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27 **Abstract**

28 The radiation of early animals during the Cambrian explosion was accompanied by
29 widespread perturbations in oceanic and atmospheric conditions. However, the cause-effect
30 relationships between evolutionary innovations, carbon (C) cycling, and continental
31 weathering are still a matter of debate. Further paleoenvironmental reconstructions of poorly
32 studied paleo-continents covering the Precambrian/Cambrian (PC/C) transition may improve
33 the correlation of the events and their impact on the Earth system on a global scale. The Tarim
34 Basin was presumably located proximal to the well-studied South China craton during the
35 Ediacaran-Cambrian boundary and is expected to contribute equally to our
36 paleoenvironmental reconstructions. Here we apply carbon and strontium (Sr) isotope records
37 ($\delta^{13}\text{C}$ and $^{87}\text{Sr}/^{86}\text{Sr}$) obtained from well-preserved marine carbonates spanning the late
38 Ediacaran to the late Cambrian from the Keping region in the northwestern Tarim Basin, NW
39 China. By comparison to other carbonate successions from various paleo-continents, our
40 obtained seawater $^{87}\text{Sr}/^{86}\text{Sr}$ curve in this study shows a stepwise first-order increase to more
41 radiogenic values superimposed by second-order presumably regional variations. Our
42 obtained $\delta^{13}\text{C}$ variations exhibit four prominent negative and three positive excursions
43 identical to the global $\delta^{13}\text{C}$ record. Based on existing paleontological evidence and our new
44 Tarim $\delta^{13}\text{C}$ and $^{87}\text{Sr}/^{86}\text{Sr}$ curves, we propose a stratigraphic correlation from the late Ediacaran
45 to the late Cambrian between the Tarim Basin and other paleo-continents. Besides, our
46 paleoenvironmental reconstruction reveals both regional and global sea-level changes that
47 may have controlled the influx of radiogenic Sr and presumably the influx of nutrients
48 controlling primary production and the rate of organic carbon burial in the Tarim Basin at the
49 time of deposition. These feedbacks may have ultimately influenced the rapid diversification
50 of metazoans and the oxygenation of the Cambrian ocean and atmosphere approaching-
51 present $p\text{O}_2$ levels.

52 **Keywords:** Carbon isotopes, Strontium isotopes, Ediacaran-Cambrian, Tarim Basin,
53 Paleoenvironment

54

55 **1. Introduction**

56

57 The transitional interval between the terminal Proterozoic and the early Paleozoic is
58 marked by the sudden appearance of abundant and diverse metazoans (Valentine et al., 1999;
59 Conway-Morris, 2003; Maloof et al., 2010; Kouchinsky et al., 2011). This early animal evolution
60 was accompanied by extensive perturbations to climate, carbon (C) cycling, continental
61 weathering and changes to the composition of the oceans (Shields-Zhou and Zhu, 2013).
62 However, the cause-effect relationships between this evolutionary innovation and these
63 environmental perturbations are still a matter of broad scientific debate and
64 paleoenvironmental reconstructions of less studied paleo-continents covering the
65 Precambrian/Cambrian (PC/C) transition can help this situation. Most studies from China
66 covering this interval focus on sedimentary successions in South China. For instance, Li et al.
67 (2013) suggested that muted continental weathering during the early Cambrian was indicated
68 by decreasing seawater strontium isotope ratios ($^{87}\text{Sr}/^{86}\text{Sr}$). Ediacaran-Cambrian marine
69 carbonate successions outcropping in the Tarim Basin were deposited on a shallow-water
70 continental shelf, which was presumably situated close to the South China craton and at a
71 similar low-middle latitude position (Fig. 1A) (McKerrow et al., 1992; Merdith et al., 2017;
72 Huang et al., 2000; Zhan et al., 2007). In the past, Ediacaran-Cambrian Tarim Basin sediment
73 successions have only received minor attention (Zhou et al., 2018; Zhu et al., 2019) but a few
74 recent chemo-stratigraphic studies provided some preliminary results on their stratigraphic
75 subdivision and paleoenvironmental conditions. He et al. (2007) and Guo et al. (2017) provided
76 the first C isotope stratigraphic correlations for the late-Ediacaran Cambrian strata in the Tarim
77 Basin between the Sugetbrak, the Penglaiba, the Wushi phosphorite and the Dongergou
78 sections, while Wang et al. (2011b) presented paired C and Sr isotope trends from the
79 Cambrian Series 2 to Series 3 at the Penglaiba section. However, there is still a chemo-
80 stratigraphic gap (especially in regard of the Sr isotope stratigraphy) spanning the late-
81 Ediacaran to the late-Cambrian time interval, which limits the stratigraphic correlation of the
82 Tarim Basin with other parts of the world.

83 C and Sr isotope compositions obtained from well-preserved (i.e. low diagenetic fluid flow
84 overprinted) ancient carbonate rocks may retain primary compositions of the

85 contemporaneous seawater and provide constraints on the isotope-based chemo-
86 stratigraphic correlation (DePaolo and Ingram, 1985; Montañez et al., 1996; Montañez and
87 Banner, 2000; Zhu et al., 2007; Wang et al., 2011b; McArthur et al., 2012; Li et al., 2013; Zhou
88 et al., 2018; Zhu et al., 2019). For instance, the first appearance datum of small shelly fossils
89 (SSFs) is globally accompanied by a negative $\delta^{13}\text{C}$ excursion and a declining trend in seawater
90 $^{87}\text{Sr}/^{86}\text{Sr}$ ratios (Cowie and Glaessner, 1975; Landing, 1989; Zhu et al., 2006; Li et al., 2013; Zhu
91 et al., 2019), although there are still debates about the exact boundary definition between the
92 Ediacaran and the Cambrian (Geyer and Landing, 2017).

93 Secular variations in seawater $^{87}\text{Sr}/^{86}\text{Sr}$ ratios have been widely applied to reflect
94 continental uplift, silicate weathering and variations in atmospheric carbon dioxide
95 concentration (Burke et al., 1982; Keto and Jacobsen, 1987; Kaufman et al., 1993; Derry et al.,
96 1994; Montañez et al., 1996; Denison et al., 1998; Montañez and Banner, 2000; Ebner et al.,
97 2001; Thomas et al., 2001; Melezhik et al., 2009; Sawaki et al., 2010). The $^{87}\text{Sr}/^{86}\text{Sr}$ ratio of
98 seawater is controlled by a balance between continental weathering-derived Sr, hydrothermal
99 Sr input and seafloor weathering (Palmer and Edmond, 1989). Continental source generally
100 yields more radiogenic $^{87}\text{Sr}/^{86}\text{Sr}$ ratios due to the more radiogenic ^{87}Sr resulting from the excess
101 decay of incompatible ^{87}Rb (rubidium) in evolved differentiated crustal rocks, while
102 hydrothermal and seafloor weathering origin Sr deplete in radiogenic ^{87}Sr and therefore low
103 $^{87}\text{Sr}/^{86}\text{Sr}$ ratios. Neglectable isotopic fractionation of the Rb/Sr system occurs during marine
104 authigenic carbonate formation (Banner and Kaufman, 1994), together with its long residence
105 time in the modern oceans, Sr is homogeneously distributed in the oceans and the seawater
106 $^{87}\text{Sr}/^{86}\text{Sr}$ ratio ideally represents a global signature (DePaolo and Ingram, 1985). Thus, well-
107 preserved marine carbonate minerals likely preserve the $^{87}\text{Sr}/^{86}\text{Sr}$ signature of the ambient
108 seawater during the carbonate deposition.

109 In this study, we present C and Sr isotope compositions obtained in carbonates from the
110 late Ediacaran to the late Cambrian at the Shiirik and Xiaoerbrak sections in the Keping region,
111 Tarim Basin, NW China. The main objective of this study is to 1) document the variations of
112 seawater C and Sr isotope compositions recorded in carbonates deposited in the Tarim Basin
113 covering the late Ediacaran-late Cambrian interval; 2) refine the stratigraphic correlation of the
114 sedimentary successions within the Tarim Basin; 3) interpret the data in the context of

115 reconstruction of continental weathering and sea-level changes. Our new interpretations
116 further aim to contribute to understanding of mechanisms linking environmental changes and
117 animal evolution during this critical period.

118

119 **2. Geological setting**

120

121 The Tarim Basin was presumably paleogeographically situated in a low to middle latitude
122 position between Australia and South China (see Fig. 1A) during the late Ediacaran to the late
123 Cambrian. Both Tarim and its adjacent South China craton developed widespread shallow
124 water carbonate platforms (McKerrow et al., 1992; Merdith et al., 2017; Huang et al., 2000;
125 Zhan et al., 2007). Our studied Keping region is located at the north-western Tarim Basin, NW
126 China (Fig. 1B) where the Ediacaran-Cambrian carbonate successions are outcropped at the
127 Xiaoerbrak and Shiirik sections (GPS locations: N 40° 55' 23", E 79° 53' 53" and N 40° 59' 12",
128 E 79° 59' 35" respectively).

129

130 **2. 1 Shiirik section**

131

132 The Shiirik section is located 30 km SW of Aksu city, NW Xinjiang Uygur Autonomous
133 Region (Fig. 1B). This section records an Ediacaran-Cambrian succession of more than 200 m
134 sedimentary rocks including the Ediacaran Sugetbrak and Chigebrak Formation (abbreviated
135 throughout the manuscript Fm.) as well as the Cambrian Yurtus Fm. The Ediacaran Chigebrak
136 and the overlying Cambrian Yurtus Fm. are separated by a disconformity between them (Fig.
137 2).

138 The Sugetbrak Fm. is mainly composed of reddish quartz sandstones interbedded with
139 mudstone and dolostone layers and is in conformable contact with the overlying Chigebrak Fm.
140 Rhythmic stratification, which reflects the influence of tidal currents, is widely developed in
141 the middle and upper part of the Sugetbrak Fm., suggesting a tidal flat depositional
142 environment (Deng et al., 2019). The Chigebrak Fm. contains thin-layered limestones
143 interbedded with brown sandstones at the bottom, overlain by medium-thick (10-50 cm)
144 layers of dolostones in the middle part and grey and yellowish medium-thick-layered

145 dolostones in the upper part (He et al., 2007; Wang et al., 2010).

146 The Chigebrak Fm. is in contact with the lower Cambrian Yurtus Fm. by a widely
147 distributed unconformity in the Tarim Block (Wu et al., 2018). In the strata just below the
148 unconformity, various features of karstification including breccias, cave-sediment infills,
149 solution vugs and pores, were observed in the uppermost layers of dolostones in the Chigebrak
150 Fm., indicating a possible erosional surface (He et al., 2018).

151 The lower Cambrian Yurtus Fm. mainly comprises black shales and phosphatic siliceous
152 rocks with barite concretions, syn-genetic paleo-weathering crusts and abundant benthic SSFs
153 (Qian and Xiao, 1984; Qian et al., 2000; He et al., 2007; Peng, 2009; Qian et al., 2009; Chen et
154 al., 2010; Yao et al., 2014).

155 In the Shiirik section, 49 carbonate samples were collected from the 156 m-thick
156 Ediacaran Chigebrak Fm. (n = 41) and the top part of the underlying Sugetbrak Fm. (n = 8),
157 respectively (Fig. 2).

158

159 **2. 2 Xiaoerbrak section**

160

161 The Xiaoerbrak section is located 50 km SW of Aksu city, NW Xinjiang Uygur Autonomous
162 Region and comprises more than 800 m sedimentary rocks subdivided in ascending
163 stratigraphic order into the Cambrian Yurtus, Xiaoerbrak, Wusonger, Shayilik, Awatagh and
164 Xiaqiulitagh Fm. (Fig. 1 & 2), which are stratigraphically continuous without any depositional
165 gaps.

166 The 21 m thick Yurtus Fm. is subdivided into two parts: the lower Yurtus Fm. consists of
167 black shales, phosphatic siliceous rocks with the occurrence of sponge spicules (Peng, 2009;
168 Chen et al., 2010) and rare SSFs (*Protohertzina Anabarica Missarzhevsky* and *Anabarites* (Zhou
169 and Chen, 1990; Zhou, 2001), Fig. 2), while the upper part comprises thin-layered (1-10 cm)
170 siliceous dolostones and muddy dolostones containing abundant SSFs (Qian and Xiao, 1984;
171 Zhou and Chen, 1990; Qian et al., 2000; Zhou, 2001; Qian et al., 2009; Chen et al., 2010), such
172 as *Paragloborilus Spinatus* (Qian and Xiao, 1984) and *Lapworthella* (Zhou and Chen, 1990). The
173 sedimentary facies of the Yurtus Fm. are described in the literature as deposited in a restricted
174 basin environment inferred from sedimentologic and geophysical studies as well as from

175 molybdenum isotope compositions analyzed in the Yurtus black shales (Bai et al., 2019; Yao et
176 al., 2014; Yao et al., 2017; Yu et al., 2009).

177 The conformably overlying Xiaoerbrak Fm. consists of 138 m-thick medium-thick layered
178 micritic dolostone, which hosts three trilobite assemblages (in ascending stratigraphic order:
179 *Shizhudiscus* Zone, *Metaredlichioides* or *Ushbaspis* Zone, *Kepingaspis-Tianshanacephalus*
180 Zone) (Wang et al., 1985; Zhou and Chen, 1990; Zhou, 2001; Peng, 2009; Fig. 2).

181 The 95 m-thick Wusonger Fm. is mainly composed of fine-grained micritic dolomites and
182 muddy dolostones (Sun et al., 2004; Yu et al., 2004; Wang et al., 2010). Fossils of *Paokannia*
183 considered to survive from the Cambrian late Stage 3 to the end of Cambrian Series 2 were
184 identified in this formation (Zhou, 2001; Peng, 2009, Fig. 2 orange arrow indicates fossil range).
185 Semi-restricted carbonate platforms were indicated from thin-section studies of the muddy
186 dolostone (Bai et al., 2019).

187 The overlying Shayilik Fm. contains the *Kunmingaspis-Chittidilla* biozone and comprises
188 micritic and fine-grained dolomites in the lower part, dolomite-bearing limestones in the
189 upper part, and is topped by algal microbialites (Zhou, 2001; Peng, 2009, Fig. 2 yellow arrow
190 indicates fossil range).

191 The Awatagh Fm. spans about 200 m and comprises anhydrite and anhydrite-bearing
192 dolostones at the bottom overlain by purple medium-thin (5-30 cm) bedded micritic dolomites
193 and fine-grained dolomites in the upper part. Bai et al. (2019) studied the sedimentary
194 characteristics of the Shayilik and Awatagh Fm. and proposed their sedimentary environments
195 experienced a transgression and changed from gentle slope environments to restricted
196 platforms. The overlying Xiaqiulitagh Fm. mainly consists of dolomites, intercalated with
197 stromatolitic dolomites (Wang et al., 2011b) and siliceous dolomites exposed in the middle
198 part of this Fm.

199 At the Xiaoerbrak section, 126 carbonate samples and three phosphorite samples were
200 collected (Fig. 2). Based on regional lithostratigraphic and biostratigraphic correlation as well
201 as sedimentary facies comparison, Shaiirik and Xiaoerbrak sections are integrated into one
202 single lithological succession in this study (Fig. 1B).

203

204 **3. Analytical methods**

205

206 Hand specimen free from visible alteration such as weathering crusts and visible calcite
207 veins were collected in the field and subsequently crushed into small pieces of ~0.5 cm
208 diameter. The best-preserved pieces (i.e. showing no microscopic calcite veins) were
209 handpicked and ground into fine powder in an agate mill (200 mesh) for further geochemical
210 analyses.

211 0.5 mg of sample powders were baked for ten hours at 70 °C and then reacted with
212 orthophosphoric acid at 70 °C for 1.5 hours to extract CO₂ (McCrea, 1950; Craig, 1953) for
213 subsequent measurement of C and oxygen (O) isotope compositions. C and O isotope
214 compositions were determined using a Gas-isotope ratio mass spectrometer (IRMS) of type
215 Finnigan Delta Plus XP at the State Key Laboratory for Mineral Deposits Research, Nanjing
216 University, China. Chinese carbonate reference materials (TTB-1 and TTB-2) were analyzed
217 along with our samples for isotopic calibration and to test for memory effects. C and O isotope
218 compositions were measured relative to analyses of an in-house standard and the obtained
219 data is here expressed relative to the international V-PDB (Vienna Pee Dee Belemnite)
220 reference material against which our in-house standard was calibrated. We use the δ notation
221 throughout the manuscript, which is the permille deviation of the C and O isotope
222 compositions from the V-PDB reference material. The analytic precision of our method was
223 determined on repeated analyses of the external reference materials and is in the order of
224 ± 0.15 ‰ for $\delta^{13}\text{C}$ and ± 0.1 ‰ for $\delta^{18}\text{O}$ respectively (2SE) (Table 1).

225 For the measurements of the concentrations of calcium (Ca), magnesium (Mg),
226 manganese (Mn), Sr and Rb in the carbonate fraction of the studied samples, ~50 mg of sample
227 powder was weighed into a 15 ml centrifuge tube. First, 3 ml 1 M sub-boiling double-distilled
228 acetic acid (HAc) were added and the tubes placed in an ultrasonic bath for 30 minutes. Then
229 the samples reacted at room temperature for 12 hours in order to sufficiently dissolve
230 carbonate minerals. Finally, the solutions were centrifuged for 15 minutes at 4000 rpm, and
231 the supernatant was placed into a 15 ml PFA beaker.

