therein). The paleo-location of the Nanpanjiang Basin may, therefore, also explain the disparity 523 between the Pingtang syncline Hg MIF record and those previously published for other parts of 524 South China (e.g., J. Shen et al., 2021; Wang et al., 2019a; Fig. 3c) (discussed in section 4.4). 525 Third, the Hg isotope MIF values for volcanic ash samples are indistinguishable from those of 526 overlying and underlying strata, suggesting that the source of Hg for volcanic ashes (TOC = ~ 0) 527 and interbedded rocks (TOC = 0 - 3 wt.%) were the same. Consequently, the Hg and Hg/TOC 528 anomalies recorded for both localities are interpreted as dominantly reflective of pulses of 529 elevated atmospheric volcanic Hg input to these deep-water marine depositional sites. 530

531

4.2 U-Pb zircon age constraints on the Hg anomaly and C isotope excursion

Although previous works (Grasby et al., 2017; J. Shen et al., 2019; Sial et al., 2020) 532 suggested a coeval global occurrence of Hg anomalies around the PTBME, the majority of 533 sections with defined Hg anomalies around the P-T transition lack high-resolution, high-534 precision geochronology. Our study attempts to specifically bracket the age of a well-defined Hg 535 anomaly in the Early Triassic via dating of under and overlying ash beds in expanded deep-water 536 537 marine records. The new U-Pb zircon ages from the Pingtang syncline span the onset of the latest Permian negative δ^{13} C excursion, ca. 252.07 ± 0.130 Ma and brackets a Griesbachian Hg 538 anomaly. The two ash layers analyzed from Laxian straddle the PTB and the U-Pb ages are 539 540 consistent with the stratigraphy, yielding U-Pb weighted mean ages of 252.07 ± 0.13 Ma (LAX8T, latest Permian) and 251.822 ± 0.060 Ma (LAX10T, Griesbachian) respectively. These 541 U-Pb zircon ages overlap, within analytical error, with the ages of Bed 25 (251.941 ± 0.037 Ma) 542 543 and Bed 28 (251.880 \pm 0.031 Ma) of the very condensed Meishan Global Stratotype Section and Point (GSSP) (Burgess et al., 2014). Thus, the interval between LAX8T and LAX10T includes 544 the PTBME interval at Meishan and also agrees with previous suggestions that the conformable 545 lithological boundary between the Talung/Dalong and Daye/Ziyun formations accurately 546 delineates the PTB in deep-water marine successions in the Nanpanjiang Basin (Bagherpour et 547

al., 2020; Baresel et al., 2017b). Based on lithological comparisons between Potuo and Laxian, 548 549 the E1 anomaly in Potuo is stratigraphically below LAX 8T (Baghgerpour et al., 2020) and as such, likely predates the mass extinction interval. However, the lack of U-Pb ages in the strata 550 spanning E1 makes quantitative comparisons to LAX 8T uncertain. 551

The overlap between the 206 Pb/ 238 U weighted mean ages of ash beds POT 66T, 67T and 552 68T suggests the occurrence of several successive volcanic eruptions (within the time covered by 553 these volcanic ash layers) at intervals shorter than the resolving power of our ID-TIMS U-Pb 554 geochronology at the given quality of zircon available for this study. Nevertheless, as these 555 volcanic ash layers from Potuo straddle both the Hg anomaly recorded in the Daye Fm., as well 556 as the coeval nadir of the negative δ^{13} C excursion (Fig. 8), the age of both the Hg anomaly (E2) 557 and peak C-cycle perturbation during the P-T transition can be determined. The youngest age for 558 both the peak of the E2 Hg anomaly and the nadir of the δ^{13} C excursion is 251.589 ± 0.052 Ma 559 (POT66T, Fig. 8), while the oldest age estimate is 251.668 ± 0.079 Ma (POT67T). These two 560 ages are identical within the analytical (X) uncertainty, precluding an estimation of the duration 561 of the Hg anomaly. Despite these uncertainties, we can establish with confidence that the post-562 PTB Hg anomaly peak observed in Potuo is ca. 300 kyr younger (largely outside of analytical 563 uncertainty) than the Meishan PTBME interval (Burgess et al., 2014; Burgess and Bowring, 564 2015), as well as the extinction horizon in Penglaitan (251.939 ± 0.031 Ma; S. Shen et al., 2019). 565

