

1 **Fingerprinting fluid source in calcite veins: combining LA-ICP-MS U-Pb calcite dating with**
2 **trace elements and clumped isotope palaeothermometry**

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58 ABSTRACT

59 Application of geochemical proxies to vein minerals - particularly calcite - can fingerprint the source
60 of fluids controlling various important geological processes from seismicity to geothermal systems.
61 Determining fluid source, e.g. meteoric, marine, magmatic or metamorphic waters, can be
62 challenging when using only trace elements and stable isotopes as different fluids can have
63 overlapping geochemical characteristics, such as $\delta^{18}\text{O}$. In this contribution we show that by
64 combining the recently developed LA-ICP-MS U-Pb calcite geochronometer with stable isotopes
65 (including clumped isotope palaeothermometry) and trace element analysis, the fluid source of
66 veins can be more readily determined. Calcite veins hosted in the Devonian Montrose Volcanic
67 Formation at Lunan Bay in the Midland Valley Terrane of Central Scotland were used as a case
68 study. δD values of fluid inclusions in the calcite, and parent fluid $\delta^{18}\text{O}$ values reconstructed from
69 clumped isotope palaeothermometry, gave values which could represent a range of fluid sources:
70 metamorphic or magmatic fluids, or surface waters which had undergone much fluid-rock
71 interaction. Trace elements showed no distinctive patterns and shed no further light on fluid
72 source. LA-ICP-MS U-Pb dating determined the vein calcite precipitation age – 318 ± 30 Ma – which
73 rule out metamorphic or magmatic fluid sources as no metamorphic or magmatic activity was
74 occurring in the area at this time. The vein fluid source was therefore a surface water (meteoric
75 based on paleogeographic reconstruction) which had undergone significant water-rock interaction.
76 This study highlights the importance of combining the recently developed LA-ICP-MS U-Pb calcite
77 geochronometer with stable isotopes and trace elements to help determine fluid sources of veins,
78 and indeed any geological feature where calcite precipitated from a fluid that may have resided in
79 the crust for a period of time (e.g. fault precipitates or cements).

80
81 Keywords: LA-ICP-MS U-Pb calcite geochronology; clumped isotopes; fluid δD and $\delta^{18}\text{O}$; calcite
82 veins; trace elements.

83
84 INTRODUCTION

85
86 Fingerprinting the source of fluids flowing through fractures in the crust has importance in a range
87 of geological applications, including: 1) understanding the origin, and predicting sustainability, of
88 geothermal systems (e.g. Simmons and Christenson 1994; Menzies et al. 2014; Lu et al. 2017; Lu
89 et al. 2018); 2) determining the origin and concentration of economic mineral deposits (e.g. Barker
90 and Cox 2011); and 3) reconstructing fluid flow pathways responsible for seismicity (e.g. Uysal et
91 al. 2011; Sturrock et al. 2017). Evidence of fluid flow through fractures is recorded by the presence
92 of veins and application of geochemical proxies to vein minerals - particularly calcite - can enable
93 reconstruction of fluid sources.

94
95 If stable isotope signatures of vein-forming minerals can be reconstructed, then this has the
96 potential to enable fluid source identification. The hydrogen isotopic signature (δD) of vein-forming
97 fluids can be measured by decrepitation if there is a high enough volume of fluid inclusions within
98 the vein-filling calcite (Gleeson et al. 2008). Fluid $\delta^{18}\text{O}$ can be calculated by determining the calcite
99 $\delta^{18}\text{O}$ and the temperature of precipitation (e.g. Epstein et al. 1951). Precipitation temperature of
100 calcite veins can be reconstructed from fluid inclusion microthermometry (e.g. Barker and
101 Goldstein 1990; Maskenskaya et al. 2014) or the more recently-developed clumped isotope
102 palaeothermometer. Clumped isotope palaeothermometry utilises the temperature dependence of
103 different isotopologues of CO_2 , particularly the mass 47 ^{13}C - ^{18}O - ^{16}O isotopologue (e.g. Schauble et
104 al. 2006; Eiler 2007). Calcite vein precipitation temperatures have been reconstructed using
105 clumped isotopes in geothermal/hydrothermal systems (Lu et al. 2017; Lu et al. 2018; MacDonald
106 et al. 2019), sedimentary basins (Mangenot et al. 2018a; Pagel et al. 2018) and fault systems
107 (Bergman et al. 2013; Hodson et al. 2016).

108
109 Stable isotope analysis can therefore provide details of the vein-forming fluid source. Different
110 fluids (e.g. magmatic, metamorphic, meteoric, seawater) have typical compositions in δD - $\delta^{18}\text{O}$ (V-
111 SMOW) space (e.g. Craig 1961; Taylor 1974; Rollinson 1993; Sharp 2007; Hoefs 2015). However,
112 these different fluids may have overlapping compositions, or their isotopic composition may have
113 changed over time. For example, a water with δD of -50 ‰ and $\delta^{18}\text{O}$ of +8 ‰ could be a magmatic
114 water or a metamorphic water (e.g. Hoefs 2015); equally though, it could be a meteoric water

115 which has undergone significant water-rock equilibration resulting in an enrichment of $\delta^{18}\text{O}$ (e.g.
116 Menzies et al. 2014). Thus, fluid stable isotope signatures in themselves do not always provide a
117 conclusive fingerprint of palaeofluid sources, especially in settings such as veins where there is
118 scope for significant water-rock interaction, and where genetic context of the hydrothermal system
119 may be equivocal.

120
121 Previous studies attempting to determine the origin of vein-forming fluids have often analysed trace
122 element concentrations in addition to stable isotopes. Barker et al., (2006) suggested varying trace
123 element concentrations (and stable isotope values) in anti-taxial veins could be caused by cycles
124 of fluid influx, water-rock interaction, and/or crack-seal processes. Maskenskaya et al., (2014)
125 found that trace element concentrations and distribution in veins did not correlate with stable
126 isotope (C, O, Sr) values; fractionation patterns of rare earth elements (REEs) were observed but
127 again these could not be correlated with any measured chemical, physical and isotopic variables
128 and so did not help to determine fluid source or vein formation mechanisms. Kalliomäki et al.
129 (2019) compared the trace element signatures of vein calcite and their host rock and showed in
130 examples from the Hattu schist belt (Finland) that interaction between the vein-forming fluid and
131 the host rock had strongly influenced the trace element signature of the resulting vein calcite.
132 Similarly, Wagner et al. (2010) used REEs to show that veins from the Rhenish Massif (Germany)
133 formed from advecting fluids which leached the wall rocks, which was reflected in vein mineral
134 trace element signatures.