232 Subsequently, 2 ml ultrapure water was added into the 15 ml centrifuge tube to wash the
233 remaining residue. After being placed in an ultrasonic bath for another 30 minutes and
234 centrifugation for 15 minutes, the supernatant solutions were added into the 15 ml PFA

235 beakers and this rinsing step was repeated. The combined 7 ml of supernatant (dissolved
236 carbonate fraction) were then dried down at 120 °C and re-dissolved in 3% v/v double-distilled
237 nitric acid with a dilution factor of 1:1000 for concentration measurements (the detailed
238 dissolution procedure summarized in Fig. 3). Mn, Sr and Rb concentrations were determined
239 using a Thermo Finnigan Element XR sector-field ICP MS at the State Key Laboratory for Mineral
240 Deposits Research (Nanjing University, PR China) with a Scott type quartz spray chamber and
241 a 100 µl/min nebulizer. For instrument drift correction, we doped 10 ppb rhodium into the
242 sample solutions for elements analyzed in low-resolution mode. The Ca and Mg concentrations
243 were determined using a Sky-ray ICP OES at the same laboratory. Along with the ICP OES
244 analyses and the ICP MS analyses, a dolomite reference material (JDo-1) was repeated one
245 time per 10 analyses, the repeated analyses of JDo-1 on ICP MS and ICP OES yielded < 10 %
246 accuracy and < 5 % precision.

247 Samples with relatively high Sr concentrations and low Mn/Sr ratios were selected for Sr
248 isotope measurements. Solutions containing around 20 ng Sr were dried at 120 °C and
249 dissolved in 1 ml 3 M double-distilled nitric acid followed by standard Sr-spec resin ion-
250 chromatography for Sr purification. Sr isotope compositions were obtained on an MC-ICP-MS
251 (Thermo Neptune Plus) at the State Key Laboratory for Mineral Deposits Research in Nanjing
252 University, China. Mass fractionation was corrected by applying an assumed $^{86}\text{Sr}/^{88}\text{Sr}$ ratio =
253 0.1194 and the exponential law (Nier, 1938). The NIST SRM 987 standard was analyzed along
254 with the samples and yielded a mean value of 0.710271 ± 30 (2SE, n= 30). The Sr isotope ratios
255 and their corresponding standard error of the mean are listed in Table 1. Data mentioned in
256 the text has been normalized to the recommended NIST SRM 987 standard value of 0.710248
257 (McArthur et al., 2012).

258

259 **4. Results**

260 **4.1 C isotope feature at the Precambrian/Cambrian (PC/C)**

261

262 All C isotope data obtained on carbonate leachates are reported in the delta (δ) notation
263 relative to V-PDB and the $\delta^{13}\text{C}$ values of all analyzed carbonate samples are plotted in Fig. 4
264 and tabulated in Table 1, respectively. The $\delta^{13}\text{C}$ values fluctuate between -13.3 ‰ and +7.9 ‰.

265 The $\delta^{13}\text{C}$ curve obtained in carbonates from the Sugetbrak Fm. to the Xiaquilitagh Fm. shows
266 four prominent negative excursions (“B”, “C”, “E”, “F”, Fig. 4, here we define “excursion” as a
267 visible deviation, i.e. more than 10 % offset from their pre-excursion background values) and
268 three positive excursions (“A”, “D”, “G”, Fig. 4).

269 The first $\delta^{13}\text{C}$ positive excursion appears at the boundary between the Chigebrak Fm. and
270 Sugetbrak Fm. In this interval, the $\delta^{13}\text{C}$ values increase from -5.5 ‰ to +7.9 ‰ (Fig. 4, positive
271 excursion “A”), and then plunge down to $\sim +1$ ‰ resulting in a small negative excursion (Fig. 4,
272 negative excursion “B”). Above the negative excursion “B”, the $\delta^{13}\text{C}$ values increase to a steady
273 background value around +2.5 ‰ through the rest of the Chigebrak Fm.

274 The $\delta^{13}\text{C}$ values increase from -2.5 ‰ to +0.5 ‰ in the lower part of the Yurtus Fm.,
275 decrease rapidly to -5.1 ‰ in the middle part (“C-1”, Fig.4), and then increase again to +0.9 ‰
276 over a short interval in the upper part of the Yurtus Fm. (“C-2”, Fig. 4). At the top of the Yurtus
277 Fm., the $\delta^{13}\text{C}$ values slowly decrease to -0.5 ‰ (“C-3”, Fig. 4).

278 The whole Xiaerbrak Fm. demonstrates a prolonged positive C excursion (“D”, Fig. 4, $\delta^{13}\text{C}$
279 values vary from -0.5 ‰ to +3 ‰) with two negative swings. One occurs at the base of the
280 Xiaerbrak Fm., varying from +2.5 ‰ to +1.7 ‰, and then recovers to +3 ‰ (“D-1”, Fig. 4). The
281 other negative excursion occurs in the middle part of the Xiaerbrak Fm., decreasing from +3 ‰
282 to +1.8 ‰, and then recovering to +2.5 ‰ (“D-2”, Fig. 4).

283 At the boundary between the Wusonger and the Xiaerbrak Fm., the $\delta^{13}\text{C}$ values gradually
284 decrease to ~ 0 ‰ followed by a plunge to -2.0 ‰. The $\delta^{13}\text{C}$ values subsequently recover to
285 ~ 0 ‰ at the top of the Wusonger Fm., completing the negative excursion “E” (Fig. 4). The $\delta^{13}\text{C}$
286 values decrease sharply to a prominent negative nadir of -13.3 ‰ forming the most negative
287 excursion “F” in our dataset (Fig. 4) at the bottom of the Shayilik Fm. In the upper part of the
288 Shayilik Fm., the $\delta^{13}\text{C}$ values recover to ~ 0 ‰.

289 A $\delta^{13}\text{C}$ positive excursion (Fig. 4, positive excursion “G”) varying from -1.5 ‰ to +1.5 ‰
290 was identified in the mid-upper part of the Awatagh Fm. The positive excursion “G” consists
291 of positive excursion “G-1” (varying from -1.5 ‰ to +1.5 ‰) and “G-2” (varying from -1.5 ‰
292 to 0 ‰). Subsequently, almost the entire Xiaquilitagh Fm. sustains a constant $\delta^{13}\text{C}$ value at \sim -
293 1.0 ‰ with a minor rise to ~ 0 ‰ at the top of this formation.

294

295 **4.2 O isotope compositions at the PC/C**

296

297 The here presented O isotope data are obtained on carbonate leachates and reported in
298 the delta (δ) notation relative to V-PDB. The $\delta^{18}\text{O}$ values of the analyzed carbonates are plotted
299 in Fig. 4 and tabulated in Table 1. The $\delta^{18}\text{O}$ values from the Ediacaran samples (from the Shairik
300 section) systematically differ from those obtained in the Cambrian samples (from the
301 Xiaoerbrak section) (Fig. 4). The $\delta^{18}\text{O}$ values in the Ediacaran stratigraphic units range from -
302 9 ‰ to 0 ‰ and have a mean value of -3 ‰. By contrast, almost all samples in the Cambrian
303 units have $\delta^{18}\text{O}$ values ranging from -12 ‰ to -3 ‰ (mean = -7 ‰) except some samples in the
304 Yurtus Fm. which show more negative values.

305 Sawaki et al. (2010) reported $\delta^{18}\text{O}$ values ranging from -10 ‰ to 0 ‰, averaging at -4 ‰
306 for the Ediacaran Three Gorges area, South China. Besides, Wang et al. (2011b) obtained $\delta^{18}\text{O}$
307 values ranging from -7.9 ‰ to +3.5 ‰ through the Cambrian Series 2 to Series 3 in the
308 Penglaiba section of the Keping region. Thus, our $\delta^{18}\text{O}$ values of the two studied section are
309 generally consistent with existing Ediacaran and Cambrian datasets.

310

311 **4.3 Sr isotope compositions at the PC/C**

312

313 Samples selected for Sr isotope measurements yield low $^{85}\text{Rb}/^{86}\text{Sr}$ ratios (all samples <
314 0.0016) (Table 1), suggesting that the chromatographic Sr separation from matrix elements
315 using the specific Sr resin was effective. Further, samples with low Mn/Sr ratios (< 5) and
316 relatively high Sr concentrations (generally > 50 ppm) were selected to perform Sr isotope
317 analyses to minimize possible influence of diagenetic alteration (see discussion in section
318 5.1.2).

319 As shown in Fig. 4, the obtained $^{87}\text{Sr}/^{86}\text{Sr}$ ratios in our carbonate leachates vary between
320 0.7085 and 0.7102. From the uppermost part of the Sugetbrak Fm. to the middle Chigebrak
321 Fm., the $^{87}\text{Sr}/^{86}\text{Sr}$ values decrease from ~0.7095 to ~0.7085. Then the $^{87}\text{Sr}/^{86}\text{Sr}$ values increase
322 to ~0.7090 at the top of the Chigebrak Fm.

323 In the Yurtus Fm., the $^{87}\text{Sr}/^{86}\text{Sr}$ ratios change rapidly within the range of 0.7086 to 0.7102,
324 followed by a plateau at ~0.7088 through most of the Xiaoerbrak Fm. Finally, a rise to ~0.7090

325 at the top of the Xiaerbrak Fm. coincides with the positive $\delta^{13}\text{C}$ excursion “D” and a $\delta^{13}\text{C}$
326 platform at 0 ‰ above the excursion “D”. In the Wusonger Fm., the $^{87}\text{Sr}/^{86}\text{Sr}$ ratios rise slowly
327 from ~ 0.7088 to ~ 0.7090 and sharply increase to 0.7092 at the bottom of the Shayilik Fm.

328 At the bottom of the Shayilik Fm., the recorded $^{87}\text{Sr}/^{86}\text{Sr}$ ratios drop to ~ 0.7087 , coinciding
329 with the prominent negative $\delta^{13}\text{C}$ excursion “F”. Subsequently, the $^{87}\text{Sr}/^{86}\text{Sr}$ ratios gradually
330 increase to ~ 0.7092 in the mid-upper part of Awatagh Fm., coinciding with the rising limb of
331 the positive $\delta^{13}\text{C}$ excursion “G”. Apart from only one outlier at 0.7086 in the lower part of
332 Xiaqiulitagh Fm., the $^{87}\text{Sr}/^{86}\text{Sr}$ ratios recorded in carbonates from the rest of the Cambrian (i.e.
333 Xiaqiulitagh Fm. and Upper Awatagh Fm.) is constant at ~ 0.7091 .

334

335 5. Discussion

336 5.1 Assessing possible processes altering the C/Sr isotope compositions

337 5.1.1 The effect of silicate leaching and dolomitization on the $^{87}\text{Sr}/^{86}\text{Sr}$

338

339 To evaluate the influence of possible silicate leaching with our digestion method on the
340 obtained $^{87}\text{Sr}/^{86}\text{Sr}$ ratios, a cross plot of $^{87}\text{Sr}/^{86}\text{Sr}$ versus [Rb] (i.e. Rb concentration normalized
341 by $\text{CaO}+\text{MgO}$) is provided in the Fig. 6G and 6N. The absence of apparent correlations between
342 Sr concentrations as well as $^{87}\text{Sr}/^{86}\text{Sr}$ ratios and [Rb] indicates our digestion method avoids
343 substantial silicate mineral leaching as these contain much higher Rb contents and more
344 radiogenic Sr than carbonate minerals. Moreover, there are no apparent correlations in all
345 analyzed carbonates in the cross plots of $^{87}\text{Sr}/^{86}\text{Sr}$ -Mg/Ca and $^{87}\text{Sr}/^{86}\text{Sr}$ -carbonate content (%)
346 (Fig. 6E, 6F, 6L, 6M), indicating $^{87}\text{Sr}/^{86}\text{Sr}$ ratios may not be influenced by dolomitization. Thus,
347 we consider our sample digestion method as a suitable method to treat ancient marine
348 carbonates, and our obtained $^{87}\text{Sr}/^{86}\text{Sr}$, $\delta^{18}\text{O}$ and $\delta^{13}\text{C}$ values as suitable proxies to record
349 ambient seawater compositions (section 5.1.2 and 5.1.3).

350

351 5.1.2 The influence of diagenetic alteration

352

353 Any possible diagenetic overprint superimposed on the primary geochemical seawater

354 signatures of marine carbonates (e.g. late-stage fluid alteration causing carbonate dissolution
355 and reprecipitation) requires to be examined in detail, and any altered samples need to be
356 rigorously screened out. In general, typical chemical characteristics of carbonates that
357 experienced a high degree diagenetic alteration yield elevated Mn/Sr ratios, low $\delta^{18}\text{O}$ values
358 and possibly low $\delta^{13}\text{C}$ values and elevated $^{87}\text{Sr}/^{86}\text{Sr}$ ratios (e.g. Marshall, 1992; Kaufman et al.,
359 1993; Derry et al., 1994). Thus, cross-plots of Mn/Sr ratios, $\delta^{18}\text{O}$, $\delta^{13}\text{C}$ and $^{87}\text{Sr}/^{86}\text{Sr}$ ratios are
360 useful and widely used tools to evaluate the degree of alteration. Moreover, Mn/Sr ratios and
361 $\delta^{18}\text{O}$ values are considered useful diagnostic tools of late-stage alteration of water-rock
362 interaction (Marshall, 1992; Derry et al., 1994; Kaufman and Knoll, 1995). Almost all the
363 dolostone samples in this study, as well as those from other sections in the Keping area (e.g.
364 Guo et al., 2017), have low Sr concentrations (< 300 ppm) which is possibly indicative for low
365 initial seawater Sr concentrations at the time of deposition (cf. Li et al., 2011). Here we selected
366 more generous cut-offs for our studied dolostone samples during our pre-screening: samples
367 that clearly deviate from the main data group (Mn/Sr > 5 and $\delta^{18}\text{O}$ < -10 ‰, Fig. 5) are
368 considered as not representing primary seawater $^{87}\text{Sr}/^{86}\text{Sr}$ signals (Derry et al., 1994; Kaufman
369 and Knoll, 1995; Li et al., 2013; Zhou et al., 2020).

370 Nearly all Ediacaran samples show isotopically heavier $\delta^{18}\text{O}$ values (> -10 ‰) and lower
371 Mn/Sr (< 5) (Fig. 5). However, carbonates from the Sugetbrak Fm. show a positive correlation
372 between $\delta^{18}\text{O}$ and $\delta^{13}\text{C}$ ($R^2 = 0.95$; Fig. 5A). Besides, correlations between radiogenic $^{87}\text{Sr}/^{86}\text{Sr}$
373 ratios and low $\delta^{13}\text{C}$ and $\delta^{18}\text{O}$ values are present ($R^2 = 0.89$ and 0.95 respectively; Fig. 6H and
374 6I), suggesting that the Sugetbrak Fm. might have suffered substantial diagenetic alteration by
375 basin-type fluids which enriched in “light” C and O and radiogenic Sr isotopes (see Hohl et al.,
376 2015b).

377 Most Cambrian samples show no apparent correlations in the cross-plot of $\delta^{13}\text{C}$ versus
378 $\delta^{18}\text{O}$ (Fig. 5B). However, carbonate leachates of the Yurtus Fm. show elevated Mn/Sr ratios (>
379 5; circles in Fig. 6D) and a positive correlation between $^{87}\text{Sr}/^{86}\text{Sr}$ ratios and $\delta^{18}\text{O}$ values (Fig.
380 6B), which indicates that the obtained Sr isotope compositions in the Yurtus Fm. have been
381 influenced by the mixing of restricted basin Sr signals with less radiogenic fluids (see section
382 5.1.3). However, the obtained $\delta^{13}\text{C}$ values in the Yurtus Fm. carbonates show no correlation
383 with obtained $^{87}\text{Sr}/^{86}\text{Sr}$ and $\delta^{18}\text{O}$ values (Fig. 6A and 5B), suggesting that the carbonates in the

384 Yurtus Fm. likely preserved the primary seawater C isotope composition.

385 In the Wusonger Fm., the $\delta^{18}\text{O}$ value of one sample yields -10 ‰ and its $\delta^{13}\text{C}$ value is
386 similarly low (-6.7 ‰) (Fig. 5B). Therefore, the sample is excluded from the seawater C and Sr
387 evolution curve (in sections 5.2 and 5.3). Likewise, the $\delta^{18}\text{O}$ value of another data point in the
388 Shayilik Fm. is lower than -10 ‰. However, no paired decrease in the $\delta^{13}\text{C}$ excludes the
389 possibility of significant post-depositional fluid alteration for this sample.

390 Apart from these obviously altered samples, there is no clear correlation in the cross-plots
391 of $\delta^{18}\text{O}$ - $\delta^{13}\text{C}$ (Fig. 5), $^{87}\text{Sr}/^{86}\text{Sr}$ - $\delta^{18}\text{O}$, $^{87}\text{Sr}/^{86}\text{Sr}$ - $\delta^{13}\text{C}$, $^{87}\text{Sr}/^{86}\text{Sr}$ -[Sr], $^{87}\text{Sr}/^{86}\text{Sr}$ -Mn/Sr, $^{87}\text{Sr}/^{86}\text{Sr}$ -
392 Mg/Ca and $^{87}\text{Sr}/^{86}\text{Sr}$ - Carbonate content (%) (Fig. 6). Thus, we consider these samples are well-
393 preserved marine carbonates, of which their $^{87}\text{Sr}/^{86}\text{Sr}$ and $\delta^{13}\text{C}$ values may record primary
394 seawater compositions.