566

4.2.1 Hg anomalies as a PTBME correlation tool

A negative δ^{13} C excursion at the P-T transition together with a Hg anomaly are common 567 features of many PTB-straddling sedimentary successions (Fig. 9). As such, the peak of the Hg 568 anomaly and/or the nadir of the negative $\delta^{13}C$ excursion associated with the PTB are often 569 considered to be stratigraphic markers for the PTB extinction interval (e.g., Grasby et al., 2017; 570 Sial et al. 2020; J. Shen et al. 2019, 2023; Wignall et al., 1998). Although some sedimentary 571 successions, especially in high latitudes, show Hg excursions coincident with both the negative 572 δ^{13} C excursion and mass extinction (e.g., Sanei et al., 2012; Grasby et al., 2013), it is apparent 573 574 that Hg excursions in several other successions straddling the PTB vary in expression, timing, and vertical stratigraphic extent (Fig. 9). Furthermore, local post-depositional processes (e.g., 575 576 weathering and burial diagenesis) may lead to the loss of Hg sequestered in rocks, altering the Hg record of these successions (Charbonnier et al., 2020). As such, Hg loss due to post-577 depositional alteration may partly explain the locally variable expression of Hg anomalies 578 observed for many PTB successions (Fig. 9). However, a thorough assessment of the degree of 579 preservation of rock successions from which P-T transition Hg records have been published 580 would be required to fully explore this possibility. Therefore, the variability of PTB Hg records 581 582 across different localities questions the reliability of these Hg anomalies for positioning the PTBME and the use of Hg anomalies as a stratigraphic correlation tool. 583

In the case of the Pingtang syncline record, neither Hg excursion (E1 nor E2) temporally 584 corresponds to the extinction interval as calibrated in the Meishan GSSP (Fig. 3; Fig. 8). Also, no 585 Hg anomaly is recorded at the PTBME extinction interval in Laxian (i.e., between LAX 8T and 586 10T, Fig. 3), although the equivalent stratigraphical interval in Potuo is a visibility gap 587 (Bagherpour et al., 2020). As such, the presence of a Hg anomaly there cannot be formally 588 excluded. Nevertheless, our U-Pb zircon ages show that both the nadir of the PTB negative δ^{13} C 589 excursion and the peak of the stratigraphically nearest Hg anomaly to the PTB (E2) are of 590 Griesbachian age (between 251.589 ± 0.052 Ma and 251.668 ± 0.079 Ma). The peak of these 591 excursions therefore, correlate with Bed 33 of the Meishan GSSP (251.583 ± 0.086 Ma, Burgess 592

et al., 2014), which is much younger than the PTB. Thus, these data provide evidence that

- although Hg anomalies may coincide with the nadir of the PTB negative δ^{13} C excursion (Fig. 9),
- this correspondence cannot be reliably used as a stratigraphic marker for the PTBME in a single section or between different sections (and/or different basins). Similar conclusions were reached
- for the end-Triassic extinction (ETE) event by Yager et al. (2021), who documented
- ⁵⁹⁸ "mismatches in timing" between Triassic-Jurassic boundary Hg anomalies and Central Atlantic
- 599 Magmatic Province (CAMP) magmatism, which is purported to have triggered the ETE event.
- 600 Hence, positioning the PTB extinction event by means of and/or correlation based on
- 601 Hgchemostratigraphy should be treated with extreme caution.
- The δ^{13} C record from Tethyan marine successions (Fig. 9) indicates that the expression of 602 the PTB δ^{13} C excursion varies between different sections (as previously alluded to by S. Shen et 603 al., 2019). Nevertheless, the PTB (as determined for the individual sections), appears to 604 approximately correspond to the midpoint, not the nadir, of the negative δ^{13} C excursion in 605 successions not affected by stratigraphic condensation (e.g., Meishan) or a PTB hiatus (Fig. 9). 606 Therefore, it is suggested that in the absence of high-resolution U-Pb zircon age calibration, an 607 approximate correlation of the PTB extinction interval could be achieved using the midpoint of 608 the PTB negative δ^{13} C excursion. This correlation strategy could be most effective for sections 609 lacking robust biochronology and where the P-T transition δ^{13} C record is not truncated by a 610 hiatus. 611
- 612

4.3 Does the E2 Hg anomaly of Griesbachian age coincide with a 2nd extinction event?

- The onset of the Griesbachian Hg anomaly as recorded from the Laxian section (Fig. 3) postdates LAX 10T, which is dated at 251.822 ± 0.060 Ma. This age coincides, within analytical uncertainty, with that determined for Bed 28 at the Meishan GSSP (251.880 ± 0.031 Ma; Burgess et al., 2014). Because a second step of the PTB mass extinction has been postulated to have occurred within Bed 28 in Meishan (Song et al., 2013), it is pertinent to consider whether the E2 Hg anomaly is associated with this proposed second extinction step.
- Conodont biozones around the PTB are usually interval zones (IZs), with the base of each 619 IZ being defined by the first occurrence (FO) of an index species and the top defined by the base 620 of the next overlying IZ. However, this biostratigraphical correlation technique often leads to 621 diachronous correlations because the relative stratigraphic order of FOs of index species is not 622 constant across space, as demonstrated by Brosse et al. (2016) for conodonts around the PTB in 623 South China (see also Ellwood et al., 2017). Reasons for diachronous IZs include ecological 624 control over the distribution of species in time and space, sampling effort, selective preservation, 625 and hiatuses in the sedimentary record (Guex, 1991; Holland & Patzkowsky, 2015; Leu et al., 626 2022). However, following this approach, the "legal" base Triassic was defined by the FO of 627 Hindeodus parvus in the very condensed Meishan section (Yin et al., 2001). 628
- Song et al. (2013) compiled FOs and LOs (last local occurrences) for conodonts and 629 benthic taxa from seven South Chinese PTB sections (including Meishan and Shangsi), which 630 led the authors to propose two extinction steps. The main and older event was placed at the base 631 of the C. meishanensis IZ, the antepenultimate IZ below the FO of H. parvus. The proposed 632 second extinction step, of lesser magnitude, was found at the base of the I. isarcica IZ, which is 633 the third Triassic IZ above the spatially variable FO of *H. parvus*. However, a recent thorough 634 635 re-investigation of Late Permian conodont IZs in the more expanded Shangsi section (Yuan et al., 2019) led to a revision of the basal Triassic into the C. meishanensis IZ. This new placement 636
- 637 of the base of the Triassic (Yuan et al. 2019; written comm. 2022) in the *C. meishanensis* IZ in