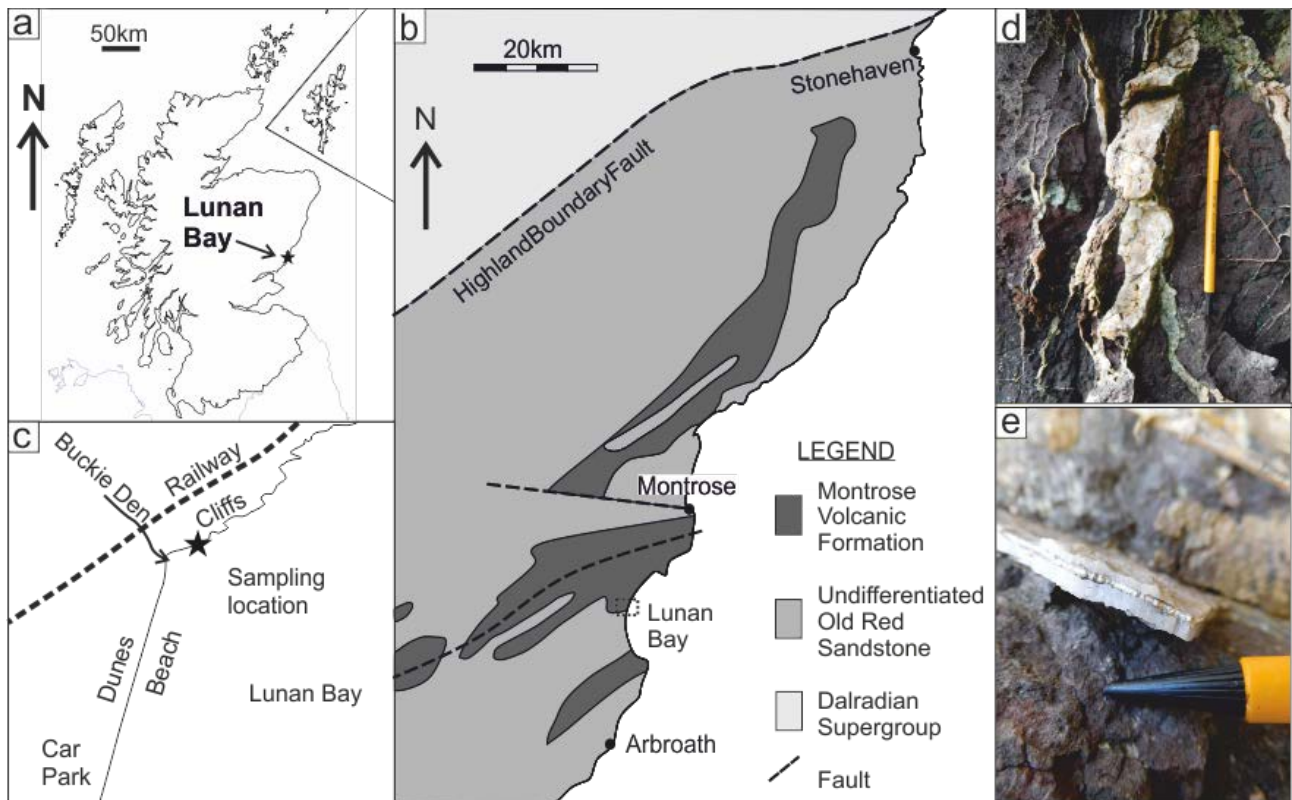
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136 The geological history of an area provides crucial context for discussion of potential fluid sources.
137 In a metamorphic terrane, clearly metamorphic fluids may be recorded. With veins, however, fluid
138 circulation may come sometime after formation of the surrounding geology and so linking vein-
139 forming fluids to host rocks is more complex. Establishing the age of precipitation of veins is
140 therefore key here to understanding the geological context of vein formation, and thus the likely
141 fluids involved. For example, if a vein with a calculated fluid $\delta^{18}\text{O}$ of +9 ‰ and δD of -50 ‰ can be
142 dated to within error of formation of nearby basalts, then a contribution of magmatic fluids to vein
143 precipitation cannot be discounted. The recent development of calcite U-Pb dating via Laser
144 Ablation Inductively Coupled Mass Spectrometry (LA-ICP-MS) has enabled precise, accurate and
145 rapid dating of calcite (e.g. Li et al. 2014; Coogan et al. 2016; Ring and Gerdes 2016; Roberts and
146 Walker 2016; Nuriel et al. 2017; Roberts et al. 2017; Drost et al. 2018). MacDonald et al., (2019)
147 used this technique to date calcite veins from ancient hydrothermal systems to show that closed-
148 system bond reordering (Passey and Henkes 2012; Henkes et al. 2014; Stolper and Eiler 2015) did
149 not affect determination of vein precipitation temperature from clumped isotopes.

150
151 In this contribution, we show that combining LA-ICP-MS U-Pb dating of calcite veins with stable
152 isotope and trace element analyses can help to fingerprint fluid source when trace elements and
153 fluid δD & $\delta^{18}\text{O}$ cannot provide an unequivocal interpretation. We use a case study of volcanic-
154 hosted veins in eastern Scotland, where this combination of proxies enables us to rule out
155 magmatic fluids, indicating the fluid source of veins was meteoric water which had undergone
156 significant water-rock interaction.

157 158 GEOLOGICAL SETTING AND SAMPLE PETROGRAPHY

159
160 Calcite veins from Lunan Bay in Angus, Scotland formed the basis of this study (Fig. 1a-b). The
161 study area is located within the northern part of the Midland Valley Terrane (e.g. Trewin 2002). The
162 host rocks to the calcite veins are the Montrose Volcanic Formation (MVF), a group of mingled
163 pahoehoe lavas, basaltic andesites, and volcanic-derived sediments deposited as part of the
164 ~2000 m thick, sandstone dominated Arbuthnott-Garvock Group (e.g. Armstrong and Patterson
165 1970; Bluck 2000; Browne et al. 2002; Hole et al. 2013). These lavas are likely sourced from the
166 northern flank of the Montrose Volcanic Centre, a north-east to south-west trending chain of
167 volcanoes active for ~15 Ma. The MVF lavas are suggested to be coeval with the Rhynie lavas to
168 the north, with a U-Pb andesite age of 411.5 ± 1.3 Ma (Parry et al. 2011). This places the MVF
169 within the late Lochkovian – early Pragian and at the boundary of the Arbuthnott and Garvock units
170 (e.g. Armstrong and Patterson 1970; Bluck 2000; Browne et al. 2002; Hole et al. 2013).

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Figure 1: location and field photographs. (a) Location of Lunan Bay case study site within Scotland; (b) the distribution of the host Montrose Volcanic Formation in the region; (c) the detailed location of the samples; Field photographs of examples veins (d) JV17-1 and (e) JV17-11.

The MVF lavas are basaltic to basaltic andesite (5.2-8.6 wt.% MgO, 52.6-57.6 wt.% SiO₂) in composition, and samples from St. Cyrus region have been described as olivine-plagioclase phyric, with olivine commonly pseudomorphed to iddingsite (Thirlwall 1981; Thirlwall 1982; Thirlwall 1983). Sub-euhedral, tabular, microphenocrystic plagioclase feldspar (labradorite to low Ab andesine, An₄₁₋₅₅) make up much of the matrix, along with abundant interstitial devitrified glass (Thirlwall 1982). Clinopyroxene is also present within some of the pahoehoe lava flows, predominantly in the form of augite (Hole et al. 2013). The lavas are also interbedded with locally sourced ephemeral playa-lake sediments, sandstones, and conglomerates, as well as air fall eruptions, creating complex sediment-lava interactions and abundant peperite formation (e.g. Hole et al. 2013). Within these mixing regions, secondary orthoclase is also present, as is sub-parallel flow alignment of feldspar laths and microphenocrysts (Thirlwall 1982; Thirlwall 1983).