395

396 **5.1.3 Fluid mixing model for Sr isotope compositions in the Yurtus Fm.**

397

398 Samples of the Yurtus Fm. show elevated Mn/Sr ratios (> 5; circles in Fig. 6D), arguing for
399 excess in Mn possibly due to pore water-redox controlled remobilization of MnO colloids
400 (Cowie et al., 1995; Thomson et al., 1998; Hohl et al., 2017). Besides, carbonate leachates of
401 the Yurtus formation show a positive correlation between $^{87}\text{Sr}/^{86}\text{Sr}$ ratios and $\delta^{18}\text{O}$ values (Fig.
402 6B) with two endmembers, one with the least-radiogenic $^{87}\text{Sr}/^{86}\text{Sr}$ ratios and light $\delta^{18}\text{O}$ values,
403 while the other endmember has more radiogenic Sr and heavier O isotopic compositions.
404 However, diagenetic basin-type fluid-alteration usually results in a negative correlation
405 between $^{87}\text{Sr}/^{86}\text{Sr}$ ratios and $\delta^{18}\text{O}$ values (Hohl et al., 2015b).

406 In order to explore the mechanisms that may generate the full range in O and Sr isotope
407 compositions observed in the Yurtus Fm., a simple open-system fluid mixing model (Jacobsen
408 and Kaufman, 1999) is applied. As inferred from the paired Sr and O isotope results, two
409 endmembers have been identified: one is a radiogenic localized basin-type seawater ($^{87}\text{Sr}/^{86}\text{Sr}$
410 = 0.7102, $\delta^{13}\text{C}$ = 0.5 ‰, $\delta^{18}\text{O}$ = 0.5 ‰ and Sr = 50 ppm), while the other endmember is a
411 meteoric fluid with $^{87}\text{Sr}/^{86}\text{Sr}$ of 0.7085, $\delta^{13}\text{C}$ of -5 ‰, $\delta^{18}\text{O}$ of -14 ‰ and 600 ppm Sr (Fig. 6A,
412 6B and 6C). Data of the Yurtus carbonates falls on a mixing trend line between the localized

413 seawater and a lighter $\delta^{18}\text{O}$ bearing meteoric fluid, exhibiting mixing ratios between 0.01 to 5
414 (Fig. 6A, 6B, 6C).

415 In general, meteoric fluids comprise more radiogenic Sr and lower Sr concentrations than
416 contemporaneous seawater (e.g. Xu and Han, 2009). However, the hypothetical global
417 regression at the Precambrian/Cambrian boundary presumably causing the 'great
418 unconformity' (Brasier, 1996; Ishikawa et al., 2008; Sawaki et al., 2008; He et al., 2017) might
419 have exposed extensive amounts of Precambrian marine carbonates on the shelf, and thus,
420 meteoric fluids leaching these lithologies, might have inherited their Ediacaran seawater-like
421 Sr isotopic compositions and Sr concentrations. He et al. (2018) suggested a wide geographical
422 distribution of paleo-karsts at the top of the Chigebrak Fm. and speculated that there is an
423 Ediacaran carbonate paleo-upland adjacent to the Xiaoerbrak and Shiirik sections during this
424 time interval. Furthermore, similar Sr isotope composition and Sr concentration of the
425 Chigebrak Fm. carbonates with those of our modelled meteoric fluid ($^{87}\text{Sr}/^{86}\text{Sr}$ ratio ~ 0.7085 ;
426 Sr concentration ~ 600 ppm) argue for a contribution to the carbonate $^{87}\text{Sr}/^{86}\text{Sr}$ in the Yurtus
427 Fm. from such an endmember.

428 On the other hand, the Sr isotope composition of the seawater endmember (~ 0.7102) in
429 this study is more radiogenic when compared to the reported $^{87}\text{Sr}/^{86}\text{Sr}$ seawater ratio from the
430 correlative strata in South China (~ 0.7085 , Li et al., 2013), which may be explained by a local
431 rather than global signature, arguing for a restricted basin depositional environment on the
432 shelf of the Tarim craton during this period (cf. Bai et al., 2019; Yao et al., 2017).

433 Several syn-genetic ancient weathering crusts in the Yurtus Fm. imply subaerial exposure
434 environments and the occurrence of black shale in the lower part of the Yurtus Fm. indicate
435 facies change to deep-water deposition. Hence the two depositional settings probably
436 represent sea-level fluctuations. Because the early Cambrian Tarim Basin was presumably
437 restricted and underwent several second-order sea-level shifts (Yao et al., 2017), we argue that
438 meteoric fluids leaching from aerial exposed Ediacaran shelf carbonates are mixed with
439 confined basin waters. The basin may be localized at the late-Ediacaran and the basin water
440 may be influenced by enhanced silicate weathering of hinterland during the global regression
441 at the late-Ediacaran (Haq and Schutter, 2008).

442

443 **5.2 C isotope stratigraphic and bio-stratigraphic correlation**

444

445 Before performing a stratigraphic correlation between the Tarim block and other time-
446 equivalent paleo-continents, it is essential to note that our obtained C isotope variation curve
447 can be correlated to the existing datasets from other sections in the Tarim Basin. For example,
448 similar rapid increasing $\delta^{13}\text{C}$ trends in the top part of the Sugetbrak Fm. have been reported at
449 a composite profile from seven sections in the Quruqtagh area and at the Penglaiba section in
450 the Keping area by Xu et al. (2003) and He et al. (2007), respectively. Further, Guo et al. (2017)
451 reported extreme negative $\delta^{13}\text{C}$ values (down to -14.3‰) at the boundary of the
452 Wusonger/Shayilik Fm. in the Penglaiba section in the Keping area, similar to our negative $\delta^{13}\text{C}$
453 excursion "F" with respect to the total magnitude of the excursion (down to -13.3‰).

454 During the PC/C, the Tarim Basin was situated in a low latitude location on the continental
455 margin shelf (Yu et al., 2009), leading to the deposition of shallow water carbonate sediments.
456 This makes the Tarim Basin sections lithologically and paleogeographical similar to other
457 shallow marine PC/C sections on the Yangtze Platform (McKerrow et al., 1992; Merdith et al.,
458 2017; Huang et al., 2000; Zhan et al., 2007; Hohl et al., 2015a). Therefore, we here apply the
459 Ediacaran/Cambrian bio-chemostratigraphic division of South China as the primary reference
460 frame for the correlation with our studied bio-chemostratigraphic data from the two sections.
461 Fig. 7 shows the carbonate C isotope evolution curve from the Keping region and our proposed
462 C isotope stratigraphic correlation with the reported composite carbonate C isotope evolution
463 curves of South China from the late Ediacaran (Jiang et al., 2007), across the Ediacaran-
464 Cambrian transition (Li et al., 2013), and through the Cambrian Series 2 to Series 3 (Guo et al.,
465 2010). On a global scale, we also compare our Tarim Basin carbonate C isotope evolution curve
466 with records from North India during late Ediacaran (Kaufman et al., 2006) and the Cambrian
467 global seawater standard $\delta^{13}\text{C}$ variations proposed by Zhu et al. (2006), which is mainly derived
468 from South China shallow-water carbonates. Similarly, carbonate C isotope datasets from
469 Laurentia (Montañez and Banner, 2000) and Siberia (Wotte et al., 2011) spanning the
470 Cambrian Series 2 – Series 3 strata and carbonate C isotope data from Southwest Mongolia
471 spanning the Ediacaran-Cambrian transition (Brasier et al., 1996) are used for correlation.

472 According to the current Ediacaran chronostratigraphy proposed by Zhu et al. (2007), the

473 upper Doushantuo negative $\delta^{13}\text{C}$ excursion (DOUNCE, equivalent to the Shuram/Wonoka
474 excursion) ranges from ca. 560 Ma to 551 Ma and the following positive $\delta^{13}\text{C}$ excursion that
475 ended at around 548 Ma, which provide a general baseline reference for C isotope stratigraphy
476 in the mid-late Ediacaran. In South China, a sharp $\delta^{13}\text{C}$ rise predates the positive $\delta^{13}\text{C}$ excursion
477 at the boundary between the Doushantuo and the Dengying Fm. (e.g. Jiang et al., 2007).
478 Similar $\delta^{13}\text{C}$ variations were observed at the boundary between the Sugetbrak and Chigebrak
479 Fm. in the Tarim Basin which lead to the hypothesis that the Chigebrak Fm. is stratigraphically
480 equivalent to the Dengying Fm. in South China (He et al., 2007). The studied Shairik section is
481 merely 40 km away from the site investigated by He et al. (2007) and similarly obtained
482 carbonate $\delta^{13}\text{C}$ curves at both sites suggest that the Chigebrak carbonates have only
483 experienced minimal diagenetic alteration and thus, may archive the $\delta^{13}\text{C}$ signals of coeval
484 Ediacaran paleo-seawater. As mentioned above, the positive excursion "A" occurring at the
485 boundary between the Chigebrak and Sugetbrak Fm. is consistent with the Ediacaran division
486 plan proposed by Zhu et al. (2007) and He et al. (2007). Since, late Ediacaran South China strata
487 bears no other obvious negative $\delta^{13}\text{C}$ excursion above the ca. 548 Ma old positive $\delta^{13}\text{C}$
488 excursion (Fig. 7I) while North Indian strata of the same time reveals two negative $\delta^{13}\text{C}$
489 excursions (Fig. 7J), the negative excursion "B" obtained in the Tarim Basin carbonates cannot
490 be considered as a global $\delta^{13}\text{C}$ excursion.

491 A slight $\delta^{13}\text{C}$ decline at the top of Chigebrak Fm. is similar to the $\delta^{13}\text{C}$ excursion at the top
492 of the Dengying Fm. (Fig. 7I) and a subsequent sharp decrease (negative Excursion "C") across
493 the top of the Chigebrak Fm. and the lower Yurtus Fm. is suggested as equivalent to the
494 globally identified BACE (BASal Cambrian C isotope Excursion) (Fig. 7B, 7G, 7H). This
495 interpretation is consistent with previous studies in which the parallel unconformity between
496 the Chigebrak Fm. and the Yurtus Fm. is considered to represent the boundary between the
497 Ediacaran and Cambrian in the Tarim Basin (Yao et al., 2014; He et al., 2018).

498 The appearance of SSFs, that are widely distributed during the Terreneuvian Series (Zhu
499 et al., 2006), can further be used for robust biostratigraphic correlation. In South China, the
500 *Anabarites-Protohertzina-Arthochites* assemblage, the first Terreneuvian SSFs assemblage in
501 the Southwest Chinese Province of Yunnan, agrees well with the earliest skeletal faunas on the
502 Siberian Platform and the Canadian Cordillera (Qian and Bengtson, 1989; Steiner et al., 2004;

503 Yang et al., 2014a; Yang et al., 2014b). The lower Yurtus Fm. in the Tarim Basin, which contains
504 an *Anabarites-Protohertzina* assemblage (Qian and Xiao, 1984), can be correlated to the lower
505 Terreneuvian. Moreover, because the second assemblage (*Siphogonuchites-Paragloborilus*)
506 and the last assemblage (*Lapworthella-Tannuolina-Sinosachites*) found in Yunnan (Qian and
507 Xiao, 1984; Qian and Bengtson, 1989; Yang et al., 2014a) also appear in the upper Yurtus Fm.
508 (i.e. *Paragloborilus-Lapworthella* assemblage), we suggest that all or most of the Yurtus Fm.,
509 conformably underlying the trilobite-bearing Xiaerbrak Fm. can be best assigned to the
510 Terreneuvian Series.

511 The Terreneuvian Series (the first series of the Cambrian) is characterized by more than
512 three carbonate $\delta^{13}\text{C}$ excursions recorded in South China sections, i.e. BACE, ZHUCE
513 (ZHUjiaqing C isotope Excursion) and SHICE (SHIyantou C isotope Excursion) (Fig. 7B). One
514 negative $\delta^{13}\text{C}$ excursion (Fig. 4, "C-2") and two positive $\delta^{13}\text{C}$ excursions (Fig. 4, "C-1" and "C-3")
515 are recorded in the Yurtus Fm. carbonates, not perfectly matching the reported excursions'
516 amounts and magnitudes during the Terreneuvian epoch in South China. As syn-genetic
517 ancient weathering crusts were recognized in field observation, several depositional hiatuses
518 in the Yurtus Fm. may explain the differences in the early Cambrian $\delta^{13}\text{C}$ profiles between
519 South China and the Tarim Basin.

520 The Cambrian Stage 3 is marked by the first appearance of trilobites and two positive $\delta^{13}\text{C}$
521 excursions (Cambrian Arthropod Radiation isotope Excursion (CARE) and Mingxinsi C Isotope
522 Excursion (MICE), Zhu et al., 2006). Due to the appearance of *Shizhudiscus* at the bottom of
523 the Xiaerbrak Fm., it may be assigned to the base of the Cambrian Series 2. Besides, due to
524 the consistency in $\delta^{13}\text{C}$ variation trends (a major positive excursion with a sub-negative
525 excursion) and magnitudes (increase from ~ 0 ‰ to +3 ‰) between the upper part of the
526 positive $\delta^{13}\text{C}$ excursion "D" (with the sub-negative excursion "D-2") and the MICE (Fig. 7), this
527 study correlates the upper part of the excursion "D" as equivalent to the MICE in South China.
528 Although the lower part of the excursion "D" is not perfectly matching in shape to the CARE,
529 the similar magnitude of the excursion "D-1" (decreasing from +2.5 ‰ to +1.7 ‰) and the
530 appearance of *Shizhudiscus* broadly confirms that the negative $\delta^{13}\text{C}$ excursion "D-1" likely
531 corresponds to the falling limb of the CARE. Therefore, combining the evidence that the
532 appearance of *Shizhudiscus* occurs slightly later than the start of the Cambrian Series 2 (Fig.

533 7B, the green arrow represents the strata with *Shizhudiscus* occurrence), the boundary
534 between the Yurtus Fm. and the Xiaerbrak Fm. is considered to be stratigraphically higher
535 than the base of the Cambrian Stage 3 in the Tarim Basin.

536 The ROECE (Redlichiid–Olenellid Extinction C isotope Excursion; Zhu et al., 2006), the most
537 negative $\delta^{13}\text{C}$ excursion throughout the Cambrian Series 2 to Furongian (Fig. 7C, 7D, 7E, 7F, e.g.
538 Fan et al., 2011; Guo et al., 2017), is considered to represent the boundary between the
539 Cambrian Stage 4 and the Wuliuan Stage (Montañez and Banner, 2000; Zhu et al., 2006; Guo
540 et al., 2010; Wotte et al., 2011; Guo et al., 2017). Accordingly, we consider that the most
541 negative $\delta^{13}\text{C}$ excursion “F” in our studied carbonates is equivalent to the ROECE and the
542 bottom of the Shayilik Fm. probably represents the base of the Cambrian Series 3 (Fig. 7). This
543 correlation is consistent with the stratigraphic correlations by Guo et al. (2017) and Wang et
544 al. (2011b) and the appearance of the *Kunmingaspis-Chittdilla* (Fig. 2) in the Shayilik Fm. (Zhou,
545 2001; Peng, 2009), the biozone range of which overlaps with the Wuliuan Stage (Wang et al.,
546 2011a, Fig. 7, the yellow arrow represent the strata with *Kunmingaspis-Chittdilla* occurrence).
547 Below the negative $\delta^{13}\text{C}$ excursion “F”, by comparing the $\delta^{13}\text{C}$ variation magnitude, the
548 negative $\delta^{13}\text{C}$ excursion “E” in the Wusonger Fm. may be equivalent to the Archaeocyathid
549 Extinction C isotope Excursion (AECE, Zhu et al. (2006)). This hypothesis is further supported
550 as *Paokannia* is considered to survive from mid-Cambrian Series 2 (late Stage 3) to the end of
551 Cambrian Series 2 (Zhou, 2001; Peng, 2009) (Fig. 7, the orange arrow represents the
552 *Paokannia*-bearing strata).

553 The positive $\delta^{13}\text{C}$ excursion “G” is the only apparent positive excursion overlying the
554 negative $\delta^{13}\text{C}$ excursion “F”, and this temporal distribution is similar to the chronological order
555 between SPICE (Steptoean C-Isotope Positive Excursion) and ROECE. However, the positive
556 $\delta^{13}\text{C}$ excursion “G” is not as distinct as the SPICE reported from other paleo-continent and its
557 magnitude of 3‰ is smaller than the magnitude of the SPICE recorded elsewhere (~4-6‰) (Li
558 et al., 2018), which may suggest depositional hiatuses existing at the base of the Furongian in
559 the Keping area. Considering that no depositional hiatus is reported in the Awatagh Fm., we
560 tentatively speculate that the positive excursion “G” is identical to SPICE but suggest further
561 verification by independent sedimentological and geochemical proxies. Because the SPICE is
562 the typical characteristic of the start of the Paibian stage of the Furongian Series (Saltzman et

563 al., 1998; Peng et al., 2004; Zhu et al., 2006; Fan et al., 2011), a correlation of the positive
564 excursion “G” with SPICE would mean that the upper Awatagh Fm. in the NW Tarim Basin may
565 be preliminarily assigned to the Furongian Series.