638 Shangsi and Meishan has the intrinsic benefit of coinciding with both the main extinction event

- and the lithostratigraphic boundary between Permian and Triassic rock units, which is marked by
- a hiatus in outer shelves and shallower depositional settings in South China and elsewhere
- (Bagherpour et al., 2017; Yin et al., 2014). Close examination of the raw biostratigraphic data
- (Table S2 of Song et al. 2013) from which a second extinction step was postulated shows no
 consistent extinction in the relative timing of the different species across the data set. Extinction
- 643 consistent extinction in the relative timing of the different species across the data set. Extinction 644 of a given clade occurs in a single section, or pair of sections at the very best, thus undermining
- arguments in favor of a second extinction event of global significance. For instance, only the
- 646 condensed Meishan section displays an apparent second step for benthic foraminifers and
- bivalves. In Shangsi, no second extinction emerges for any benthic clade around the base of *the*
- 648 *I. isarcica* IZ.649 Furthermore, the second second
- Furthermore, the compositing of local extinctions patterns (Song et al. 2013) relies on the implicit assumption of synchronous conodont IZs. However, as the seven sections of this data set 650 cover a very broad range of water depths, ranging from lagoon-shoals to lower slope-basin 651 (Table S1 of Song et al. 2013,), the assumption of synchronous conodont IZs becomes untenable. 652 It is also at variance with the fact that the respective bathymetric distribution of segminate and 653 segminiplanate conodonts, both involved as index species of IZs, is known to be spatially 654 controlled by temperature - i.e., depth of water masses (Joachimski et al., 2012; Leu et al., 655 2019). Moreover, the general development of a hiatus spanning approximately the entire C. 656 657 meishanensis IZ in shallower depositional settings as clearly established by Yin et al. (2014) automatically excludes any synchronicity of the neighboring IZ when comparing to hiatus-free 658
- 659 deep-water sections.

660 Consequently, because (i) the presence of a stratigraphic hiatus in some sections will 661 inevitably generate a spurious extinction event and (ii) conodont IZs across a depth gradient 662 ranging from lagoonal to basinal depositional settings can hardly be synchronous, there is 663 insufficient biostratigraphic evidence to support a second extinction event in South China. 664 Consequently, it is concluded that the E2 Griesbachian Hg anomaly reported here is not

associated with a second extinction pulse in South China.

666 4.4 Provenance of volcanic Hg input

Our U-Pb geochronological results allow us to place the Pingtang syncline Hg record 667 within the temporal framework of Siberian Traps Large Igneous Province magmatism (Burgess 668 & Bowring, 2015; Burgess et al., 2017), a prominently discussed source of volatiles and toxic 669 elements (such as Hg) during the P-T transition (Black et al., 2012; Broadley et al., 2018; Sibik 670 et al., 2021; Svensen et al., 2018). A maximum duration for intrusive and extrusive magmatic 671 STLIP activity is given by the bracketing ages of 252.27 ± 0.11 Ma (Burgess & Bowring, 2015) 672 to 250.60 ± 0.22 Ma (Augland et al., 2019), the latter from syenitic intrusions with an uncertain 673 relationship to the STLIP. Therefore, STLIP magmatism, being active during the studied 674 interval, is a potential source of volcanic Hg input to South China during the P-T transition. 675 However, several studies have demonstrated that more proximal regional volcanic 676 activity related to convergent plate tectonism and subduction magmatism occurred during the P-677 T transition, which led to elevated Hg concentrations and may have contributed to the PTBME in 678 South China (Gao et al., 2013; He et al., 2014; Jiao et al., 2022; H. Zhang et al., 2021; Zhao et 679 al., 2019; Zheng et al., 2020). In addition to Hg/TOC anomalies close to the PTBME horizon, 680 Hg/TOC excursions and corresponding Hg isotope compositions have been documented for the 681

earliest Triassic (Griesbachian) from South China and northern India (Wang et al., 2019a, 2018).