Samples of MVF-hosted calcite veins were taken from the low cliffs just at the head of Lunan Bay at NO 69549 52488 (56°39'47.5"N, 02°29'54.1"W) (Fig. 1c). Images of vein petrography and relations to geochemical analysis are provided in Supplementary Figures S1-2. Most veins appear to be randomly oriented, with abundant stock work veining present; there is no clear field evidence of difference generations of veins. Five samples – JV17-1, -2, -9, -11 & -12 – were collected for analysis. The veins analysed in this study varied from >50 mm in width to less than 5 mm (Fig. 1d-e). Primary vein formation is along a singular opening (JV17-1, JV17-2, JV17-9), although some veins also occur as a bundle of connected sub parallel veins (JV17-12). Within these veins, multiple forms of calcite growth were recognized including bladed (JV17-1) and toothy (JV17-2, JV17-9) calcites along host rock contacts (Supp. Fig. S1), while euhedral, scalenohedral, and blocky crystals making up the bulk of most vein matrices (JV17-1, JV17-2, JV17-12) (Supp. Fig. S1). Crosscutting relationships and alteration are readily observed in JV17-1a, where a primarily syntaxial, bedrock growth phase is crosscut by a secondary, anhedral growth phase, and in JV17-11, where a stretched vein is crosscut by an iron rich vein. Multiple phases are also evidenced by bedrock fragments and remnant calcite crystal growth along these fragments that have been sealed within the vein during subsequent vein sealing (JV17-1, JV17-12) (Supp. Fig. S1).

205 JV17-11 contains the only formation of stretched beef-veining (appearing similar to beef tendons),
206 progressing from stretched/bladed crystals with vein opening (Supp. Fig. S1), although other
207 samples not included within this study were also observed to have significant beef veining. Minor
208 veins are prevalent in many of the samples (JV17-1, JV17-2, JV17-9, JV17-11), predominantly
209 sealed with fine, euhedral calcite crystals only a few mm in size. Accessory minerals (quartz and
210 chlorite) are visible along the vein-bedrock contacts, while reddish sutures are visible during the
211 final stage of vein formation/closure in JV17-1 and JV17-2, consisting primarily of iron oxides and
212 other rare carbonate phases (Supp. Fig. S1). These suture-defined vugs are filled with
213 predominantly cloudy, anhedral calcites. Despite the variations in texture, only JV17-11 exhibits
214 antitaxial vein growth (Supp. Fig. S1).
215 Minor variations in CL can be seen within the larger individual euhedral-anhedral blocky crystals
216 that make up the bulk of the vein matrix in JV17-1 and JV17-2 (Supp. Fig. S1). However, CL
217 signatures are generally uniform across individual veins and amygdales despite textural variation,
218 with primary excitation associated with calcite cleavage planes and extinction (Supp. Fig. S1).

219 220 METHODS

221
222 Cathodoluminescence (CL) petrography was undertaken using a Lumin HC4-LM hot-cathode CL
223 microscope at Saint Marys University. Plane-polarized and CL imagery was taken using an
224 incorporated Olympus BXFM focusing unit and Kappa DX40C peltier cooled camera, controlled by
225 the DX40C-285FW software package. The samples were analyzed under a vacuum, with an
226 accelerating voltage of ~6 KV, a beam current of 0.25 mA, and a 1 s camera exposure time with a
227 6 db camera gain.

228 $\delta^{13}\text{C}$ and $\delta^{18}\text{O}$ measurements were made at either the Scottish Universities Environmental
229 Research Centre (SUERC) or Memorial University Newfoundland's TERRA Stable Isotope Lab. At
230 SUERC, 1 mg of powdered sample was digested in 100% phosphoric acid in a 25°C water bath
231 prior to analysis in a VG OPTIMA mass spectrometer. A marble standard, as well as replicate
232 analyses, were used to calibrate the results, with reproducibility in $\delta^{13}\text{C}$ of $-9.2\pm 0.1\text{‰}$ and $\delta^{18}\text{O}$ of
233 $-15.0\pm 0.1\text{‰}$ (2σ) (Supp. Table S3). At Memorial, 0.2 mg of powdered sample was digested in
234 100% phosphoric acid in a 25°C water bath prior to analysis in a using a DeltaVPlus isotope ratio
235 mass spectrometer (IRMS) equipped with a Thermo Electron GasBench II unit. NBS19, plus two
236 internal standards, were used to calibrate the results, with NBS19 $\delta^{13}\text{C}$ of $2.0\pm 0.1\text{‰}$ and $\delta^{18}\text{O}$ of
237 $-2.2\pm 0.1\text{‰}$ (2σ), within error of accepted values (Friedman et al. 1982; Coplen et al. 2006) (Supp.
238 Table S3).

239 δD values of fluid inclusions in calcite vein chips were measured by *in vacuo* decrepitation
240 following the procedures outlined by Gleeson et al. (2008) at the Scottish Universities
241 Environmental Research Centre (Supp. Table S4). Procedural reproducibility was tested with 3 in-
242 house standards (Gleeson et al. 2008) and values were within 3 ‰ of long-term averages.

243 Carbonate clumped isotope (Δ_{47}) measurements were carried out in the Isotopologue
244 Paleosciences Laboratory at the University of Michigan, Ann Arbor. Samples were powdered using
245 a dental drill. For Δ_{47} analysis, ~ 8 mg of sample powder was reacted in an automated preparation
246 line previously described in Henkes et al., (2014). Carbonate powder was reacted under vacuum
247 with 104% phosphoric acid at 90 °C for 10 min. Vapour-phase water generated during the reaction
248 was separated from the produced CO_2 using liquid nitrogen swapped out an ethanol-liquid
249 nitrogen mixture held at -85 °C . The water remained frozen while the CO_2 was passed through a
250 Poropak Q chromatography trap held at -20 °C . The purified CO_2 was measured using a Nu
251 Instruments Perspective isotope ratio mass spectrometer in dual inlet mode, with a measurement
252 time of c. 2 h. All analyses were run as triplicates. Masses 44–49 were measured. Carrara marble,
253 NBS19 and an in-house carbonate standard (102-GCAZ) were used to verify the results. Carrara
254 Marble Δ_{47} averaged $0.417\pm 0.022\text{‰}$ (2σ , $n=3$), NBS19 Δ_{47} averaged $0.441\pm 0.021\text{‰}$ (2σ , $n=10$)
255 and 102-GCAZ averaged $0.650\pm 0.012\text{‰}$ (2σ , $n=14$) during the analytical window. (Supp. Table
256 S5).

257 All carbonate clumped isotope (Δ_{47}) values in this study are presented on an absolute
258 reference frame, also termed a 'carbon dioxide equilibrium scale' or CDES, which empirically
259 corrects for instrumental nonlinearities and changes in the ionization environment during mass
260 spectrometry (Dennis et al. 2011; Henkes et al. 2013). This reference frame was established by
261 periodically analyzing aliquots of CO_2 that were isotopically equilibrated at 25 or 1000 °C (Dennis

262 et al. 2011). Temperatures were calculated from Δ_{47} using the empirical “high temperature” Δ_{47} -
263 temperature relationship from Bonifacie et al., (2017). Fluid $\delta^{18}\text{O}$ values were calculated using the
264 equation of Friedman and O’Neil (1977).