566

567 **5.3 Sr isotope stratigraphic correlation**

568

569 Based on the carbonate C isotope stratigraphic correlation, a carbonate-based seawater
570 $^{87}\text{Sr}/^{86}\text{Sr}$ curve (Fig. 8 red dashed line) is constructed using a “lower limit” Sr isotope data
571 approach (Jones et al., 1994). Fig. 8 shows the obtained least-diagenetically altered $^{87}\text{Sr}/^{86}\text{Sr}$
572 ratios from carbonate leachates in the Tarim Basin of this study (Fig. 8D), a proposed standard
573 Cambrian global seawater $^{87}\text{Sr}/^{86}\text{Sr}$ curve (Fig. 8A, Peng et al., 2012; Zhu et al., 2019)
574 constructed after the work of Derry et al. (1994); Montañez and Banner (2000); Ebner et al.
575 (2001); Kouchinsky et al. (2008) as well as the $^{87}\text{Sr}/^{86}\text{Sr}$ values recorded in carbonates from the
576 Yangtze Platform, South China during the Ediacaran-Cambrian transition period (Fig. 8B, C,
577 Sawaki et al., 2008; Sawaki et al., 2010).

578 In this study, our inferred seawater $^{87}\text{Sr}/^{86}\text{Sr}$ ratios from the Keping region decrease from
579 ~ 0.7090 to ~ 0.7085 in the lower part of the Chigebrak Fm. with a slight rebound to more
580 radiogenic values (~ 0.7091) at the top of this formation. This trend mimics the variation
581 recorded in the terminal Proterozoic Dengying and Doushantuo Fm. on the South China
582 Yangtze Platform (Sawaki et al., 2010). Sawaki et al. (2008) and Hohl et al. (2015a) reported
583 increasing $^{87}\text{Sr}/^{86}\text{Sr}$ ratios in the upper Dengying Fm., coinciding the similar increasing trends
584 recorded in the contemporarily deposited upper Chigebrak Fm. carbonates (Fig. 8). However,
585 the magnitude of the total rise in $^{87}\text{Sr}/^{86}\text{Sr}$ ratios is smaller than those recorded by Sawaki et
586 al. (2008) but comparable to that reported by Hohl et al. (2015a). This may be explained with
587 erosion and denudation of the uppermost Chigebrak carbonate platform (He et al., 2018) and
588 the incomplete outcrop situation at the Xiaofenghe section in South China (Hohl et al., 2015a).
589 As a summary from the above discussion, we consider the Chigebrak Fm. to be equivalent to
590 the Dengying Fm., revealing consistent Sr and C isotope stratigraphy (Fig. 7).

591 As discussed above, the scattering $^{87}\text{Sr}/^{86}\text{Sr}$ ratios obtained in carbonates from the Yurtus
592 Fm. are likely results of restricted-basin seawater mixed with carbonate weathering fluid

593 (section 5.1.3, Fig. 6A, 6B and 6C) and thus may not be compared to the global seawater
594 $^{87}\text{Sr}/^{86}\text{Sr}$ curve. We, therefore, exclude these data points from our Sr isotope composition
595 curve as they may represent a restricted-basin signal (Bai et al., 2019; Yao et al., 2017) that is
596 unsuitable for global correlations.

597 From the bottom of the Xiaerbrak Fm. to the bottom of Shayilik Fm., the obtained
598 $^{87}\text{Sr}/^{86}\text{Sr}$ ratios in carbonate leachates rise from ~ 0.7087 to ~ 0.7090 , consistent with the
599 reported seawater $^{87}\text{Sr}/^{86}\text{Sr}$ evolutionary curve during the Cambrian Series 2 (Montañez and
600 Banner, 2000; Ebner et al., 2001; Peng et al., 2012; Zhu et al., 2019) (Fig. 8). At the bottom of
601 the Shayilik Fm., the $^{87}\text{Sr}/^{86}\text{Sr}$ ratios in the carbonates leachates decrease from ~ 0.7090 to
602 ~ 0.7087 , analogous to the global seawater $^{87}\text{Sr}/^{86}\text{Sr}$ curve at the Stage 4/Wuliuan Stage
603 boundary (Peng et al., 2012; Zhu et al., 2019) (Fig. 8). Subsequently, $^{87}\text{Sr}/^{86}\text{Sr}$ ratios increase
604 again at the middle part of the Shayilik Fm. and reach a peak of ~ 0.7092 in the middle part of
605 the Awatagh Fm., coinciding the starting point of the $\delta^{13}\text{C}$ positive excursion "G" (Fig. 7). The
606 SPICE and the maximum $^{87}\text{Sr}/^{86}\text{Sr}$ peak at around ~ 0.7092 characterize the boundary between
607 the Guzhuangian and Paibian Stage (Montañez and Banner, 2000; Ebner et al., 2001;
608 Kouchinsky et al., 2008; Fan et al., 2011). As we inferred hereinbefore, based on the
609 coincidence between the $^{87}\text{Sr}/^{86}\text{Sr}$ peak and the $\delta^{13}\text{C}$ positive excursion "G", this study
610 correlates the middle-upper part of the Awatagh Fm. approximately to the boundary between
611 the Guzhuangian and Paibian Stage. Apart from only one outlier with a much lower $^{87}\text{Sr}/^{86}\text{Sr}$
612 value (~ 0.7086) in the lower part of Xiaqiulitagh Fm., the $^{87}\text{Sr}/^{86}\text{Sr}$ ratios slowly decrease from
613 ~ 0.7092 to ~ 0.7091 from the middle part of the Awatagh Fm. to the Xiaqiulitagh Fm., which
614 are broadly similar to those ratios reported by Ebner et al. (2001) and Montañez and Banner
615 (2000).

616 In this study, as a whole, the $^{87}\text{Sr}/^{86}\text{Sr}$ ratios and the C isotope variations obtained in
617 carbonates from two sections of the Keping region spanning the late-Ediacaran to the late-
618 Cambrian show global consistency (Fig. 7 and Fig. 8), except those analyses performed in
619 presumably diagenetically altered Sugetbrak Fm. and the Yurtus Fm. carbonates. Our results
620 from the stratigraphic correlation between the Keping area and other parts of the world from
621 the late-Ediacaran to the late-Cambrian are as follows:

622 i. The Chigebreak Fm. is presumably equivalent to the Dengying Fm. of the South China

623 Yangtze Platform.

624 ii. The base of the Xiaerbrak Fm., which contains *Shizhudiscus* fossils, is slightly
625 stratigraphically higher than the base of the Cambrian Stage 3.

626 iii. The most negative $\delta^{13}\text{C}$ excursion “F” together with a decrease in $^{87}\text{Sr}/^{86}\text{Sr}$ characterize
627 the bottom of the Shayilik Fm., which can be regarded as the boundary between the Cambrian
628 Stage 4 and Cambrian Wuliuan in the Tarim Basin. This stratigraphic correlation is consistent
629 with the existing paleontological constraints in the Tarim Basin (Peng, 2009).

630 iv. The carbonate samples of the upper-middle part of the Awatagh Fm. archive a positive
631 $\delta^{13}\text{C}$ excursion “G” (likely equivalent to SPICE) and a peak value of $^{87}\text{Sr}/^{86}\text{Sr}$ ratios at ~ 0.7092 ,
632 presumably marking the base of the Cambrian Paibian Stage.

633

634 **5.4 Paleoenvironmental implications for the Tarim Basin at the PC/C**

635

636 Overall, the obtained seawater $^{87}\text{Sr}/^{86}\text{Sr}$ ratios in carbonates from this study and those
637 recorded in numerous carbonate successions from different paleo-continent show a stepwise
638 increase from late Ediacaran to late Cambrian (Furongian) which is superimposed by several
639 minor declines (Fig. 8 and Fig. 9). Meanwhile, the obtained $\delta^{13}\text{C}$ values in our study show four
640 prominent negative and three positive excursions that are identical to trends found in various
641 carbonate-based C isotope records at a global scale (Fig. 9).

642

643 **5.4.1 Late Ediacaran paleoenvironmental reconstruction of the Tarim Basin**

644

645 The integrated global carbonate $\delta^{13}\text{C}$ trend of late Ediacaran sections (Fig. 7) suggests that
646 the carbonate $\delta^{13}\text{C}$ record from the Keping region probably archives a primary global seawater
647 $\delta^{13}\text{C}$ signal during this interval. According to our C and Sr isotope stratigraphic correlation (Fig.
648 7 and Fig. 8), the lower part of the positive $\delta^{13}\text{C}$ excursion “A” in this study probably
649 corresponds to the uppermost Doushantuo negative $\delta^{13}\text{C}$ excursion which ended at around
650 551 Ma (Zhu et al., 2007). It is controversial whether DOUNCE/Shuram-Wonoka is
651 representative of the primary coeval seawater chemistry or not. Some researchers explained
652 the extreme negative $\delta^{13}\text{C}$ values of the DOUNCE/Shuram-Wonoka event by global changes in

653 the C isotopic composition of Ediacaran seawater as a result of the remineralization of a vast
654 dissolved organic carbon reservoir (cf. Rothman et al., 2003; Fike et al., 2006). Others
655 suggested a large flux of methane released from the seafloor (Bjerrum and Canfield, 2011) as
656 the cause, while Derry (2010) argued for a large-scale post-depositional diagenetic alteration.

657 In this study, samples from the Sugetbrak Fm., which constitute for the most of the lower
658 part of the positive excursion "A", exhibit a strong positive correlation in the $\delta^{13}\text{C}$ - $\delta^{18}\text{O}$ plot (Fig.
659 5A) and negative correlations in the $^{87}\text{Sr}/^{86}\text{Sr}$ - $\delta^{13}\text{C}$, $^{87}\text{Sr}/^{86}\text{Sr}$ - $\delta^{18}\text{O}$ plots (Fig. 6H, 6I) in the
660 Sugetbrak Fm. and the correlations support a severe diagenetic alteration imprint. $^{87}\text{Sr}/^{86}\text{Sr}$
661 data of the diagenetically altered samples of this study is also comparable to the altered
662 samples in the Doushantuo Fm. from South China (Sawaki et al., 2010) (Fig. 8). However,
663 although this diagenetic alteration may relate to Derry's large-scale or even global post-
664 depositional diagenetic $\delta^{13}\text{C}$ variation hypothesis (Derry, 2010), it requires further diagenesis
665 evaluation by other proxies at a global scale (e.g. lithological evidence of widespread
666 deposition of authigenic carbonate).

667 Immediately above the altered samples from the Sugetbrak Fm., samples from the base
668 of the Chigebrak Fm. delineate the peak of the positive $\delta^{13}\text{C}$ excursion "A" and its recovery
669 towards lower values. Wood et al. (2015) also reported a positive $\delta^{13}\text{C}$ excursion (ca. 548 Ma,
670 equivalent to the excursion "A") in the carbonates of the Nama group from Namibia postdating
671 the appearance of calcified metazoans. The steep increase of calcified metazoans in the total
672 primary production could have amplified organic carbon burial and led to a net drawdown of
673 atmospheric CO_2 , ultimately shifting the seawater $\delta^{13}\text{C}$ to more positive values (Knoll et al.,
674 1996). In this study, the positive $\delta^{13}\text{C}$ excursion "A" is accompanied by an $^{87}\text{Sr}/^{86}\text{Sr}$ decrease,
675 coinciding a lithological transition from more siliciclastic carbonates to micritic carbonates (Fig.
676 9) that may relate to a transgression event (Deng et al., 2019). Therefore, the recorded $^{87}\text{Sr}/^{86}\text{Sr}$
677 decrease in carbonates may imply suppressed continental silicate weathering due to reduced
678 continental silicates being exposed to physical-chemical weathering on land (Jiang et al., 2007)
679 and possible associated atmospheric CO_2 drawdown (Knoll et al., 1996).

680 Carbonates deposited in the Keping region above the positive excursion "A" show a
681 negative $\delta^{13}\text{C}$ excursion "B" accompanied by a decrease in seawater $^{87}\text{Sr}/^{86}\text{Sr}$ ratios (Fig. 9),
682 which was also reported in North India (Fig. 7, Jiang et al., 2007; Kaufman et al., 2006). The

683 further decrease in seawater $^{87}\text{Sr}/^{86}\text{Sr}$ might result from continued continental flooding and
684 suppressed continental weathering. At the same time, the $\delta^{13}\text{C}$ negative excursion “B” could
685 be explained by global ocean stratification by upwelling of deep-sea ^{13}C depleted suboxic to
686 euxinic waters in the terminal Proterozoic (cf. Jiang et al., 2007). Finally, the upper Chigebrak
687 Fm. carbonates record a rising seawater $^{87}\text{Sr}/^{86}\text{Sr}$ trend, consistent with the global seawater
688 record in the late Ediacaran pristine carbonate (Brasier et al., 1996; Sawaki et al., 2008),
689 possibly indicating enhanced silicate weathering as a result of increased silicates exposure
690 during large-scale regression (Brasier, 1996; Haq and Schutter, 2008).

691

692 **5.4.2 Terrenevian Series and Cambrian Series 2 paleoenvironmental reconstruction of** 693 **the Tarim Basin**

694

695 As discussed in section 5.1.3, $^{87}\text{Sr}/^{86}\text{Sr}$ variations recorded in the Yurtus Fm. carbonates
696 are possibly caused by restricted basin seawater mixing with meteoric waters (Fig. 6). Further,
697 the widespread karst structures may represent a widely distributed hiatus between the
698 Chigebrak and Yurtus Fm. (Wu et al., 2018), which indicates a large scale sea-level drop during
699 the Ediacaran-Cambrian transition and is consistent with a proposed global sea-level shift
700 during this interval (Haq and Schutter, 2008; He et al., 2017). He et al. (2018) studied the
701 geographical distribution of the paleo-karst at the top of the Chigebrak Fm. and speculated
702 that there was an Ediacaran carbonate paleo-upland near to the Xiaerbrak and Shiirik
703 sections during this time. Additionally, the paleo-karst structures in the uppermost of the
704 Chigebrak Fm. may indicate that meteoric fluids (with light O isotopic compositions) and
705 dissolution of Ediacaran carbonates (and their Sr isotope compositions) might have influenced
706 the Yurtus Fm. carbonate Sr isotope values.

707 The obtained $\delta^{13}\text{C}$ values in the Yurtus Fm. carbonates show no variation with $^{87}\text{Sr}/^{86}\text{Sr}$,
708 however, given the existence of a paleo-upland which might contribute to the “great
709 unconformity” at the PC/C boundary (He et al., 2017; He et al., 2018), the negative $\delta^{13}\text{C}$
710 excursions in the Yurtus Fm. may be a result of enhanced dissolved organic carbon (with low
711 $\delta^{13}\text{C}$ signals) remineralization as a result of oceanic ventilation during regression (He et al.,
712 2019).

713 During the Cambrian Series 2, the global seawater $^{87}\text{Sr}/^{86}\text{Sr}$ trend shows a slow rise from
714 ~ 0.7087 to ~ 0.7090 (Kaufman et al., 1993; Derry et al., 1994; Montañez et al., 1996; Montañez
715 and Banner, 2000; Ebner et al., 2001, Fig. 8). This rise may record an increased radiogenic ^{87}Sr
716 input from enhanced continental weathering. Further, increased weathering enhanced organic
717 matter burial, and eventually caused the distinct positive $\delta^{13}\text{C}$ excursion “D”. Finally, organic
718 matter burial could have induced a net increase in oxygen in the deep waters (Guo et al., 2017)
719 which potentially triggered the appearance of *Shizhudiscus* (Fig. 9, Qian and Xiao, 1984), that
720 has higher oxygen demand for its cellular respiration and motile activities.

721

722 **5.4.3 Causes of the ROECE event**

723

724 The prominent global decrease in $^{87}\text{Sr}/^{86}\text{Sr}$ ratios at the boundary between the Cambrian
725 Stage 4 and the Cambrian Wuliuan Stage is commonly explained by one of the following
726 hypotheses:

727 i. Large-scale sea-level rise (Montañez and Banner, 2000; Hough et al., 2006). The flooding
728 of continental areas decreases the input of radiogenic Sr to the oceans and therefore lower
729 the $^{87}\text{Sr}/^{86}\text{Sr}$ ratios of seawater.

730 ii. Widespread rifting along with the Weddell Sea-South African sector of the paleo-Pacific
731 margin of Laurentia (Curtis et al., 1999) exposes mafic rocks (enriched in ^{86}Sr) resulting in a net
732 decrease in radiogenic Sr input.

733 Considering that the negative $\delta^{13}\text{C}$ excursion “F” obtained in the carbonates from the
734 Keping region (down to ~ -13 ‰) is much lower than -6 ‰, we rule out explanation “ii” as
735 hydrothermal fluid CO_2 exhalation of mid-ocean ridge-type (with C isotope compositions
736 usually > -6 ‰; Ishibashi et al., 1995) shows no such negative C isotope compositions. By
737 contrast, the sea-level rise hypothesis represents a more plausible mechanism for the $^{87}\text{Sr}/^{86}\text{Sr}$
738 decrease in the contemporaneous seawater and is supported by the transgression event
739 revealed in the sedimentary characteristics (Bai et al., 2019). Moreover, the extreme negative
740 $\delta^{13}\text{C}$ recorded in the carbonates, which exceeds the magnitude of other ROECE-equivalent
741 negative excursions in South China (Guo et al., 2010), can be hardly explained alone by a
742 reduced dissolved inorganic carbon input into the ocean (Kump and Arthur, 1999). The

743 extreme negative $\delta^{13}\text{C}$ values observed in our studied carbonates are therefore probably
744 superimposed by an upwelling of ^{13}C -depleted deep-waters alongside the transgression,
745 leading to extreme low $\delta^{13}\text{C}$ values in the regional water-mass and the deposited carbonates
746 (Deng et al., 2019; Guo et al., 2017).