Wang et al. (2019a) documented a pair of Hg/TOC anomalies in the latest Permian and earliest 683 Triassic respectively, accompanied by a decreasing trend of Δ^{199} Hg values from the 684 Changhsingian to the Griesbachian. Based on this Δ^{199} Hg trend, they interpreted their 685 Griesbachian Hg/TOC excursion as having resulted from terrestrial Hg input due to elevated 686 Early Triassic continental weathering. Recently, J. Shen et al. (2021) reported several Hg 687 enrichment intervals predating the PTBME (named ME1-ME3), as well as one interval coeval 688 with the PTBME (ME4), from three marine sections in South China. These authors reported the 689 same pattern of decreasing Δ^{199} Hg values from the Changhsingian to Griesbachian. The Hg 690 anomalies preceding the PTBME were interpreted to reflect regional subduction-related 691 volcanism due to their spatial restriction to the Tethys region, the occurrence of numerous 692 volcanic ash layers in Upper Permian rocks across South China, and the geochemical 693 composition of zircons in these ash layers supporting a subduction-zone volcanic arc origin. 694

In contrast with previous studies, the Δ^{199} Hg values in the current study remain near-zero 695 throughout the studied interval, and are slightly positive within the Griesbachian E2 anomaly 696 (Fig. 8). This Δ^{199} Hg trend, despite variations in OM content and detrital flux, is consistent with 697 a constant, dominantly atmospheric volcanic Hg source relatively close to the Nanpanjiang Basin 698 during the studied interval. The disparity between the Hg isotope MIF record of the Pingtang 699 syncline and other deep-water marine records in South China (e.g., Wang et al., 2019a; J. Shen et 700 al., 2021) (Fig. 9), however, suggests that: 1) Hg sequestered in PTB-straddling marine 701 702 successions in different parts of South China do not have the same source/depositional pathway, and 2) Hg isotope compositions alone may not be sufficient for discriminating between different 703 potential volcanic sources of Hg (e.g., Siberian Traps volcanism or regional arc volcanism) to 704 deep-water depositional sites during the P-T transition in South China. 705

Given that i) Hg excursions in the Pingtang syncline successions are recorded in strata 706 with numerous interbedded volcanic ash layers, ii) the Δ^{199} Hg values of these ashes are 707 indistinguishable from those of interbedded rocks, and iii) the Nanpanjiang Basin was situated 708 close to a volcanic center during the P-T transition, we postulate that the major and trace element 709 geochemical properties of these volcanic ashes can be used to trace the origin of volcanic Hg 710 inputs to these successions. The analyzed volcanic ash beds from Potuo have major and trace 711 element characteristics similar to previously studied volcanic ashes from South China (Fig. 6) 712 (He et al., 2014; Wang et al., 2019b). Their primitive mantle-normalized trace element 713 714 compositions (Sun & McDonough, 1989) are characterized by depletions in Ta, Nb, Sr and Ti, similar to rocks from subduction settings (Pearce et al., 1995). The ashes show intermediate to 715 acidic chemical compositions, plotting in the field of basaltic andesite, trachy-andesite and 716 rhyolite/dacite (Fig. 6a), and are chemically distinct from volcanic rocks originating from the 717 STLIP (Callegaro et al., 2021; Reichow et al., 2005; Sibik et al., 2015; Sobolev et al., 2009). In 718 addition, a mid-oceanic ridge basalt (MORB)-normalized trace element (Th vs. Nb) discriminant 719 plot (Saccani et al., 2015, 2018) of the volcanic ashes from the Pingtang syncline suggests that 720 721 they originate from a continental margin volcanic arc tectonic setting (Fig. 6b). Similar conclusions were reached for other volcanic ashes from successions straddling the PTB in South 722 China, in that these ashes have no genetic link to the Siberian Traps, but instead derived from 723 subduction zone arc volcanism in the Tethys region (Gao et al., 2013; He et al., 2014; Jiao et al., 724 2022; Song et al., 2022; Zhao et al., 2019). 725 Consequently, we conclude that episodic regional arc volcanism associated with 726

convergent plate tectonics in the Tethys region best explains the elevated Hg input to the
 Nanpanjiang Basin as recorded in the Pingtang syncline during Changhsingian to Griesbachian

times. Furthermore, based on general atmospheric circulation models, previous studies have

- suggested that the Canadian Arctic was favorably positioned to receive volatiles (including
- volcanic ash) from the STLIP during the Permian to Triassic because it was situated downwind
 relative to the location of the STLIP eruptions (Dal Corso et al., 2022; Grasby et al., 2011, 2013).
- In this scenario, the South China region, being located towards the southeast and several
- thousands of kilometers away from Siberia, would have been poorly situated to receive volatiles
- from the STLIP. Nevertheless, STLIP Hg contributions to South China cannot be ruled out.
- Finally, as noted by J. Shen et al. (2023), volcanic arc magmatism along the eastern margin of
- the PaleoTethys was especially active during the Permian to Triassic, as inferred from abundant
- volcanic ash beds in coeval strata, the distribution of volcanic rocks, as well as a number of
- associated Hg anomalies across the PTB in this region (Fig. 9).
- 740