265 Minor and trace element LA-ICP-MS analyses were undertaken at the Dalhousie
266 Laboratory for Experimental High Pressure Geological Research using a New Wave Research
267 frequency quintupled laser operating at 213 nm, coupled to a quadrupole mass spectrometer (PQ
268 Excell or Thermo X-series) with He flushing. The analyses occurred as both linescans and spot
269 analyses, with a 100 μm spot size, ablated at 4-5 Hz with a 20% total energy. Concentrations of
270 ^{43}Ca , ^{55}Mn , ^{57}Fe , ^{85}Rb , ^{86}Sr , ^{87}Sr , ^{89}Y , ^{137}Ba , ^{139}La , ^{140}Ce , ^{141}Pr , ^{146}Nd , ^{147}Sm , ^{153}Eu , ^{157}Gd , ^{159}Tb ,
271 ^{163}Dy , ^{165}Ho , ^{166}Er , ^{169}Tm , ^{172}Yb , and ^{175}Lu were measured in blocks of sixteen analyses, with two
272 NIST 610 bounding each block for a total of 20 analyses per run. Total run times were 140 s, with
273 20 s laser warm-up, 60 s ablation, and 60 s He-gas flushing time. However, due to calcite burn
274 through, many analyses were between 20-30 s in order to prevent damage to the slide.

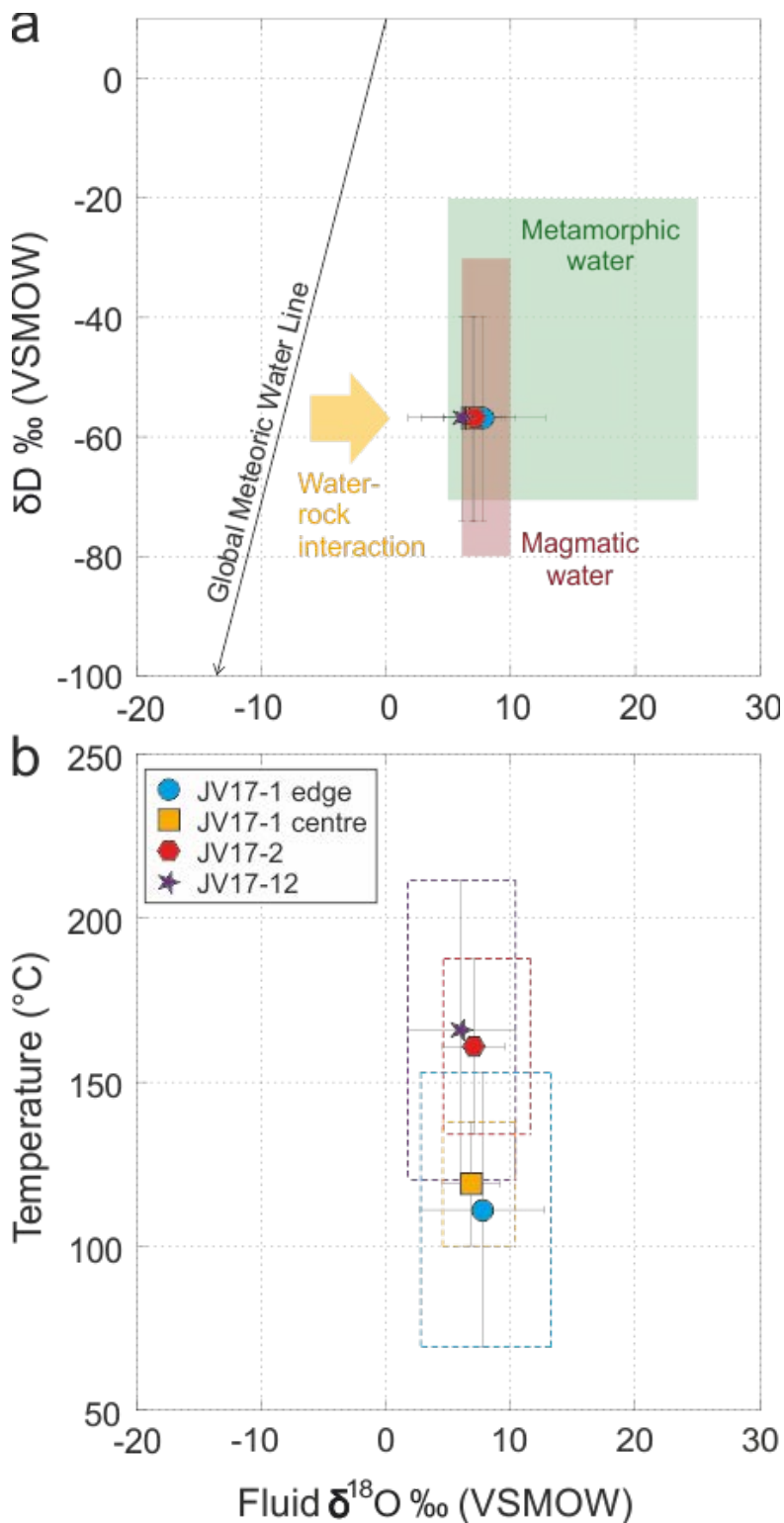
275 Data reduction was conducted off-line using Iolite software. Base levels were determined
276 through ^{43}Ca peak analysis making sure to avoid anomalous intensities but also including washout
277 periods. Analytical drift was addressed by running a linear regression through average ^{43}Ca
278 intensities in the NIST SRM610 runs before and after unknown analyses; reproducibility in was
279 better than 5% for all elements analysed in NIST SRM610. Average concentrations of all elements
280 were within error of published values (Jochum et al. 2011) (Supp. Table S6). REE values were
281 normalized to chondrite (McDonough and Sun 1995) using Microsoft Excel, following methods
282 outlined in Rollinson (1993).

283 LA-ICP-MS U-Pb calcite dating was conducted at the Geochronology & Tracers Facility,
284 British Geological Survey (Nottingham, UK) using a New Wave Research 193UC excimer laser
285 ablation system, coupled to a Nu Instruments Attom single-collector sector-field ICP-MS following
286 the methods outlined by Roberts and Walker (2016). Samples were pre-ablated with a 150 μm
287 spot for 30 pulses. Full ablation conditions comprise a 100 μm spot for 30 seconds, at 10 Hz and a
288 fluence of ca. 8 J/cm^2 . A gas blank of ca. 60 seconds is measured at the beginning of each run.
289 Normalisation uses NIST614 for $^{207}\text{Pb}/^{206}\text{Pb}$ and WC-1 for $^{206}\text{Pb}/^{238}\text{U}$, with data reduction and
290 uncertainty propagation following Roberts et al., (2017) and the recommendations of Horstwood et
291 al., (2016), and conducted using an in-house spreadsheet and the Nu Attolab Time Resolved
292 Acquisition software. Spot analyses with low count rates (< 100 cps) or high uncertainties (>7.5%
293 1σ) are removed from age calculations. Age calculations and plotting were conducted using Isoplot
294 4.15 (Ludwig 2012). Duff Brown Tank limestone was analysed during the session as a validation
295 material; an age of 63.5 ± 1.7 Ma (MSWD = 2.9) was obtained (Supp. Table S7), which overlaps
296 the published age of 64.04 ± 0.67 Ma (Hill et al. 2016).