747

748 **5.4.4 Cambrian Series 3 (Miaolingian) and Furongian Series paleoenvironmental** 749 **reconstruction of the Tarim Basin**

750

751 During the Cambrian Series 3 (Miaolingian), the ambient seawater $^{87}\text{Sr}/^{86}\text{Sr}$ ratio increases
752 steadily to ~ 0.7092 , while $\delta^{13}\text{C}$ values likewise rise to zero per mil succeeding the excursion "F"
753 (Fig. 9). Following Montañez and Banner (2000), this increase to more radiogenic seawater Sr
754 ratios may have resulted from orogenic events in Antarctica and Australia which is presumably
755 proximal to the Tarim Basin during the Miaolingian (Huang et al., 2000) as they exposed more
756 silicates on the continents prone to physicochemical weathering. This process may have
757 further enhanced the influx of key-nutrients (e.g. phosphorus) into the ocean (Montañez et
758 al., 1996), and ultimately leading to an amplification in the primary productivity and organic
759 carbon burial. As a result, the $\delta^{13}\text{C}$ values of co-precipitating carbonates from surface waters
760 could have increased towards higher values.

761 The beginning of the Furongian is characterized by a peak in both $\delta^{13}\text{C}$ and $^{87}\text{Sr}/^{86}\text{Sr}$ in
762 well-preserved carbonates (e.g. Montañez and Banner, 2000; Ebneith et al., 2001, Fig. 8). In
763 this study, contemporarily precipitated carbonates from the Tarim Basin yield $^{87}\text{Sr}/^{86}\text{Sr}$ ratios
764 between 0.7092 to 0.7091, which are close to modern seawater Sr isotope compositions (i.e.
765 present seawater $^{87}\text{Sr}/^{86}\text{Sr}$: 0.70916 (Ebneith et al., 2001)) and may imply a contribution from
766 a more radiogenic Sr-bearing rocks during the denudation of Damara Belt as part of the Pan-
767 African orogeny (Montañez and Banner, 2000; Ebneith et al., 2001). Moreover, a slight decrease
768 in the $^{87}\text{Sr}/^{86}\text{Sr}$ towards the end of the Furongian in the Keping region as well as in the global
769 record may imply the waning of the Pan-African orogeny (Fig. 9).

770

771 **6. Conclusions**

772

773 This study presents an integrated C and Sr isotope stratigraphic framework from late
774 Ediacaran (ca. 551 Ma) to late Cambrian (< ca. 497 Ma) fully marine strata from the Keping
775 region in the Tarim Basin, North-West China. The C and Sr isotope curves obtained from well-
776 preserved carbonates are similar in their overall trends to those obtained in sections from
777 other carbonate platforms worldwide. A chemo-stratigraphic correlation of our C and Sr
778 isotope curves with globally reported isotope curves from various PC/C sections provides a
779 stratigraphic framework for the late Ediacaran-late Cambrian strata in the Keping area, NW
780 Tarim Basin. Furthermore, the obtained curves may contribute to the global seawater $\delta^{13}\text{C}$ and
781 $^{87}\text{Sr}/^{86}\text{Sr}$ evolution curves and provide a reference for future stratigraphic correlations in the
782 Tarim Basin and elsewhere.

783 Finally, our paleoenvironmental reconstruction shows bio-geochemical variations of the
784 Tarim Basin seawater that were presumably governed by fluctuations in the hinterland
785 weathering regime and temporal changes in the connection to the open ocean. The studied
786 carbonates record both local as well as global seawater level changes, which may have
787 controlled the influx of radiogenic Sr and key nutrients, restricting primary production, into
788 the Tarim Basin. These cause-effect relationships provided in this study may have ultimately
789 impacted the extent of organic C burial and the rapid diversification of metazoans.

790

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792

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801

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1087 **Figure captions:**

1088 **Fig. 1** A) Paleogeographic maps of the Tarim Basin (yellow) and South China (orange)
1089 during the Ediacaran (A-1) and the middle Cambrian (A-2) (modified from Merdith et al., 2017),
1090 abbreviations: T for the Tarim Basin and SC for South China. B) Proterozoic and Phanerozoic
1091 geological units of the Keping region (red square on the inlet map B-1) with locations of the
1092 sections indicated by red stars (modified from Zhou et al., 2015).

1093 **Fig. 2** Comprehensive stratigraphic log of sections from the Keping region (Modified after
1094 Zhou and Chen, 1990; Zhou, 2001; Peng, 2009; Wang et al., 2011b). The markers in the
1095 “Sampling Location” column represent the heights/distances of samples (the height scale of
1096 50 m can be found in the upper-right corner), the Ediacaran samples are from the Shiairik
1097 section (red dashes), and the Cambrian samples are from the Xiaerbrak section (black dashes).
1098 The dotted lines represent less defined formation boundaries; the colored arrows mark the
1099 biozone and fossil occurrence range.

1100 **Fig. 3** Schematic flow diagram of the carbonate dissolution procedure.

1101 **Fig. 4** C, O and Sr isotopic isotope compositions obtained in carbonate leachates from the
1102 late Ediacaran to Cambrian at the studied sections in the Keping region. The green/orange
1103 fields represent positive/negative C isotope excursions, respectively. The inserted column
1104 shows the $\delta^{13}\text{C}$ variation in the Yurtus Fm. in 10-fold height magnification for better visibility.

1105 **Fig. 5** C, O isotope compositions and Mn/Sr ratios for the late-Ediacaran samples (A) and
1106 Cambrian samples (B). The yellow layer represents $\delta^{18}\text{O} < -10\text{‰}$.

1107 **Fig. 6** Sr isotopic composition vs. $\delta^{13}\text{C}$, $\delta^{18}\text{O}$, Sr concentration, Mn/Sr, Mg/Ca, carbonate
1108 content (% = CaO (%) + MgO (%)) and [Rb] for the Cambrian samples (A-G) and late Ediacaran
1109 samples (H-N). For the fluid-carbonate mixing model in Yurtus Fm., a meteoric fluid
1110 composition of $^{87}\text{Sr}/^{86}\text{Sr} = 0.7085$, $\delta^{18}\text{O} = -14$, $\delta^{13}\text{C} = -5$ and Sr concentration of 600 ppm was
1111 chosen together with a presumed localized seawater of $^{87}\text{Sr}/^{86}\text{Sr} = 0.7102$, $\delta^{18}\text{O} = 0.5$, $\delta^{13}\text{C} =$
1112 0.5 , and Sr concentration of 50 ppm was used in the model.

1113 **Fig. 7** Comparison of $\delta^{13}\text{C}$ signatures recorded in carbonate rocks spanning late-Ediacaran
1114 to late-Cambrian strata in A) the Tarim Basin (this study), B) the presumable global Cambrian
1115 seawater standard $\delta^{13}\text{C}$ variations, mainly derived from South China (Zhu et al., 2019), C-D-G-
1116 I) South China (Jiang et al., 2007; Guo et al., 2010; Li et al., 2013), H) South-western Mongolia

1117 (Brasier et al., 1996) E) Laurentia (Montañez and Banner, 2000), F) Siberia (Wotte et al., 2011)
1118 and J) North India (Kaufman et al., 2006). Red dots represent screened samples with severe
1119 diagenetic alteration; the dotted line indicates uncertain boundaries; the red line indicates the
1120 first appearance of trilobites. Paleontological data are derived from Qian and Xiao (1984); Qian
1121 and Bengtson (1989); Peng (2009); Wang et al. (2011b). The age of the beginning of the Paibian
1122 stage is set to 497 Ma (Ogg et al., 2016).

1123 **Fig. 8** Comparison of $^{87}\text{Sr}/^{86}\text{Sr}$ ratios from the late-Ediacaran to the late-Cambrian in A)
1124 Cambrian global seawater $^{87}\text{Sr}/^{86}\text{Sr}$ standard proposed by Peng et al. (2012); Zhu et al. (2019);
1125 of which data is derived from Derry et al. (1994); Montañez and Banner (2000); Ebnet et al.
1126 (2001); Kouchinsky et al. (2008); dotted lines represent less defined formation boundaries, B)
1127 & C) Sr isotope data from the Yangtze Platform, South China (Sawaki et al., 2008; Sawaki et al.,
1128 2010) and D) Keping Area, NW Tarim Basin (this study) where the data from the altered
1129 carbonates from the Yurtus and Sugetbrak Fm. is excluded.

1130 **Fig. 9** C and Sr isotope evolution of seawater in the Keping region, NW Tarim. This figure
1131 includes the stratigraphic correlation and supposed age constraints. Colored (green/orange)
1132 rectangular boxes represent the durations of the $\delta^{13}\text{C}$ excursions. Colored arrows pointed out
1133 the occurrences of the fossil-bearing strata; red dots indicate diagenetically altered $^{87}\text{Sr}/^{86}\text{Sr}$
1134 values and the diagenetically altered $^{87}\text{Sr}/^{86}\text{Sr}$ values from the Yurtus and Sugetbrak
1135 carbonates are excluded in this figure.

1136

Table 1. Analytical results of C, O, Sr isotope compositions and Ca, Mg, Mn, Sr, Rb concentrations of samples from the Keping area, NW Tarim Basin, NW China.

Sample	Formation	Lithology	Height (m)	$\delta^{13}\text{C}$	$\delta^{18}\text{O}$	$^{87}\text{Sr}/^{86}\text{Sr}$	$\pm\sigma$	Ca (%)	Mg (%)	Mn (ppm)	Rb (ppm)	Sr (ppm)	Mn/Sr	Mg/Ca	$^{85}\text{Rb}/^{86}\text{Sr}$
				(V-PDB, ‰)											
SARK-1	Sugetbrak	limestone	-217.9	-5.49	-9.88	0.709471	0.000005	32.4	0.52	2180	1.56	700	3.11	0.02	0.000341
SARK-2	Sugetbrak	limestone	-215.0	-4.68	-9.44					2340	1.94	590	3.96		
SARK-3	Sugetbrak	dolostone	-156.2	1.00	-5.90	0.709365	0.000005	19.7	6.42	781	1.29	171	4.57	0.33	0.001164
SARK-4	Sugetbrak	dolostone	-153.0	3.67	-4.20					436	0.83	160	2.73		
SARK-5	Sugetbrak	dolostone	-151.4	3.90	-3.96					700	4.17	148	4.72		
SARK-6	Sugetbrak	dolostone	-150.5	4.14	-1.83					642	2.13	62.1	10.3		
SARK-7	Sugetbrak	dolostone	-148.5	4.67	-2.53	0.709158	0.000004	13.4	7.39	669	1.60	147	4.55	0.55	0.000280
SARK-8	Sugetbrak	dolostone	-146.0	5.60	-0.85					488	1.06	175	2.78		
SARK-9	Chigebrak	dolostone	-143.0	6.37	-3.02	0.708849	0.000005	15.3	8.64	219	0.69	235	0.93	0.57	0.000458
SARK-10	Chigebrak	dolostone	-142.1	6.61	-3.56					277	0.65	171	1.62		
SARK-11	Chigebrak	dolostone	-141.0	6.56	-3.49	0.709069	0.000005	12.2	6.71	369	1.46	253	1.46	0.55	0.000726
SARK-12	Chigebrak	dolostone	-140.0	6.96	-0.98					205	0.69	201	1.02		
SARK-13	Chigebrak	dolostone	-138.0	7.94	-2.50					182	0.81	227	0.80		
SARK-14	Chigebrak	dolostone	-135.0	5.59	-5.32	0.708934	0.000005	16.3	9.12	499	1.31	141	3.54	0.56	0.001235
SARK-15	Chigebrak	dolostone	-133.5	5.01	-2.41					312	1.23	180	1.73		
SARK-16	Chigebrak	dolostone	-132.0	5.68	-2.78					241	0.57	174	1.38		
SARK-17	Chigebrak	dolostone	-130.5	5.54	-2.60	0.708962	0.000006	15.7	9.16	259	0.74	171	1.51	0.58	0.000738
SARK-18	Chigebrak	dolostone	-129.6	5.93	-2.63					363	1.16	207	1.76		
SARK-19	Chigebrak	dolostone	-129.0	5.69	-2.36					243	0.62	203	1.20		
SARK-20	Chigebrak	dolostone	-127.0	4.87	-3.65	0.708994	0.000005	13.4	7.83	402	1.67	226	1.78	0.58	0.000579
SARK-21	Chigebrak	dolostone	-125.2	5.88	-2.16					279	0.65	180	1.55		
SARK-22	Chigebrak	dolostone	-124.2	5.93	-2.58					218	0.78	176	1.24		
SARK-23	Chigebrak	dolostone	-122.0	4.41	-0.72	0.708806	0.000005	12.6	7.27	410	1.23	191	2.15	0.58	0.000467
SARK-24	Chigebrak	dolostone	-119.0	4.40	-2.30					297	1.42	115	2.58		
SARK-25	Chigebrak	dolostone	-117.2	4.12	0.15					222	0.77	142	1.56		

Sample	Formation	Lithology	Height (m)	$\delta^{13}\text{C}$	$\delta^{18}\text{O}$	$^{87}\text{Sr}/^{86}\text{Sr}$	$\pm\sigma$	Ca (%)	Mg (%)	Mn (ppm)	Rb (ppm)	Sr (ppm)	Mn/Sr	Mg/Ca	$^{85}\text{Rb}/^{86}\text{Sr}$
				(V-PDB, ‰)											
SARK-26	Chigebrak	dolostone	-114.0	3.50	-2.80	0.708816	0.000004	17.5	10.1	190	0.69	158	1.20	0.58	0.000260
SARK-27	Chigebrak	dolostone	-111.0	3.63	-2.47					156	0.72	220	0.71		
SARK-28	Chigebrak	dolostone	-109.0	2.06	-3.56					152	0.54	170	0.89		
SARK-29	Chigebrak	dolostone	-107.0	0.97	-3.10	0.708750	0.000005	16.8	9.80	149	0.54	169	0.88	0.58	0.000474
SARK-30	Chigebrak	dolostone	-100.3	1.70	-2.87	0.708611	0.000005	18.9	11.1	207	0.51	104	1.99	0.59	0.000455
SARK-31	Chigebrak	dolostone	-99.6	1.69	-2.98					18.7	0.45				
SARK-32	Chigebrak	dolostone	-98.0	2.35	-3.11					199	0.50	161	1.23		
SARK-33	Chigebrak	dolostone	-95.0	2.57	-3.12					108	0.96	73.5	1.47		
SARK-34	Chigebrak	dolostone	-90.0	2.05	-5.08	0.708746	0.000005	17.9	10.1	396	0.68	127	3.11	0.57	0.000641
SARK-35	Chigebrak	dolostone	-86.0	2.13	-5.16					464	0.47	144	3.23		
SARK-36	Chigebrak	dolostone	-83.0	2.52	-1.52					256	0.44	130	1.96		
SARK-37	Chigebrak	dolostone	-80.0	3.17	-2.45					165	0.44	123	1.34		
SARK-38	Chigebrak	dolostone	-75.0	3.05	-1.04	0.708464	0.000005	19.8	11.5	173	0.44	138	1.25	0.58	0.000689
SARK-39	Chigebrak	dolostone	-66.0	2.70	-2.10					179	0.43	163	1.10		
SARK-40	Chigebrak	dolostone	-60.0	2.75	-0.10					177	0.47	440	0.40		
SARK-41	Chigebrak	dolostone	-56.0	2.86	-0.05	0.708544	0.000005	19	11.0	229	0.50	125	1.83	0.58	0.000469
SARK-42	Chigebrak	dolostone	-53.0	3.22	0.42					110	0.45	107	1.03		
SARK-43	Chigebrak	dolostone	-48.2	2.88	-1.91					136	0.61	78.8	1.73		
SARK-44	Chigebrak	dolostone	-45.0	2.98	-0.91	0.708775	0.000005	18.2	10.5	155	0.61	116	1.33	0.58	0.001286
SARK-45	Chigebrak	dolostone	-40.0	2.75	-1.78					185	0.59	104	1.78		
SARK-46	Chigebrak	dolostone	-34.5	2.33	-3.92					336	0.55	94.2	3.57		
SARK-47	Chigebrak	dolostone	-6.0	1.97	-5.54	0.708880	0.000005	13.3	7.70	557	0.94	143	3.90	0.58	0.000579
SARK-48	Chigebrak	dolostone	-3.0	1.65	-6.49	0.709089	0.000004	14.6	8.20	604	1.98	208	2.91	0.56	0.000610
SARK-49	Chigebrak	dolostone	-2.0	1.98	-5.97					546	0.92	167	3.26		
XBK-1	Yurtus	phosphorite	0.5							1988	1.31	386	5.15		
XBK-2	Yurtus	phosphorite	2.5			0.709950	0.000008	1.04	0.19	0.36	0.25	58.4	0.01	0.18	0.000483