4.5 Hg anomalies, δ^{13} C excursions and volcanism during the P-T transition

741 Owing to the paucity of U-Pb zircon ages for deep-water sections from which Hg anomalies have been reported in South China, it is difficult to confidently correlate our recorded 742 743 Hg anomalies with those reported for other localities in the Tethys region. However, the E1 anomaly recorded from the studied successions together with ME2 and ME3 episodes of J. Shen 744 et al. (2021), indicate that Hg anomalies preceding the PTB extinction are recorded in both 745 shallow and deep-marine settings in South China. In contrast, the E2 Griesbachian Hg anomaly 746 is not recorded from any shallow-water marine section nor deep-water sections in South China, 747 except for deep-water sections in the Nanpanjiang Basin (e.g., Xinmin, Kejiao; Fig. 9). 748 749 Nevertheless, E2 may be coeval with the Hg anomaly recorded between the *I. staeschei* and *I.* isarcica conodont zones (Wang et al., 2019a), which also coincides with the nadir of the PTB-750 straddling negative δ^{13} C excursion in the southwestern Tethys Guryul Ravine section (Fig. 9). 751 However, further work from other marine successions with precise U-Pb zircon ages is required 752 to confirm the spatial extent of the Griesbachian Hg anomaly. 753

The (stratigraphically variable) coincidence of the nadir of δ^{13} C excursions with Hg 754 anomalies in the Pingtang syncline and other Tethyan successions (Fig. 9) hints at a common 755 driving factor – volcanism. The global distribution of $\delta^{13}C$ excursions and Hg anomalies close to 756 the PTB in Permian-Triassic successions (Baud et al., 1996; Grasby et al., 2013; Korte & Kozur, 757 2010; Sanei et al., 2012; J. Shen et al., 2019) argues in favor of a large-scale volcanic degassing 758 episode, such as that of the STLIP, as the ultimate cause of drastic environmental changes of 759 global extent. The onset of the negative δ^{13} C excursion in the Pingtang syncline starts just before 760 deposition of the ash layer (LAX8T) at 252.07 ± 0.130 Ma (Fig. 8) and is characterized by an 761 initial 2 ‰ decrease in δ^{13} C values in the upper Changhsingian followed by a further 3 ‰ 762 decrease in the lower Griesbachian. This onset of δ^{13} C excursion may temporally overlap with 763 the transition from the extrusive Stage 1 to intrusive Stage 2 of STLIP magmatism, proposed to 764 have occurred at about 251.9 Ma (Burgess et al., 2017). Stage 2 STLIP magmatism was 765 characterized by intrusion of dykes and sills into country rocks in the Tunguska Basin (Burgess 766 & Bowring, 2015; Burgess et al., 2017). The intrusions, and subsequent sill complex formation, 767 are proposed to have caused injection of massive amounts of both CO₂ and CH₄ into the 768 atmosphere via contact metamorphism of coal beds, shales and petroleum-bearing evaporites in 769 the Tunguska Basin (Burgess et al., 2017; Svensen et al., 2009, 2018), which collectively are 770 thought to result in a > 5 ‰ negative $\delta^{13}C$ excursion. 771

However, Davydov (2021) recently questioned the validity of this sill-rock thermal
 interaction model, pointing out that there is no correlation between coal metamorphism and the

distribution of sills in the Tunguska Basin. This author also argued that the role of contact 774 775 metamorphism by undifferentiated intrusions, which constitute > 95 % of the Tunguska Basin intrusions, was limited and insignificant to the general coal metamorphism in the Tunguska 776 777 Basin; and that coal metamorphism in the Tunguska Basin was probably related to regional tectonic deformation instead of magmatism. Furthermore, the role of intrusive STLIP 778 magmatism as a potential driver of the global carbon cycle perturbations and mass extinction 779 during the P-T transition is questioned. This stems from the uncertainty regarding the age of the 780 explosion pipes in the Tunguska Basin (which could have transported CO₂ and other gases such 781 as Hg to the atmosphere), as well as the U-Pb ages of the sill intrusions that violate stratigraphic 782 superposition (Davydov, 2021). Consequently, additional U-Pb zircon geochronologic 783 calibrations of STLIP intrusive rocks are required to resolve these questions. 784

Pending the resolution of these questions, an additional but not mutually exclusive 785 explanation that could reconcile the coincidence of negative δ^{13} C excursions and Hg anomalies 786 recorded at a global scale during the P-T transition can be considered. This is that concurrent 787 regional arc volcanism in different palaeocontinents was responsible for both CO₂ and Hg 788 release resulting in the δ^{13} C excursions and Hg anomalies. In addition to South China, extensive 789 regional arc volcanism during the P-T transition has been documented for many spatially 790 disparate localities. Some of these include: the northern Patagonian Massif, Argentina (Luppo et 791 al., 2018), Antarctica (Nelson & Cottle, 2019), Sydney Basin, Australia (Metcalfe et al., 2015), 792 793 Karoo Basin, South Africa (Gastaldo et al., 2020), with coeval Hg anomalies also recently documented for the latter two (J. Shen et al., 2023). The suggestion of concurrent regional arc 794 volcanism in several paleocontinents during the P-T transition is congruent with recent 795 suggestions of a Pangean 'ring of fire' (subduction-related volcanism along the convergent 796 Panthalassan margin of Pangea) as an explanation for marine and terrestrial environmental 797 perturbations during the latest Permian (Vajda et al., 2020). 798