297 298 RESULTS

299
300 The location of analyses of all types in the vein samples are shown in Supplementary Figures S1-2
301 and standard and sample geochemical data are given in Supplementary Tables S3-6. Across the 5
302 samples analysed, $\delta^{13}\text{C}$ ranged from -1.90 ‰ to -9.87 ‰ but with the majority between -3 and -4 ‰
303 (Table 1, Supp. Table S3). There was no clear correlation in $\delta^{13}\text{C}$ values and calcite crystal
304 shape/vein texture or vein width at the point of analysis (Supp. Fig. S2). Vein calcite $\delta^{18}\text{O}$ (V-PDB)
305 values ranged from -1.36 ‰ to -13.21 ‰ (Table 1, Supp. Table S3). Narrower veins tended to have
306 more depleted $\delta^{18}\text{O}$ values, although this did not hold true for all samples, and $\delta^{18}\text{O}$ varied by
307 several permil in single veins (up to ~10.5 ‰ between two adjacent analyses in vein JV17-1)
308 (Table 1, Supp. Fig. S2). A δD value of -56.8 was obtained from fluid inclusions in calcite chips
309 from JV17-1 (Table 1, Fig. 2a). Four clumped isotope temperatures were determined from three of
310 the samples. The edge of the large vein in sample JV17-1 yielded a temperature of 111 ± 42 °C
311 while the centre of the same vein recorded 119 ± 19 °C. A temperature of 161 ± 27 °C was recorded
312 from the centre of the large vein in JV17-2, and the set of sub-parallel linked veins in JV17-12
313 yielded a temperature of 166 ± 46 °C (Table 1, Fig. 2b).

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 320 Figure 2: stable isotope data. a) Fluid δD and calculated fluid $\delta^{18}O$ values using Friedman and
 321 O'Neil (1977) equation with the temperature and calcite $\delta^{18}O$ from clumped isotope analyses; $\delta^{18}O$
 322 error bars are 1 standard error propagated from the clumped isotope analysis while δD error bars
 323 represent the range of δD values obtained during the analysis; Global Meteoric Water Line from
 324 Craig (1961); range of typical isotopic composition of metamorphic water (e.g. Taylor, 1974; Hoefs,
 325 2015) and magmatic water (e.g. Hoefs, 2015). b) temperature calculated using Bonifacie et al.,
 326 (2017) Δ_{47} -T calibration plotted against fluid $\delta^{18}O$; temperature error bars are 95% confidence level
 327 while fluid $\delta^{18}O$ error bars are the conventional 1 standard error; dashed boxes constrain the 95%
 328 confidence level temperature error and the maximum range of fluid $\delta^{18}O$ calculated from the
 329 clumped isotope temperatures and the range of calcite $\delta^{18}O$ measured in the different samples.
 330 Blue circle = JV17-1 edge; orange square = JV17-1 centre; red hexagon = JV17-2; purple star =
 331 JV17-12.

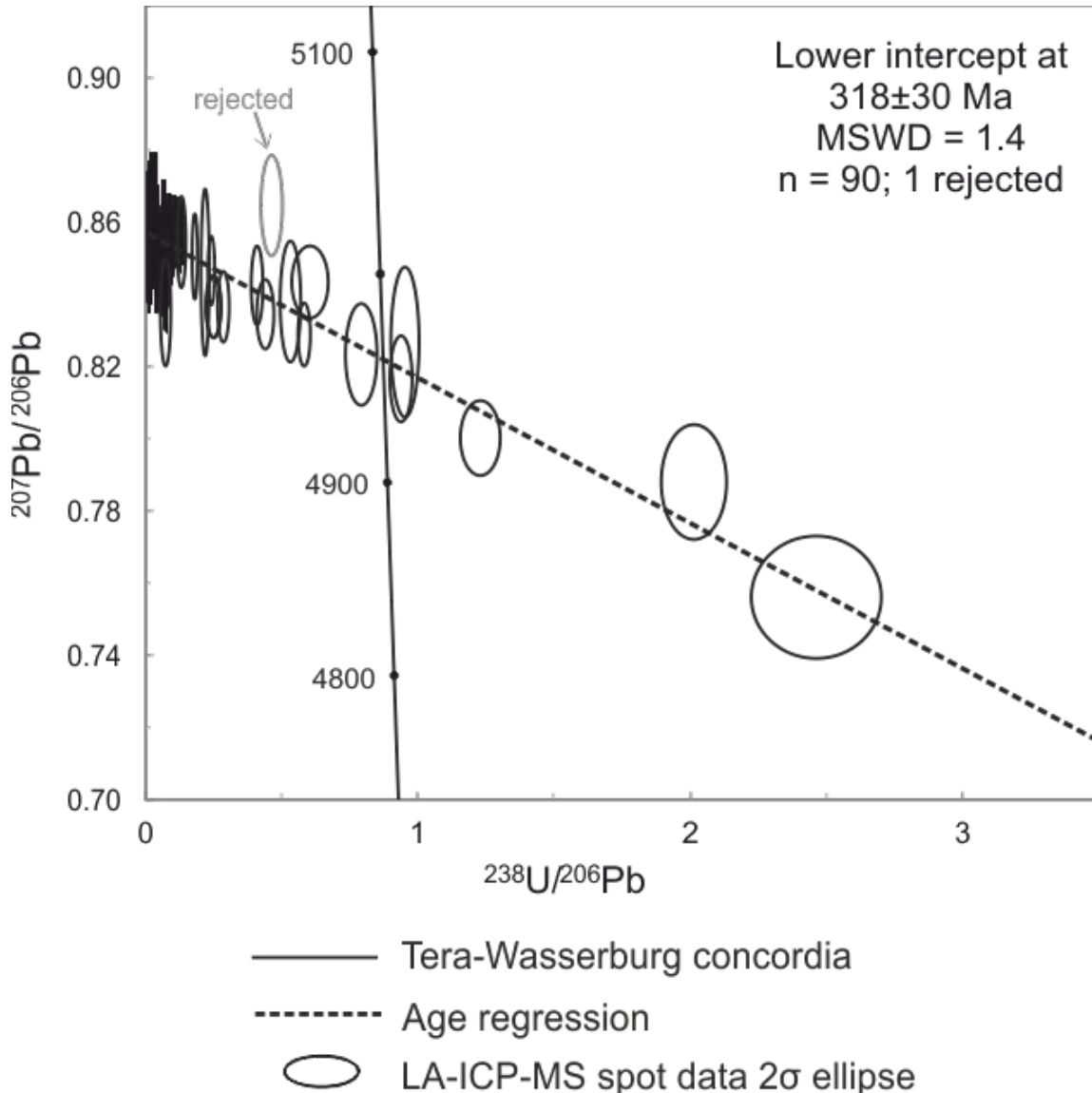
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333 Fe concentrations were ~300-10000 ppm, with the majority <1000 ppm; Mn concentrations were
 334 ~700-12000 ppm. There was no clear correlation between Fe or Mn concentration and position
 335 across veins (i.e. edge to centre) and cathodoluminescence intensities were fairly uniform across
 336 all veins (Table 1, Supp. Fig. S2). Total Rare Earth Elements (Σ REE) values were ~1-1750 ppm
 337 (Table 1). In vein JV17-1, there was slight pattern of higher REE concentration at the vein edges
 338 than the core; however, this pattern was not present in the other wide vein (JV17-2) (Supp. Fig.
 339 S2). The other veins were too narrow for an assessment of REE concentration across the vein. All
 340 analyses had higher light REE concentrations than heavy REEs. A number of analyses had flat
 341 normalised LREE-MREE patterns; La/Gd ratios were usually lower than Gd/Lu ratios (Table 1). Ce
 342 anomalies were negligible, with Ce/Ce* values of 0.7-1.2, representing slight negative to slight
 343 positive anomalies (Table 1). Eu anomalies were also mainly negligible, with slight negative (0.7) to
 344 slight positive (1.2) values; a small number of analyses recorded more positive anomalies (1.5-2.0)
 345 (Table 1, Supp. Table S6).