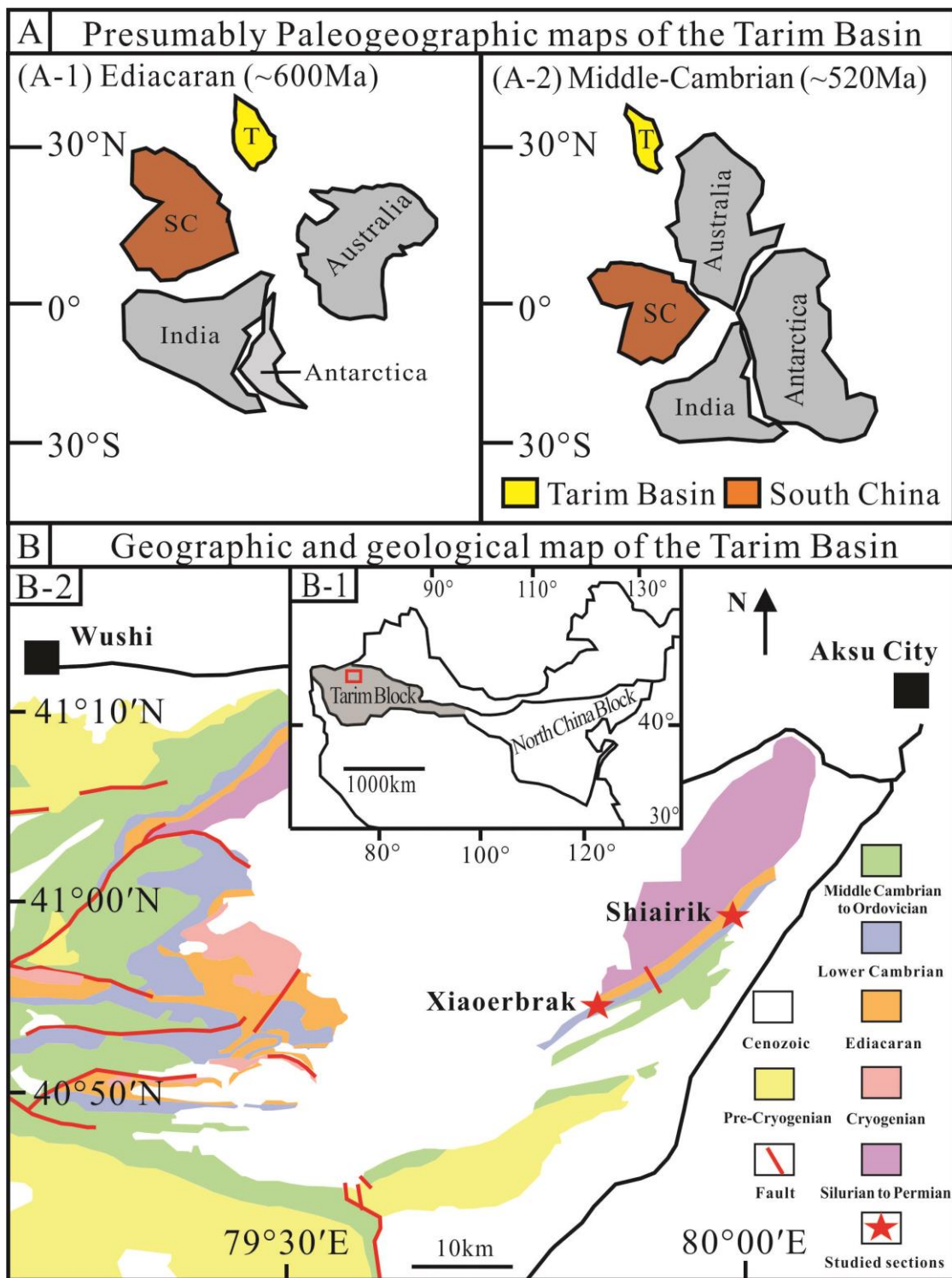
Sample	Formation	Lithology	Height (m)	$\delta^{13}\text{C}$	$\delta^{18}\text{O}$	$^{87}\text{Sr}/^{86}\text{Sr}$	$\pm\sigma$	Ca (%)	Mg (%)	Mn (ppm)	Rb (ppm)	Sr (ppm)	Mn/Sr	Mg/Ca	$^{85}\text{Rb}/^{86}\text{Sr}$
				(V-PDB, ‰)											
XBK-3	Yurtus	phosphorite	4.5			0.708852	0.000008	1.33	0.55	1.50	0.13	578		0.41	0.000917
XBK-4	Yurtus	dolostone	6.5	-2.63	-6.32	0.709560	0.000007	34.6	17.7	415	2.30	362	1.15	0.51	0.000787
XBK-5	Yurtus	limestone	8.5	-0.58	-12.1					344	0.34	206	1.67		
XBK-6	Yurtus	limestone	9.0	0.46	-13.2	0.708695	0.000011	36.2	0.20	213	0.43	123	1.74	0.01	0.000400
XBK-7	Yurtus	dolostone	9.3	-5.05	-9.20					496	0.35	103	4.84		
XBK-8	Yurtus	dolostone	10.2	-1.90	-7.77	0.709264	0.000009	2.74	0.75	44.9	0.56	63.6	0.70	0.27	0.000506
XBK-9	Yurtus	dolostone	11.0	-0.77	-6.45	0.710168	0.000009	3.27	1.63	121	0.70	24.2	4.98	0.50	0.001347
XBK-10	Yurtus	dolostone	12.0	-1.12	-9.94					950	0.45	68.9	13.8		
XBK-11	Yurtus	dolostone	13.0	-0.05	-8.33					935	0.40	46.9	19.9		
XBK-12	Yurtus	dolostone	13.5	0.30	-8.22					881	0.32	50.2	17.6		
XBK-13	Yurtus	dolostone	14.2	0.73	-7.67	0.709472	0.000009	18.3	10.2	1062	0.37	73.8	14.4	0.56	0.000990
XBK-14	Yurtus	dolostone	15.6	0.61	-6.85					0.10	0.00				
XBK-15	Yurtus	dolostone	17.8	0.30	-7.06					1017	0.34	50.0	20.3		
XBK-16	Yurtus	dolostone	18.7	0.22	-6.99					902	0.33	44.9	20.1		
XBK-17	Yurtus	dolostone	19.5	-0.12	-7.18					966	0.13	43.6	22.1		
XBK-18	Yurtus	dolostone	21.5	-0.53	-7.11					1428	0.34	41.9	34.1		
XBK-19	Yurtus	dolostone	21.5	0.21	-7.53	0.709451	0.000009	18.4	10.8	472	0.43	58.5	8.07	0.59	0.000861
XBK-20	Yurtus	dolostone	21.6	-0.04	-7.73					361	0.08	44.8	8.07		
XBK-21	Xiaoerbrak	dolostone	21.7	0.08	-7.34					277	0.15	53.6	5.16		
XBK-22	Xiaoerbrak	dolostone	24.9	1.21	-7.7	0.708971	0.000009	21.8	12.5	250	0.11	71.7	3.49	0.57	0.000793
XBK-23	Xiaoerbrak	dolostone	25.9	1.26	-7.52					172	0.03	75.9	2.27		
XBK-24	Xiaoerbrak	dolostone	31.6	2.20	-6.79	0.708817	0.000011	23.8	13.9	136		91.4	1.49	0.58	0.000392
XBK-25	Xiaoerbrak	dolostone	33.4	2.54	-7.05					147		89.2	1.65		
XBK-26	Xiaoerbrak	dolostone	36.1	1.73	-7.39					129		67.4	1.92		
XBK-27	Xiaoerbrak	dolostone	41.4	2.78	-7.26	0.708790	0.000009	19.0	11.3	142		81.7	1.73	0.59	0.001545
XBK-28	Xiaoerbrak	dolostone	45.9	2.59	-6.71					139		92.2	1.51		

Sample	Formation	Lithology	Height (m)	$\delta^{13}\text{C}$	$\delta^{18}\text{O}$	$^{87}\text{Sr}/^{86}\text{Sr}$	$\pm\sigma$	Ca (%)	Mg (%)	Mn (ppm)	Rb (ppm)	Sr (ppm)	Mn/Sr	Mg/Ca	$^{85}\text{Rb}/^{86}\text{Sr}$
				(V-PDB, ‰)											
XBK-29	Xiaoerbrak	dolostone	52.4	2.55	-6.97	0.708765	0.000009	21.9	12.9	117		73.7	1.58	0.59	0.000309
XBK-30	Xiaoerbrak	dolostone	59.5	2.69	-7.32					124		67.3	1.84		
XBK-31	Xiaoerbrak	dolostone	63.1	2.99	-7.33	0.708731	0.000009	21.2	12.5	132	0.09	87.7	1.50	0.59	0.000555
XBK-32	Xiaoerbrak	dolostone	63.3	2.89	-7.23					111		73.2	1.51		
XBK-33	Xiaoerbrak	dolostone	65.4	3.18	-7.59					96.0		83.9	1.14		
XBK-34	Xiaoerbrak	dolostone	71.6	3.12	-6.94	0.708733	0.000009	20.8	12.2	122		112	1.08	0.58	0.000373
XBK-35	Xiaoerbrak	dolostone	75.9	3.17	-7.24					101		74.5	1.36		
XBK-36	Xiaoerbrak	dolostone	80.3	2.87	-8.49					112		55.5	2.02		
XBK-37	Xiaoerbrak	dolostone	85.5	2.96	-7.96	0.708803	0.000008	21.4	12.8	124		65.2	1.91	0.59	0.000649
XBK-38	Xiaoerbrak	dolostone	90.3	2.66	-8.78					141		64.4	2.18		
XBK-39	Xiaoerbrak	dolostone	103.7	2.73	-8.27					134		53.9	2.49		
XBK-40	Xiaoerbrak	dolostone	113.7	3.15	-7.12	0.708817	0.000015	20.0	11.8	82.8		45.3	1.83	0.59	0.000562
XBK-41	Xiaoerbrak	dolostone	118.0	2.70	-8.24					109	0.03	43.8	2.48		
XBK-42	Xiaoerbrak	dolostone	123.5	2.47	-7.32	0.708888	0.00001	20.6	11.6	91.1		47.7	1.91	0.56	0.000990
XBK-43	Xiaoerbrak	dolostone	126.5	1.73	-7.21					95.8		36.5	2.62		
XBK-44	Xiaoerbrak	dolostone	132.6	2.07	-6.90	0.708810	0.000009	19.5	11.5	92.5		43	2.15	0.59	0.000599
XBK-45	Xiaoerbrak	dolostone	135.6	2.66	-6.44					72.6		30.9	2.35		
XBK-46	Xiaoerbrak	dolostone	142.6	2.85	-6.41					95.4		35.2	2.71		
XBK-47	Xiaoerbrak	dolostone	145.0	2.86	-6.45	0.708788	0.000009	17.8	10.6	86.7		36.0	2.41	0.60	0.000567
XBK-48	Xiaoerbrak	dolostone	149.4	2.33	-6.81					120		39.8	3.02		
XBK-49	Xiaoerbrak	dolostone	151.6	2.43	-6.44					112		39.5	2.83		
XBK-50	Xiaoerbrak	dolostone	155.5	2.45	-6.49	0.708795	0.000009	20.3	12.0	114		57.5	1.98	0.59	0.000615
XBK-51	Xiaoerbrak	dolostone	159.8	2.46	-6.74					104	0.10	51.9	2.00		
XBK-52	Xiaoerbrak	dolostone	165.3	2.51	-6.45					116	0.06	64.1	1.82		
XBK-53	Xiaoerbrak	dolostone	166.0	2.54	-5.50	0.708831	0.000008	19.7	11.5	98.9	0.11	67.0	1.48	0.59	0.00033
XBK-54	Xiaoerbrak	dolostone	167.5	2.42	-5.60					88.4	0.02	64.6	1.37		

Sample	Formation	Lithology	Height (m)	$\delta^{13}\text{C}$	$\delta^{18}\text{O}$	$^{87}\text{Sr}/^{86}\text{Sr}$	$\pm\sigma$	Ca (%)	Mg (%)	Mn (ppm)	Rb (ppm)	Sr (ppm)	Mn/Sr	Mg/Ca	$^{85}\text{Rb}/^{86}\text{Sr}$
				(V-PDB, ‰)											
XBK-55	Xiaoerbrak	dolostone	168.9	2.50	-5.61	0.708827	0.00001	5.53	3.01	14.8		24.7	0.60	0.54	0.000830
XBK-56	Xiaoerbrak	dolostone	170.1	2.25	-3.53	0.708796	0.000008	17.3	10.1	79.3	0.12	86.1	0.92	0.58	0.000534
XBK-57	Xiaoerbrak	dolostone	171.7	1.22	-4.56					111	0.09	91.5	1.21		
XBK-58	Xiaoerbrak	dolostone	174.1	0.71	-6.7	0.708962	0.000009	18.8	10.7	135	0.01	62.2	2.17	0.57	0.000761
XBK-59	Xiaoerbrak	dolostone	174.0	1.08	-6.12					139	0.00	60.4	2.31		
XBK-60	Xiaoerbrak	dolostone	176.8	-0.69	-6.33	0.708989	0.000008	17.9	10.3	137	0.14	69.0	1.99	0.58	0.000492
XBK-61	Xiaoerbrak	dolostone	178.2	0.19	-8.09					157	0.32	112	1.40		
XBK-62	Xiaoerbrak	dolostone	179.0	-0.19	-6.05	0.708990	0.000008	19.9	11.3	120	0.04	66.1	1.81	0.57	0.000509
XBK-63	Xiaoerbrak	dolostone	182.3	0.33	-6.08					151	0.19	81.9	1.84		
XBK-64	Xiaoerbrak	dolostone	186.7	0.34	-6.12	0.709057	0.00001	19.3	11.1	110	0.14	77.8	1.41	0.58	0.000517
XBK-65	Xiaoerbrak	dolostone	188.9	-0.14	-6.36					143	0.14	68.7	2.09		
XBK-66	Xiaoerbrak	dolostone	192.5	0.62	-6.10	0.708863	0.00001	18.5	10.4	139	0.03	89.6	1.55	0.56	0.000506
XBK-67	Wusonger	dolostone	196.1	-1.05	-6.47					241	0.19	72.3	3.34		
XBK-68	Wusonger	dolostone	202.5	-1.77	-5.80					323	1.73	106	3.04		
XBK-69	Wusonger	dolostone	210.1	-1.21	-6.44	0.708902	0.000007	16.8	9.45	260	0.21	81.9	3.17	0.56	0.000587
XBK-70	Wusonger	dolostone	213.2	-1.89	-6.33					214	0.94	77.9	2.75		
XBK-71	Wusonger	dolostone	244.7	-1.71	-5.14	0.708934	0.000008	13.5	7.88	207	1.17	97.9	2.11	0.58	0.000747
XBK-72	Wusonger	dolostone	254.7	-1.11	-5.47	0.708980	0.000007	19.1	10.9	273	1.36	209	1.30	0.57	0.000628
XBK-73	Wusonger	dolostone	256.2	-0.21	-5.54	0.708941	0.000007	19.8	10.8	298	0.95	219	1.36	0.55	0.000425
XBK-74	Wusonger	dolostone	257.7	-0.27	-5.76	0.708974	0.000007	20.4	11.8	243	0.92	257	0.94	0.58	0.000378
XBK-75	Wusonger	dolostone	259.2	-0.54	-5.84	0.709179	0.000008	18.1	10.2	335	1.26	203	1.65	0.56	0.000242
XBK-76	Wusonger	dolostone	260.7	-0.31	-5.67	0.709054	0.000008	19.3	10.4	295	1.19	220	1.34	0.54	0.000378
XBK-77	Wusonger	dolostone	262.2	0.98	-5.27	0.709169	0.000009	18.5	10.8	276	0.99	213	1.29	0.58	0.000508
XBK-78	Wusonger	dolostone	263.7	-6.66	-12.9	0.709150	0.000006	2.32	0.15	57.9	3.50	1213	0.05	0.06	0.000209
XBK-79	Wusonger	dolostone	265.2	-0.67	-5.59					314	1.13	343	0.92		
XBK-80	Wusonger	dolostone	266.7	-1.18	-5.80					276	0.96	240	1.15		