Reasons for a global increase in regional arc volcanism concurrent with STLIP volcanism 799 are still uncertain. Jiao et al. (2022) recently suggested that the ascent of the STLIP mantle plume 800 may have triggered volcanic activity in several subduction zone systems. While the effect of a 801 large mantle plume on global plate tectonics is highly debatable, STLIP volcanism occurred in a 802 context of global increase of accretion and subduction rates from the Permian to Triassic (Vérard 803 et al. 2015a, their Fig. 11a), which has also been linked with Early Triassic sea-level rise (Vérard 804 et al., 2015b, their Fig.17). However, such global increase in subduction rates is likely to occur 805 over tens of millions of years. Hence, increased regional arc volcanism over hundreds of 806 thousands of years between the latest Permian to earliest Triassic may be totally disconnected 807 from global tectonics. Irrespective of the reasons for this global increase in arc volcanism over 808 the P-T transition, our results from the Pingtang syncline suggest that volcanic activity linked to 809 Hg anomalies and C-isotope excursion probably peaked between 251.589 ± 0.052 Ma and 810

811 251.668 ± 0.079 Ma in the Nanpanjiang Basin, South China (Fig. 8).

4.6 Implications for the PTBME

While it is generally accepted that STLIP magmatism exerted a major control on the global carbon budget and mercury cycle during the Paleozoic to Mesozoic transition, it is clear that South China was substantially influenced by nearby volcanic centers throughout the Late Permian to Triassic (He et al., 2014; H. Zhang et al., 2021). This is evident from the older and well-known Emeishan LIP (Huang et al., 2022), the occurrence of Permian-Triassic volcanic rocks especially in southwest South China (Gao et al., 2013), the abundant volcanic ash layers

- 819 within Permian-Triassic marine sedimentary successions, shown to be genetically distinct from
- STLIP rocks (Gao et al., 2013; He et al., 2014; Yang et al., 2012), as well as Changhsingian
- Hg/TOC anomalies restricted to the Tethys region (J. Shen et al., 2021; this study). In addition,
- regional arc volcanism has been linked to notably decreased carbonate and biogenic silica production, as well as decreased water column oxygenation in South China (J. Shen et al., 2013).
- production, as well as decreased water column oxygenation in South China (J. Shen et al., 20
 The temporal overlap between regional intermediate to felsic volcanism and the basaltic
- The temporal overlap between regional intermediate to felsic volcanism and the basaltic volcanism from the STLIP, thus supports a scenario in which STLIP magmatism and concurrent
- subduction-related regional arc volcanism in the Tethys region may have acted in concert to
- generate pernicious environmental conditions for marine and terrestrial faunas in South China
- during the Late Permian (J. Shen et al., 2013, 2021; S. Shen et al., 2019).
- The finding of several sudden increases in mercury concentration that both predate and post-date the PTBME event is significant. This is because these Hg enrichments suggest that elevated volcanic activity, which is thought to have triggered the PTBME in South China, was
- not restricted only to a short interval e.g., within Stage 2 STLIP magmatism (Burgess et al.,
- 2017), but rather, took place over several thousands of years during the P-T transition. Thus,
- repeated bursts of regional felsic and intermediate volcanism (and resulting increases in mercury
- concentration) could have had a cumulative adverse effect on the environment and the species
- that lived at the time, making them more vulnerable to extinction (J.Shen et al., 2013; 2021; S.
- 837 Shen et al., 2019). In this scenario, significant environmental stresses resulting from STLIP
- magmatism would have quite easily pushed faunas in South China "over the edge" to bring about
- the mass extinction at the Paleozoic-Mesozoic transition.

840 **5 Conclusions**

The present study establishes the Hg and δ^{13} C record of two deep-water marine 841 successions in the Nanpanjiang Basin, South China, spanning the Changhsingian to 842 Griesbachian. U-Pb zircon ages and Hg/TOC ratios indicate elevated volcanic activity before 843 252.07 ± 0.130 Ma (preceding the PTBME) and during several episodes in the Griesbachian, 844 peaking between 251.589 ± 0.052 Ma and 251.668 ± 0.079 Ma. The latter range in ages is coeval 845 with the nadir of a large negative C isotope excursion, similar in magnitude to the global 846 excursion at the PTB. Based on evidence from the coupled δ^{13} C and Hg records, major and trace 847 element geochemistry of volcanic ashes, and the U-Pb zircon age constraints, we conclude that 848 the recorded Hg enrichments are primarily sourced from subduction-related arc volcanism in the 849 Tethys region. Consequently, our Hg record together with recent results from South China 850 documenting several episodes of Hg enrichment during the P-T transition, are compatible with 851 previous suggestions that arc volcanism contributed to environmental deterioration and 852 biodiversity decline in South China leading up to the PTBME, which was triggered by STLIP 853 magmatism. Our study provides an absolute timeframe within which Late Permian to Early 854 Triassic Hg and δ^{13} C records from different deep-water successions (which are less likely to 855 contain stratigraphic gaps) in South China can be calibrated. This precise, radioisotopic 856 timeframe allows for a more in-depth assessment of the relation between volcanism, 857 environmental changes, and the mass extinction event at the P-T transition. 858