346

347 One sample (JV17-2) yielded a calcite U-Pb age. The age of 318 ± 30 Ma (MSWD = 1.4) was
 348 derived from regression of 89 spot analyses in that vein, with one analysis lying off the regression
 349 being rejected. This age includes propagation of the systematic uncertainties (Table 1, Fig. 3).

350



351

352 Figure 3: Tera-Wasserburg concordia plot showing age regression through LA-ICP-MS analytical
 353 spots (blue ellipses; rejected spot in grey); precipitation age defined as the lower intercept age of
 354 318 ± 30 Ma (2σ).

355

Sample	Description	δD (VSMOW, ‰)	$\delta^{13}C$ (VPDB, ‰)	$\delta^{18}O_{calc}$ (VPDB, ‰)	T_{47} (°C)	$\delta^{18}O_{fluid}$ (VSMOW, ‰)	Mn (ppm)	Fe (ppm)	ΣREE (ppm)	La/Gd	Gd/Lu	Ce/Ce*	Eu/Eu*	U-Pb Age (Ma)
JV17-1	wide	-56.55	-1.90 to -5.21	-1.36 to -11.83	edge 111±42 centre 119±19	3.0 to 13.7 4.9 to 10.6	712-4230	373-846	4-1742	1.4-7.0	4.5-13.8	0.7-1.0	0.8-1.9	-
JV17-2	wide	nd	-3.46 to -4.55	-6.89 to -12.10	161±27	6.9 to 12.2	1996-6290	382-687	24-376	0.6-7.3	2.6-6.0	0.8-1.1	0.9-1.7	328±27
JV17-9	narrow	nd	-3.8 to -3.55	-8.25 to -9.51	-	-	7070-9380	719-1320	248-769	8.7-10.7	13.1-19.6	0.7-0.9	0.7-0.8	-
JV17-11	narrow	nd	-5.64 to -9.87	-10.77 to -12.89	-	-	3228-7001	447-5110	59-159	10.1-27.0	2.8-5.6	0.7-0.8	0.9-1.7	-
JV17-12	complex of connected sub-parallel narrow veins	nd	-3.04 to -3.35	-12.18 to -13.21	166±46	6.1 to 7.1	1780-5390	509-670	53-280	4.1-11.0	1.9-10.7	0.7-0.9	0.9-2.0	-

Table 1: summary data table; 'nd' denotes not enough water was recovered from these samples to make a measurement.

DISCUSSION

Stable Isotopes

Calcite which has resided in the subsurface at high temperatures (ca. >100 °C) for a long period (ca. >100 Myr) is susceptible to solid-state bond reordering (Passey and Henkes 2012; Henkes et al. 2014; Shenton et al. 2015; Stolper and Eiler 2015). Passey and Henkes (2012) interpreted a two-stage bond reordering process of an initial phase of defect annealing followed by solid-state diffusion. Stolper and Eiler (2015) proposed different mechanisms: an initial rapid change of ~1-40 °C at ambient temperatures of ~75-120 °C sustained for ~100 Myr due to diffusion of isotopes through the crystal lattice; after a period of stability, a secondary stage of slow isotope exchange reactions between adjacent carbonate groups at >~150 °C sustained for >~100 Myr which may bring the clumped isotope temperatures to the ambient temperature.

Given that the calcite U-Pb dating indicates that the veins are much older than ca. 100 Myr, and clumped isotope thermometry yields temperatures of ca. 100 °C in all samples, thermal history reordering models (THRMs) were run to test for bond reordering (Supp. Table S8). The THRM approach developed by Shenton et al., (2015) involves modelling temporal evolution in Δ_{47} based on kinetic parameters (e.g., activation energy, E_a and pre-exponential factor, K_0) derived from Arrhenius regressions of experimental data from Passey and Henkes (2012). THRMs require knowledge or assumptions about the temperature history of the analysed sample. This temperature history is divided into a series of time steps with a specified ambient temperature (converted back to Δ_{47}) at each time step. The bond reordering reaction (reaction 13 in Passey and Henkes (2012)) is then used to calculate the extent of clumped isotope reordering during each step. The 'new' Δ_{47} value at the end of each time step is treated as the 'initial' Δ_{47} value for the next step and the model is run iteratively from the time of initial calcite precipitation to the present day (Shenton et al. 2015). Additionally, calcite of different origin (e.g. brachiopods vs spar calcite vs optical calcite) were found to have different reordering kinetics (activation energy and pre-exponential factor) during laboratory experiments (Passey and Henkes 2012; Henkes et al. 2014).