Sample	Formation	Lithology	Height (m)	$\delta^{13}\text{C}$	$\delta^{18}\text{O}$	$^{87}\text{Sr}/^{86}\text{Sr}$	$\pm\sigma$	Ca (%)	Mg (%)	Mn (ppm)	Rb (ppm)	Sr (ppm)	Mn/Sr	Mg/Ca	$^{85}\text{Rb}/^{86}\text{Sr}$
				(V-PDB, ‰)											
XBK-81	Wusonger	dolostone	268.2	-3.65	-5.48	0.708749	0.000008	20.0	11.4	265	0.74	209	1.27	0.57	0.000483
XBK-82	Wusonger	dolostone	269.7	-2.23	-5.77					281	1.16	214	1.31		
XBK-83	Wusonger	dolostone	271.2	-11.7	-5.48	0.708771	0.000008	18.8	10.6	352	0.89	248	1.42	0.56	0.000609
XBK-84	Shayilik	dolostone	273.0	-13.3	-7.75					471	1.48	184	2.56		
XBK-85	Shayilik	dolostone	277.9	-2.60	-6.17	0.708923	0.000009	20.3	9.50	320	0.96	213	1.50	0.47	0.001105
XBK-86	Shayilik	dolostone	285.6	-2.45	-4.81			13.6	7.90	196	2.70	153	1.28	0.58	
XBK-87	Shayilik	dolostone	301.7	-2.44	-6.37	0.708723	0.000008	13.6	7.80	287	1.80	128	2.24	0.57	0.000928
XBK-88	Shayilik	dolostone	308.7	-1.35	-6.98					614	2.52	131	4.68		
XBK-89	Shayilik	limestone	326.2	-0.35	-10.5	0.709021	0.000009	35.3	0.40	330	0.99	615	0.54	0.01	0.000329
XBK-90	Shayilik	limestone	336.4	-0.65	-8.05					217	0.64	201	1.08		
XBK-91	Shayilik	limestone	340.0	-0.08	-8.13	0.708853	0.000010	30.2	3.40	192	0.52	570	0.34	0.11	0.000260
XBK-92	Shayilik	limestone	350.4	-0.61	-8.47					369	0.81	425	0.87		
XBK-93	Awatagh	limestone	369.1	0.67	-5.94					598	3.16	130	4.60		
XBK-94	Awatagh	limestone	371.1	0.97	-8.59	0.708880	0.000007	36.3	0.50	135	0.61	5906	0.02	0.01	0.000746
XBK-95	Awatagh	dolostone	386.0	-0.66	-5.76	0.708892	0.000007	12.0	6.80	484	2.14	144	3.37	0.56	0.000542
XBK-96	Awatagh	dolostone	421.5	0.16	-8.81	0.708988	0.000007	19.2	6.50	1085	0.69	183	5.93	0.34	0.000628
XBK-97	Awatagh	dolostone	449.4	-1.50	-7.03					1109	1.21	180	6.17		
XBK-98	Awatagh	dolostone	449.6	-0.25	-6.47	0.709189	0.000008	15.2	7.10	545	0.70	287	1.90	0.47	0.000849
XBK-99	Awatagh	dolostone	468.1	-0.54	-7.47					82.1	0.55	38.9	2.11		
XBK-100	Awatagh	dolostone	483.0	0.70	-6.31	0.709083	0.000009	11.6	6.20	493	4.76	222	2.22	0.53	0.000347
XBK-101	Awatagh	dolostone	483.0	1.11	-6.82					482	1.15	182	2.65		
XBK-102	Awatagh	dolostone	485.0	1.34	-6.45					284	1.05	268	1.06		
XBK-103	Awatagh	dolostone	498.9	-1.38	-7.07	0.709053	0.000007	14.8	8.20	169	1.31	157	1.07	0.56	0.000289
XBK-104	Awatagh	dolostone	500.2	0.06	-6.94					223	1.56	95.5	2.33		
XBK-105	Awatagh	dolostone	517.6	0.09	-6.63	0.709134	0.000008	17.2	9.60	239	1.30	112	2.14	0.56	0.000610
XBK-106	Awatagh	dolostone	519.0	-1.56	-8.69					159	0.80	81.9	1.94		

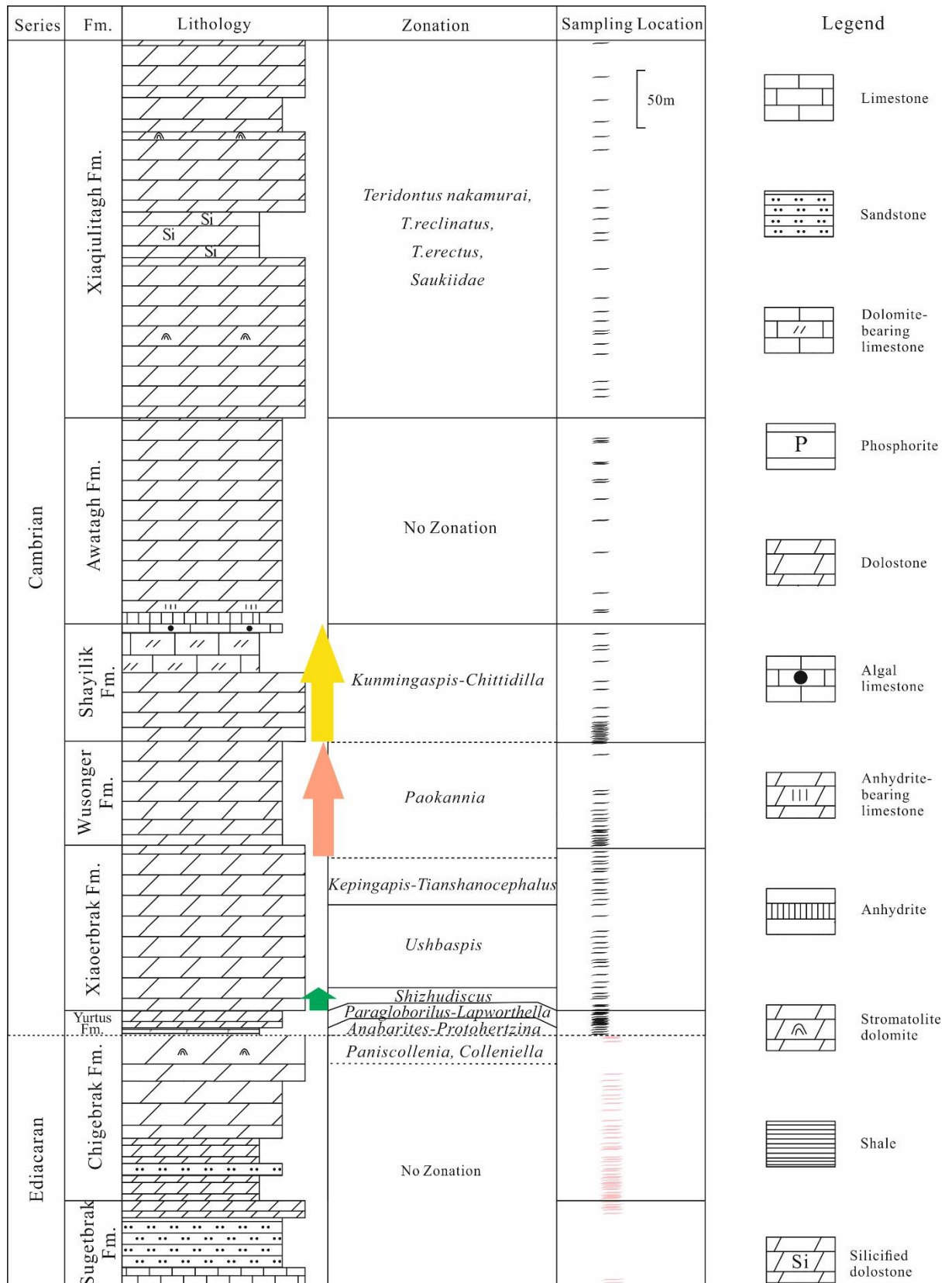
Sample	Formation	Lithology	Height (m)	$\delta^{13}\text{C}$	$\delta^{18}\text{O}$	$^{87}\text{Sr}/^{86}\text{Sr}$	$\pm\sigma$	Ca (%)	Mg (%)	Mn (ppm)	Rb (ppm)	Sr (ppm)	Mn/Sr	Mg/Ca	$^{85}\text{Rb}/^{86}\text{Sr}$
				(V-PDB, ‰)											
XBK-107	Awatagh	dolostone	521.2	-0.58	-6.61					249	1.16	183	1.36		
XBK-108	Xiaquilitagh	dolostone	557.6	-0.41	-6.37	0.709156	0.000011	19.2	11.0	106	0.82	119	0.88	0.57	0.000381
XBK-109	Xiaquilitagh	dolostone	563.9	-0.05	-7.22					149	0.76	106	1.40		
XBK-110	Xiaquilitagh	dolostone	570.8	-0.08	-7.38					105	0.69	76.2	1.38		
XBK-111	Xiaquilitagh	dolostone	594.8	-1.30	-5.04	0.708608	0.000008	19.8	11.2	96.7	0.76	91.3	1.06	0.57	0.000679
XBK-112	Xiaquilitagh	dolostone	603.7	-1.37	-4.43					104	1.12	90.7	1.14		
XBK-113	Xiaquilitagh	dolostone	612.7	-0.96	-6.23	0.709018	0.000008	19.1	10.9	68.2	0.67	116	0.59	0.57	0.000277
XBK-114	Xiaquilitagh	dolostone	615.1	-1.22	-6.50					83	0.64	141	0.59		
XBK-115	Xiaquilitagh	dolostone	623.7	-1.12	-6.32	0.709107	0.000009	20.4	9.80	84.3	1.08	126	0.67	0.48	0.000181
XBK-116	Xiaquilitagh	dolostone	631.9	-1.08	-6.25					64.5	0.74	140	0.46		
XBK-117	Xiaquilitagh	dolostone	643.7	-1.16	-6.10	0.709114	0.000008	19.2	10.7	57.5	0.69	118	0.49	0.56	0.000546
XBK-118	Xiaquilitagh	dolostone	669.2	-0.99	-6.16					65.1	0.79	115	0.57		
XBK-119	Xiaquilitagh	dolostone	694.3	-0.87	-6.55	0.709087	0.000008	15.1	8.10	73.5	0.55	100	0.73	0.54	0.000312
XBK-120	Xiaquilitagh	dolostone	700.8	-1.04	-6.16					88.9	0.72	130	0.69		
XBK-121	Xiaquilitagh	dolostone	712.9	-0.91	-6.37	0.709127	0.000008	16.5	9.00	55.1	0.71	151	0.36	0.54	0.000257
XBK-122	Xiaquilitagh	dolostone	722.6	-1.18	-6.70					59.3	1.10	146	0.41		
XBK-123	Xiaquilitagh	dolostone	738.2	-0.96	-6.37	0.709138	0.000008	18.1	10.0	104	0.80	204	0.51	0.56	0.000450
XBK-124	Xiaquilitagh	dolostone	773.2	-1.17	-6.48					118	0.74	227	0.52		
XBK-125	Xiaquilitagh	dolostone	784.9	-0.97	-6.58	0.709143	0.000009	20.3	11.4	120	0.81	245	0.49	0.56	0.000230
XBK-126	Xiaquilitagh	dolostone	797.9	-0.86	-6.49					69.5	0.77	150	0.46		
XBK-127	Xiaquilitagh	dolostone	816.8	-0.82	-7.17	0.709195	0.000008	21.1	10.4	62.8	0.81	126	0.50	0.49	0.000360
XBK-128	Xiaquilitagh	dolostone	837.0	-0.52	-5.68					73.6	0.77	129	0.57		
XBK-129	Xiaquilitagh	dolostone	870.7	-0.29	-6.03	0.709107	0.000009	18.4	10.1	88.3	0.58	136	0.65	0.55	0.000317



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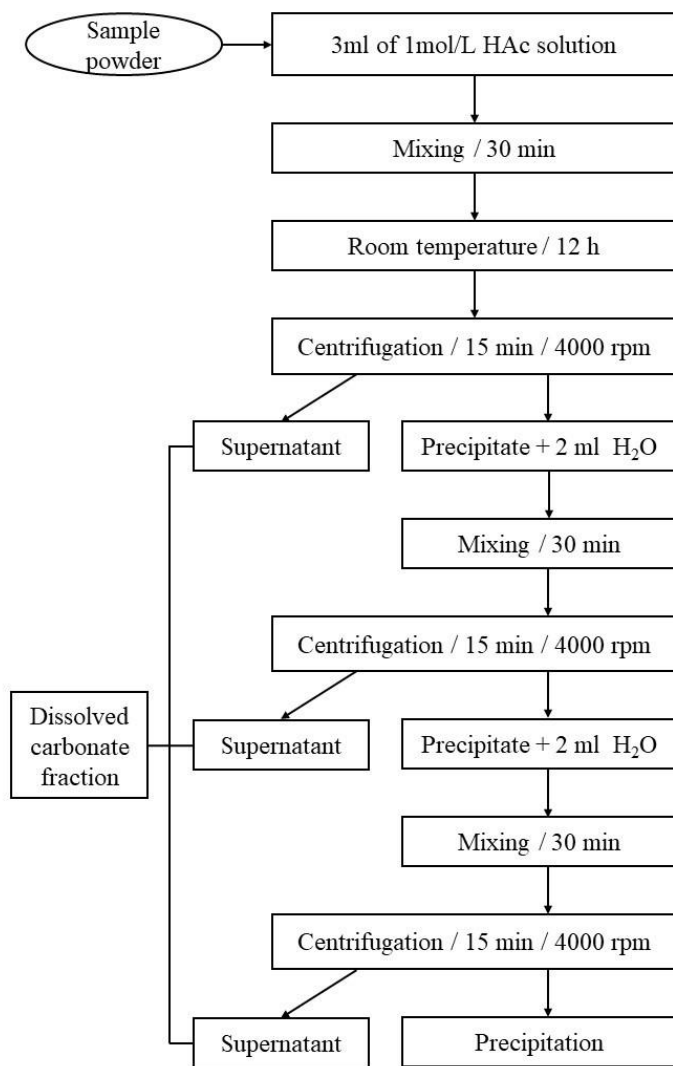
Figure 1



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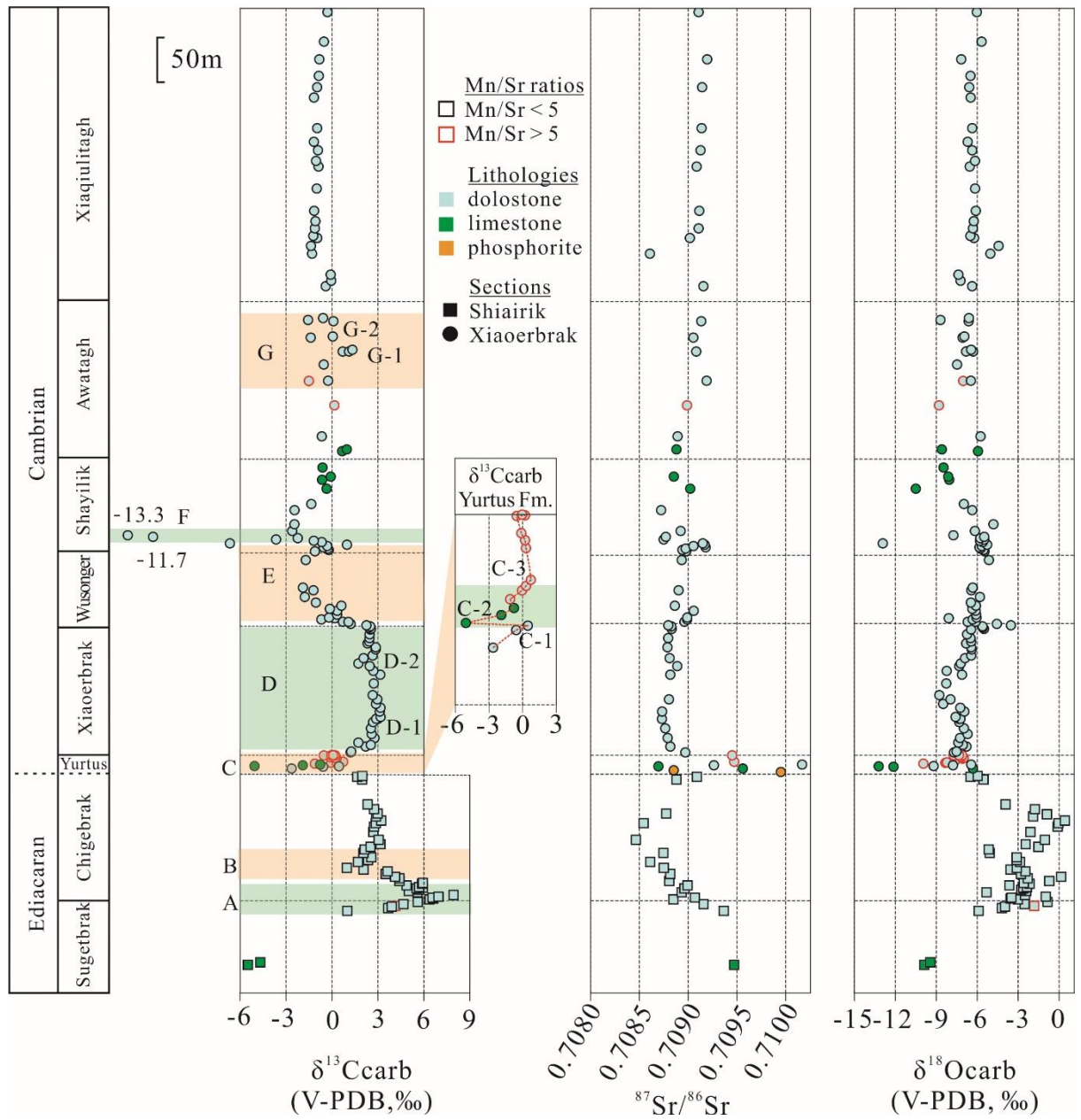
Figure 2