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867 **Conflict of interest**

868 The authors declare no conflict of interest relevant to this study.

869 Author contributions

- 870 Conceptualization & Funding acquisition: Hugo Bucher, Urs Schaltegger, Torsten Vennemann
- 871 Formal analysis: Oluwaseun Edward, André N. Paul, Christian Vérard
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876 Data availability Statement

- 877 The dataset associated with the current study is available on the open-source online data
- repository Zenodo at "[*link-to-be-given-upon paper-acceptance*]". Figure 1 was made based
- on the Panalesis model (Vérard, 2019) using ArcGIS®. Figures 2-8 were made with Matplotlib
- version 3.5.2 (Caswell et al., 2022), available under the Matplotlib license at
- 881 <u>https://matplotlib.org</u>, and seaborn version 0.11.2 (Waskom, 2021) available at
- 882 <u>https://pypi.org/project/seaborn/</u>. All figures were edited using Adobe Illustrator©.

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1294 Figure Captions

Figure 1. Maps showing the location of the studied successions, as well as other marine 1295 1296 successions in the Tethys region from which Permian-Triassic (P-T) transition mercury records have been published. a) Global paleogeographic map at the P-T transition after the Panalesis 1297 model (Vérard, 2019). Locality marked '10' represents the paleo-location of Guryul Ravine, 1298 1299 northern India. b) Map showing the present-day location of the studied sections (red circles) and 1300 other deep-water marine sections in South China (yellow circles); white circles show the main cities. The sections are represented by numbers as follows: 1 – Meishan, 2 – Majiashan, 3 – 1301 Daxiakou, 4 - Xiakou. 5 - Shangsi, 6 - Laxian, 7 - Potuo, 8 - Xinmin, 9 - Kejiao. c) Detailed 1302 1303 map of the South China area (orthogonal projection) with the paleo-location of the discussed sections. 1304

Figure 2. Single-grain zircon analyses and ²⁰⁶Pb/²³⁸U weighted mean dates for volcanic ash beds from Potuo and Laxian. Bars represent single grain zircon U-Pb dates and their uncertainty. Transparent bars are rejected for interpretation (Pb-loss, inheritance, antecrystic).

Figure 3. Carbon isotope values, Hg concentrations, Hg/TOC ratios, TOC contents and 1308 Δ^{199} Hg values from a) Laxian, b) Potuo. The gap between the Talung and Daye formations in the 1309 Potuo stratigraphic log is an observation gap due to coverage by alluvium (Bagherpour et al., 1310 2020). Black dashed lines represent the 0.2 wt.% TOC limit for Hg/TOC normalization (Grasby 1311 et al., 2019) and green dashed lines represent background $\delta^{13}C_{org}$ values (-27 ‰). Box and 1312 whisker plots of c) Δ^{199} Hg values and d) Δ^{200} Hg values for Potuo and Laxian compared to those 1313 1314 previously documented for other deep-water marine sections in South China. Grey shaded rectangle (in a, b) and red dashed lines (in c, d) represent the interval of no measurable Hg 1315 1316 isotope mass independent fractionation (MIF, 0 ± 0.05 ‰). Data sources are as follows: 1317 Majiashan - Wang et al. (2019a), Daxiakou - Wang et al. (2018), Xiakou - J. Shen et al. (2019), Shangsi - J. Shen et al. (2021). 1318

Figure 4. Scatter plot showing the relationship between Hg and a) Al, b) U/Al, c) TOC, d) Mo/Al and e) Fe in the Pingtang syncline sections. Black dashed line represents 0.2 wt.% TOC limit (Grasby et al., 2016). Volcanic ash samples are not included in the calculation of the correlation coefficient squared (r²).

1323 Figure 5. Composite plot of Hg/element ratios and Hg isotope mass dependent fractionation values for Potuo and Laxian. a) Al, b) Hg/Al, c) Hg/Fe, d) Hg/TOC, e) Mo/Al, f) 1324 U/Al, g) δ^{202} Hg. Symbols of lithological log are identical to those of Fig. 3. The composite 1325 section is based on lithological comparison of the Potuo and Laxian sections (Bagherpour et al., 1326 2020) and assumes similar sedimentation rates for both sections. Note that the Potuo and Laxian 1327 sections are only 6.4 km apart and both belong to the Pingtang syncline of the Nanpajiang Basin, 1328 1329 recording laterally continuous basinal facies belonging to the Talung Fm. and Daye Fm. 1330 (Bagherpour et al., 2020).