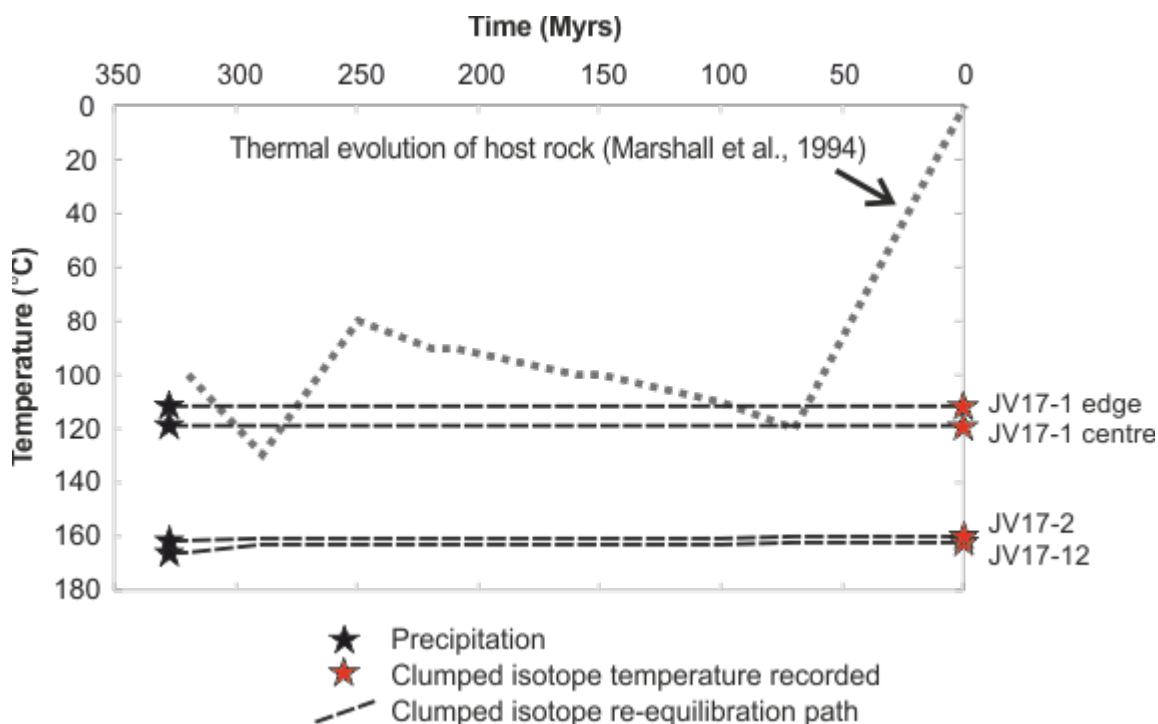
In addition to the activation energy and pre-exponential factor, the assumed initial precipitation temperature and age of precipitation are input to run the model. For sedimentary or biogenic calcite, an assumed surface temperature of ca. 25 °C (or a more accurate one based on species in biogenic calcites) is used (Henkes et al. 2013; Henkes et al. 2014). For calcite veins this is challenging as one cannot assume an initial precipitation temperature. We assumed that the temperature reconstructed from clumped isotope analysis was the initial precipitation temperature and forward modelled using an ambient thermal history to determine if bond reordering had occurred.

THRMs were run for all samples using the burial history for the local area constructed from vitrinite reflectance data and an assumed geotherm of 30 °C/km (Marshall et al. 1994), along with the calcite precipitation ages derived in this study from LA-ICP-MS calcite U-Pb dating. Kinetic parameters for both optical and spar (labile and refractory) calcite from Passey and Henkes (2012) were used but the choice of kinetic parameters did not affect the model output. This is because the THRMs indicate that negligible (much less than analytical error) bond reordering took place in any of these samples (Fig. 4).

Clumped isotope temperatures from the centre and edge of the large vein in sample JV17-1 are within error, suggesting that temperature remained relatively constant during calcite precipitation. However, the temperatures from JV17-1 are ~50 °C lower than in JV17-2 and JV17-12. The origin of this difference is unclear but may reflect an age difference – JV17-1 may be younger than JV17-2 and JV17-12 and records a cooling of the vein-forming fluid. Unfortunately, it was not possible to obtain an age from JV17-1 and so this cannot be proven. Calcite $\delta^{18}O$ (V-PDB)

409 values varied by several permil within veins, suggesting that as well as some minor temperature
 410 variation within veins, slight variation in source fluid $\delta^{18}\text{O}$ and/or interaction with oxygen from the
 411 wall rocks of the veins resulted in variability in calcite $\delta^{18}\text{O}$. As the THRM's have shown that the
 412 clumped isotope temperatures do indeed represent the calcite precipitation temperature, the $\delta^{18}\text{O}$
 413 of the parent fluids can be reconstructed.

414 In sample JV17-1, calculated fluid $\delta^{18}\text{O}$ (V-SMOW) values ranges from 3 to 14 ‰, but with
 415 the majority in the 7-9 ‰ range. Similarly for JV17-2, values are ~7-12 ‰, with most 7-9 ‰. In
 416 sample JV17-12, values are ~6-7 ‰ (Table 1, Fig. 2). Values such as these represent fluids
 417 isotopically enriched relative to VSMOW and are typical of metamorphic waters, magmatic waters,
 418 or meteoric/marine waters which have undergone significant fluid-rock interaction (e.g. Sharp
 419 2007). Barker et al., (2009) suggested that homogeneity of calcite $\delta^{18}\text{O}$ across veins may indicate
 420 the progressive reaction of fluids with host rock, with sufficient reaction occurring along discrete
 421 fluid flow pathways to fully equilibrate the fluids for these isotope systems. Samples JV17-9, JV17-
 422 11 and JV17-12 (narrow veins) display this homogeneity (<~2 ‰ variation), as does large vein
 423 JV17-2 apart from a single analysis at the very edge of the vein which is ~3 ‰ less depleted than
 424 the rest of the analyses from that vein (Supp. Fig. S2). Sample JV17-1, however, displays a wide
 425 range in calcite $\delta^{18}\text{O}$. Additionally, $\delta^{13}\text{C}$ values are all negative which suggest oxidised fluids
 426 (Barker et al. 2006) which is more likely to be a surface water but is not conclusive. The δD values
 427 are also inconclusive and could represent metamorphic, magmatic or meteoric waters. Even when
 428 taking δD and $\delta^{18}\text{O}$ together, the samples fall within the field of typical magmatic and metamorphic
 429 waters or could represent meteoric water which has undergone significant water-rock interaction
 430 (e.g. Taylor 1974; Rollinson 1993; Sharp 2007; Hoefs 2015) (Fig. 2b). Stable isotopes and
 431 reconstructed fluid $\delta^{18}\text{O}$ alone can therefore not distinguish fluid sources.
 432



433
 434 Figure 4: Thermal history reordering models for vein samples where clumped isotope temperatures
 435 were obtained showing the modelled evolution of clumped isotope temperature (calculated using
 436 the equations in Passey et al., (2012) and the approach of Shenton et al., (2015)) and ambient
 437 temperature after vein precipitation (from Marshall et al., (1994)).
 438

439 Trace Elements

440 All REEs are incompatible in basalt/andesite (the host rocks to this study) (e.g. Rollinson 1993)
 441 and so are easily scavenged by fluids. Calcite fractionates LREE over HREE during precipitation,
 442 leading to the negatively-sloping normalised REE patterns (e.g. Bau and Möller 1992; Denniston et
 443 al. 1997; Morad et al. 2010) as indicated by positive La/Gd and Gd/Lu ratios (Table 1). REE
 444 concentrations in the calcite veins in this study are higher than in typical freshwater/seawater (e.g.
 445 Rollinson 1993; Morad et al. 2010), suggesting metamorphic/magmatic fluids or

446 freshwater/seawater which has undergone significant water-rock interaction. This agrees with the
447 interpretation of the reconstructed fluid $\delta^{18}\text{O}$ values but still does not fingerprint a particular source
448 fluid.

449 Conceptually, in a wider vein, e.g. JV17-1, if there was a lot of fluid-rock interaction then
450 minor and trace element concentrations would be expected to be higher at the vein edge than the
451 centre (assuming growth out from the vein edge as the petrographic analysis suggests); initial
452 reaction of the fluid with the fracture walls would scavenge trace elements (e.g. incompatible
453 REEs) and incorporate these in precipitated calcite. As the vein infilled, initially-precipitated calcite
454 would 'armour' the wall rock from the fluid and so later calcite, precipitated in the centre of the vein,
455 would have lower REE concentrations. There is a suggestion of this in total REE in JV17-1 but not
456 clearly (Supp. Fig. S2); however, such a pattern does not occur in veins JV17-2 or JV17-12 (Supp.
457 Fig. S2). This pattern of higher trace element concentration at vein edges is also not shown clearly
458 by Fe or Mn in any of the analysed veins (Supp. Fig. S2).