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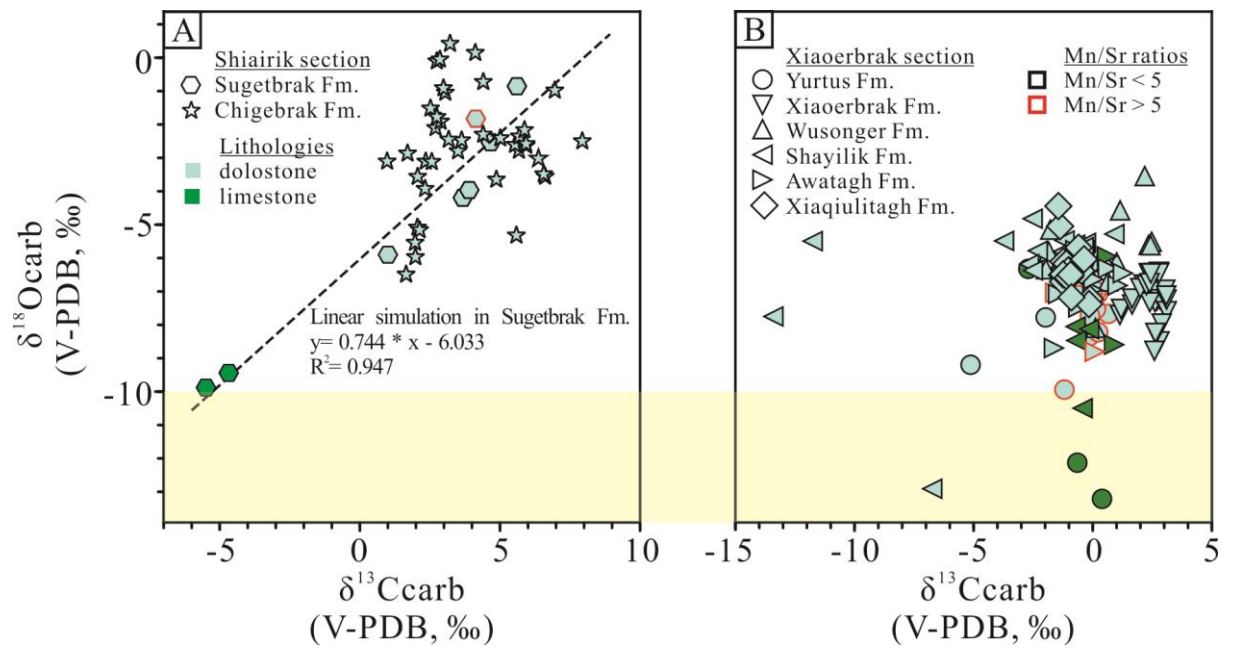
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Figure 3



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1146 Figure 4



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Figure 5

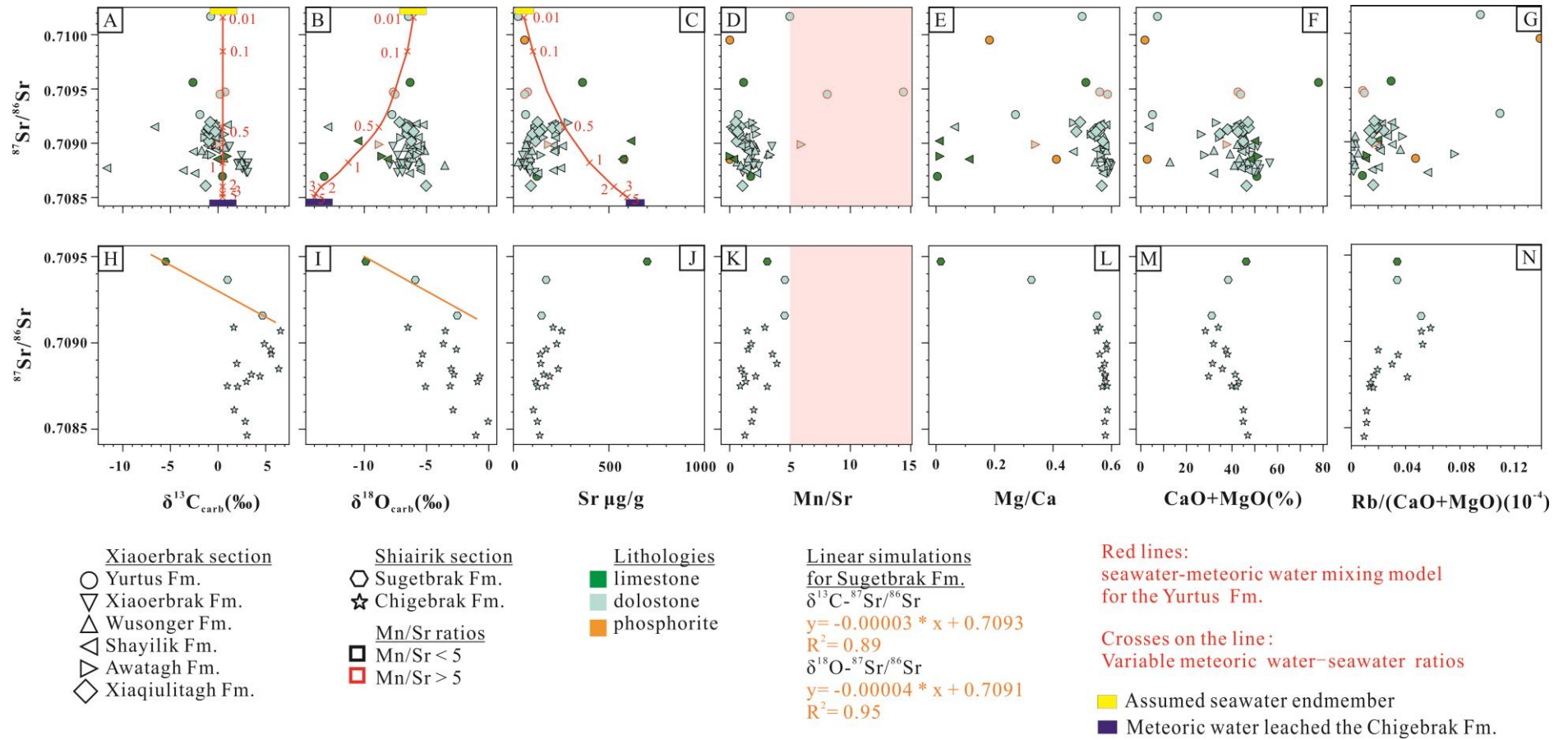
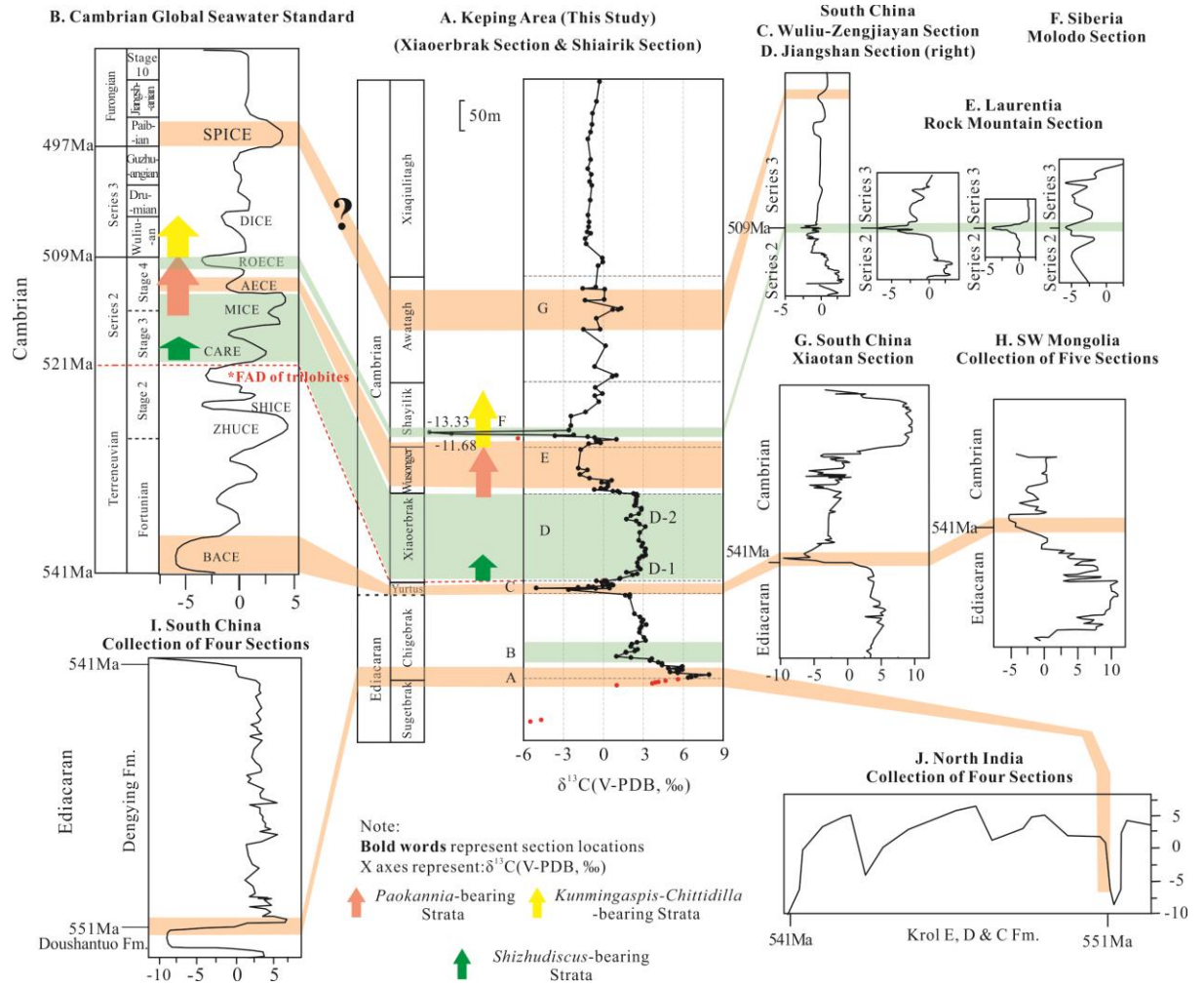


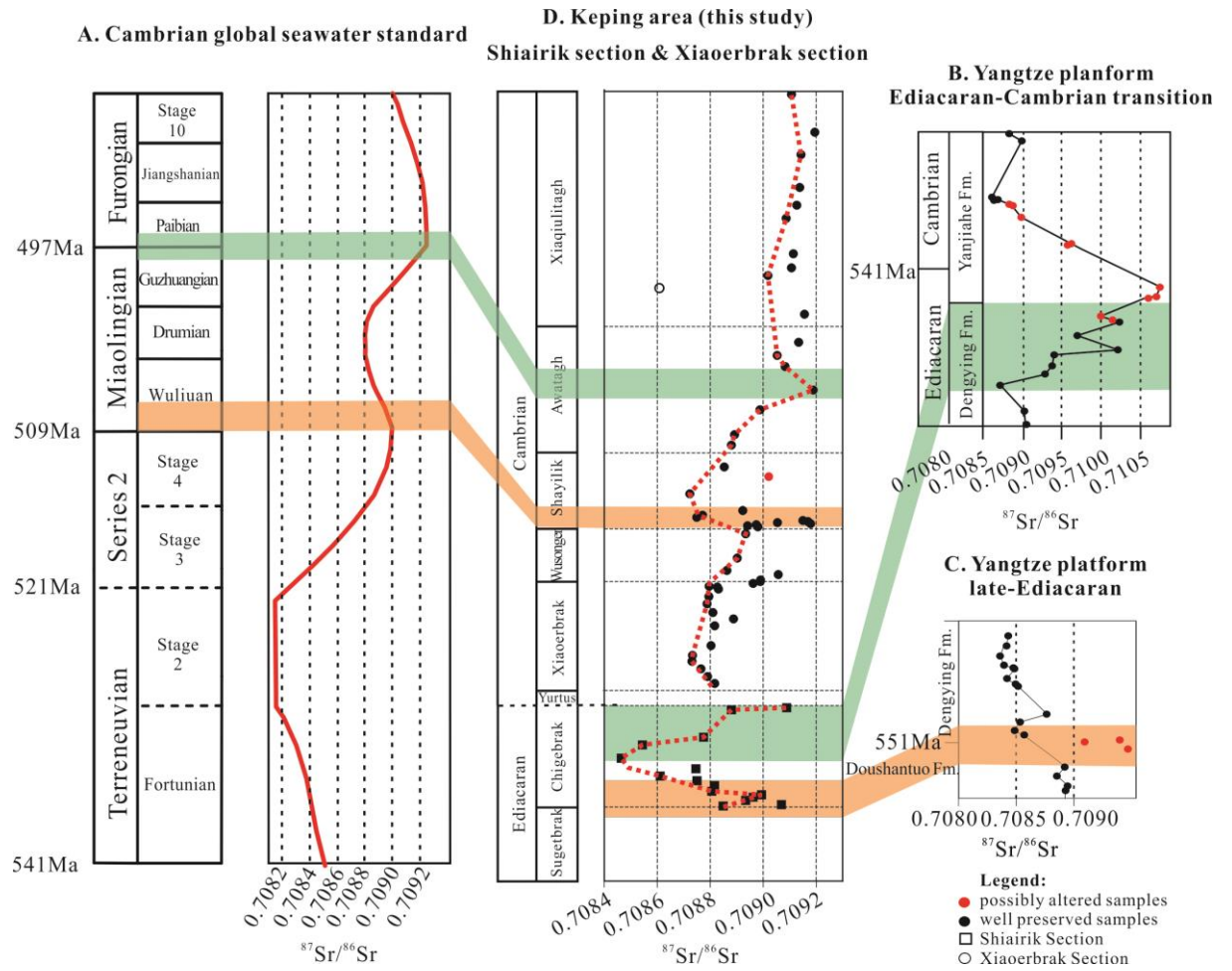
Figure 6



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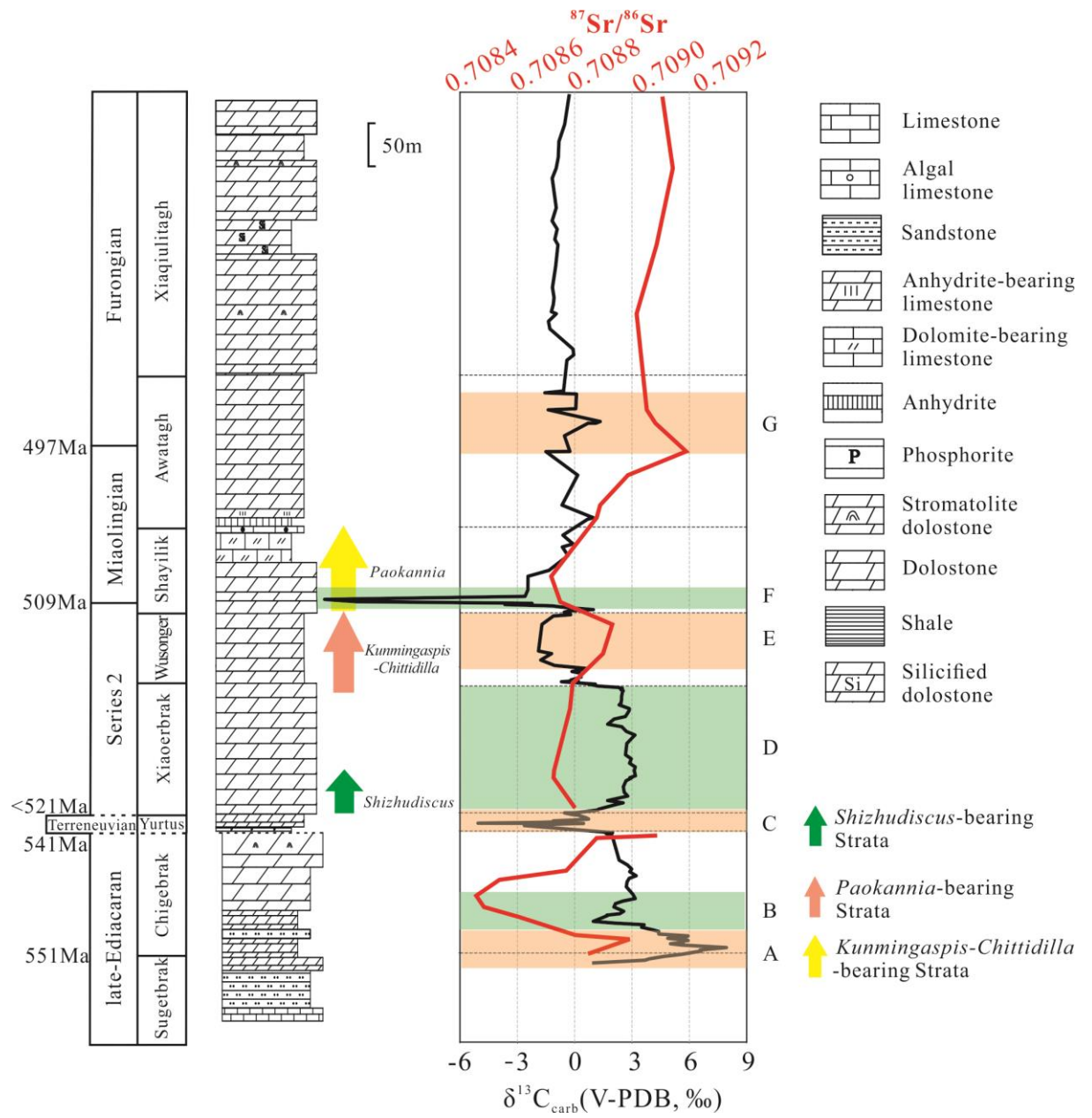
Figure 7



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Figure 8



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1157 Figure 9