Figure 6: Major and trace element geochemistry results for volcanic ash samples from the Pingtang syncline. A) Zr/Ti vs Nb/Y classification diagram (Winchester & Floyd, 1977) for volcanic ashes from South China plotted along with volcanic rocks from the Siberian Traps Large Igneous Province (STLIP) b) N-MORB-normalized Th vs. Nb discriminant plot, with tectonic setting interpretation after Saccani et al. (2015, 2018) for volcanic ashes from South

- 1336 China, together with data from dolerite sills, dykes and basalts from the STLIP (Tunguska Basin)
- 1337 plotted for comparison. N-MORB: normal-type mid-oceanic ridge basalt. C) Ti vs Zr
- 1338 classification diagram after Pearce (1982). The field illustrating typical Siberian Traps volcanic
- rock compositions is after He et al. (2014). D) Primitive mantle-normalized spider diagram for Changhsingian and Griesbachian volcanic ashes from Potuo and other localities (Meishan,
- Changhsingian and Griesbachian volcanic ashes from Potuo and other localities (Meishan,
 Chaotian, Jianshi, Rencunping, Shangsi and Dongpan) in South China. Data for South China
- localities other than Potuo are from He et al. (2014), Song et al. (2022) and Wang et al. (2019b).
- Normalization values for N-MORB and primitive mantle are from Sun & McDonough (1989).
- 1344 Data for volcanic rocks from the STLIP are from several sources as follows: basalt flows,
- dolerite sills, dykes: Sibik et al. (2015); Callegaro et al. (2021); magnesian rocks, melt
- 1346 inclusions: Sobolev et al. (2009); West Siberia Basin (WSB) basalts: Reichow et al. (2005).
- 1347 WSB: West Siberia Basin, SC: South China.
- 1348Figure 7. Cross plots of Rock-Eval TOC data for samples with TOC > 0.2 wt.% to assess1349the type and quality of preserved organic matter in Potuo and Laxian (modified after1350Charbonnier et al., 2020). A) HI vs OI, b) HI vs T_{max}, c) HI vs TOC, d) Hg vs Hg.
- Figure 8. Composite profiles for a) δ^{13} C, b) Hg concentration, c) Hg/TOC ratios, d) 1351 Δ^{199} Hg, e) Δ^{200} Hg records from the Potuo and Laxian sections. δ^{13} C values are from Bagherpour 1352 et al. (2020). The horizontal gray band marks the interval of Hg concentration spike (E2) and 1353 nadir of the negative δ^{13} C excursion and the vertical pink bands depict near-zero Hg isotope MIF 1354 (0 ± 0.05) %, Thibodeau et al., 2016). The dashed green line represents the conformable Talung-1355 1356 Daye formational boundary and the PTB. The Siberian Traps magmatism timeline is after Burgess et al. (2017) and the color gradient between the stages depicts the uncertainty in the 1357 timing of the transition between different stages. The PTB on the composite log is defined from 1358 1359 the Laxian section as this interval at Potuo is covered by recent alluvial deposits (Bagherpour et al., 2020). U-Pb zircon ages for the PTB (in green) are from (a) Burgess et al. (2014) and (b) 1360 Baresel et al. (2017b). FO – first occurrence. 1361
- Figure 9. Compilation of carbon (organic and carbonate) isotope, mercury (Hg) 1362 concentration, Hg/TOC and Hg isotope (Δ^{199} Hg) records for marine depositional environments 1363 in the Tethys region: South China and northern India. Hg/TOC ratios are shown only for 1364 localities and stratigraphic intervals where TOC content is > 0.2 wt.%. a)Shangsi, b) Xiakou, c) 1365 1366 Majiashan, d) Daxiakou, e)Xinmin, f) Kejiao, g) Pingtang syncline (composite of the Potuo and Laxian sections), h) Meishan. U-Pb zircon ages are from Burgess et al. (2014), i) Guryul Ravine, 1367 northern India. The different profiles are correlated using the Permian-Triassic boundary as 1368 placed by the authors of the original data sources. Data sources are as follows: Meishan: Cao et 1369 1370 al. (2002); Grasby et al. (2017) and J. Shen et al. (2019); Guryul Ravine and Majiashan: Wang et al. (2019a); Shangsi: J. Shen et al. (2021); Xiakou: J. Shen et al. (2019); G.J. Zhang et al. (2021); 1371 1372 Xinmin, Kejiao: J. Shen et al. (2019); Daxiakou: Wang et al. (2018); Pingtang syncline (Potuo, Laxian): this study. U-Pb zircon age for the Permian-Triassic boundary (in green) is from 1373 Burgess et al. (2014). Abbreviations: Gries.: Griesbachian; *: Hindeodus latidentatus; C.m.: 1374 Clarkina meishanensis; **: Hindeodus parvus; H.p: Hindeodus praeparvus; I.st.: Isarcicella 1375 staeschei; Nc.krv.: Neoclarkina krvstvni; C.v..: Clarkina vini; C.tav.: Clarkina tavlorae. 1376

Figure 1.



Figure 2.



Figure 3.



Figure 4.



Figure 5.



Figure 6.



Figure 7.



Figure 8.



Figure 9.