459 Similarly, conceptually, in narrow veins all the calcite might be expected to have higher
460 REE concentrations given the lower calcite-wall rock ratio. This is not borne out by the total REEs
461 but high La/Gd values – signifying strong fractionation of LREEs into the calcite – are found in
462 narrow veins JV17-9 and JV17-11 (Table 1). JV17-9 in particular also shows the highest
463 concentrations of Mn, again indicating significant fluid-rock interaction (Table 1).

464 The veins generally showed no strongly positive Eu anomalies which suggests little
465 interaction with Ca-rich bedrock, as Eu substitutes for Ca, predominantly in plagioclase (Barker et
466 al 2006). The MVF is high in Ca (in clinopyroxene and plagioclase) and so extensive water-rock
467 interaction would be expected to lead to positive Eu anomalies. This is generally not seen although
468 some spots do have strongly positive Eu anomalies (up to 2.0); these do not appear to clearly
469 correlate with position in the vein in relation to the wall rock or calcite petrography (Supp. Fig. S2).
470 The Eu anomaly is therefore inconclusive regarding the origin of the fluid.

471 The Ce anomaly can be used as a redox proxy for fluids in veins, where a negative Ce
472 anomaly indicates oxidising conditions (e.g. Göb et al. 2013). No strongly negative (or positive) Ce
473 anomalies were found in the veins in this study, indicating the fluid was not highly oxidised at the
474 time of calcite precipitation. While well-oxidised surface waters have negative Ce anomalies, fluids
475 which originate from the subsurface (magmatic/metamorphic fluids) or surface waters which have
476 resided in the crust for some time and undergone water-rock interaction show no Ce anomaly (e.g.
477 Göb et al. 2013). The Ce anomaly is therefore in agreement with the stable isotope data, in that the
478 fluid from which the veins precipitated from was either a magmatic/metamorphic fluid, or a
479 meteoric/marine water which had undergone significant water-rock interaction. No correlation was
480 found between calcite crystal microstructure and stable isotopes/trace elements (Supp. Fig. S2).
481 This lack of correlation, and inability to fingerprint the source fluid, was also encountered by
482 Maskenskaya et al., (2014) in a previous study.

483 484 *Calcite Geochronology*

485 Based on stable isotopes and trace elements, it has not been possible to distinguish the fluid
486 source which formed the calcite veins between either a deep isotopically-enriched fluid (magmatic
487 or metamorphic water) or a surface water which has undergone significant water-rock interaction.
488 Simply based on the local geology, metamorphic waters can likely be ruled out as the fluid source
489 as the nearest exposed metamorphic rocks are ~25 km away beyond the Highland Boundary Fault
490 and the age of metamorphism (~470 Ma, Viète et al. 2013) long predates the formation of the
491 Devonian MVF host rocks to the calcite veins.

492 Magmatic waters remain a viable fluid source as the host rocks are volcanic, and sporadic
493 volcanic activity occurred through time in the Midland Valley Terrane (Cameron and Stephenson
494 1985). Determining whether magmatic waters are a likely fluid source requires the age of calcite
495 precipitation in the veins to be known, so that that age can then be compared to ages of
496 volcanic/magmatic activity in the local area. If vein calcite precipitation occurred very soon after the
497 formation of the MVF from residual waters from the volcanic activity, then the calcite should yield
498 an age within error of the MVF crystallisation age. Given that the MVF is correlated with the Rhynie
499 Chert dated at 411.5 ± 1.3 Ma (Parry et al. 2011) and the calcite precipitation age is 318 ± 30 Ma, this
500 shows that the calcite did not form from a magmatic fluid associated with the formation of the host
501 MVF. There is some Lower Carboniferous (potentially within uncertainty of the age from the Lunan
502 Bay calcite veins) volcanic activity in the Midland Valley Terrane, but the nearest is located several

503 tens of kilometres to the south around St Andrews (Cameron and Stephenson 1985), and so it is
504 unlikely that magmatic fluids associated with this volcanic activity were the source fluids for the
505 calcite veins.

506 507 *Importance of Calcite Geochronology in Fingerprinting Vein Fluid Sources*

508 LA-ICP-MS U-Pb calcite dating has enabled us to rule out magmatic fluids as the source for calcite
509 veins hosted in basaltic andesites when stable isotopes and trace elements were unable to do so.
510 For the Lunan Bay calcite veins, the fluid source must be a surface water which has undergone
511 considerable fluid-rock interaction, leading to the enriched fluid $\delta^{18}\text{O}$ reconstructed from stable
512 isotope analysis and clumped isotope thermometry. Palaeogeographic reconstructions indicate
513 that the area was coastal terrestrial lowland for much of the Carboniferous (Cope et al. 1992), and
514 we interpret that this surface water was likely meteoric water, rather than seawater/brine.

515 This case study from Lunan Bay highlights, along with previous studies from other locations
516 (e.g. Maskenskaya et al. 2014), the difficulty in fingerprinting fluid source from stable isotopes
517 and/or trace elements. In this study, LA-ICP-MS U-Pb calcite geochronology helped eliminate
518 potential fluid sources, enabling determination of the most likely fluid source for the analysed
519 calcite veins. It is an additional proxy that should be used alongside stable isotopes and trace
520 elements in studies where the fluid source of veins, or indeed any other geological feature where
521 the parent fluid may have resided in the crust for a period of time such as fault precipitates (e.g.
522 Roberts and Walker 2016; Parrish et al. 2018) or cements (e.g. Mangenot et al. 2018b; Pagel et al.
523 2018).

524 525 CONCLUSIONS

526 In this contribution we have shown that combining LA-ICP-MS U-Pb calcite dating with stable
527 isotopes (including clumped isotope palaeothermometry) and trace element analysis increases the
528 likelihood of determining the fluid source of veins. Calcite veins hosted in the Devonian Montrose
529 Volcanic Formation at Lunan Bay in the Midland Valley Terrane of Central Scotland were used as
530 a case study. δD values of fluid inclusions in the calcite, and parent fluid $\delta^{18}\text{O}$ values reconstructed
531 from clumped isotope palaeothermometry, gave values which could represent a range of fluid
532 sources: metamorphic or magmatic fluids, or surface waters which had undergone much fluid-rock
533 interaction. Trace elements showed no distinctive patterns and shed no further light on fluid
534 source. LA-ICP-MS U-Pb dating determined the vein calcite precipitation age – 318 ± 30 Ma – which
535 rules out metamorphic or magmatic fluid sources as no metamorphic or magmatic activity was
536 occurring in the area at this time. The vein fluid source was therefore a surface water (meteoric
537 based on paleogeographic reconstruction) which had undergone significant water-rock interaction.
538 This study highlights the importance of combining the recently developed LA-ICP-MS U-Pb calcite
539 geochronometer with stable isotopes and trace elements to help determine fluid sources of veins,
540 and indeed any geological feature where calcite precipitated from a fluid which may have resided
541 in the crust for a period of time (e.g. fault precipitates or cements).

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