1	A new approach to geobarometry by combining fluid inclusion and clumped isotope thermometry in hydrothermal carbonates
2	and crumped isotope mermometry in nydromermar carbonates
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19	Running title :
20	Geobarometry from diagenetic carbonates
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25 ABSTRACT

26 This study presents a new approach to geobarometry by combining fluid inclusion and clumped 27 isotope (Δ_{47}) thermometry on carbonate minerals. The offset between homogenization 28 temperatures of primary fluid inclusions with known composition and Δ_{47} temperatures of the 29 host mineral allows a direct estimation of the fluid pressure at the time of carbonate 30 crystallization. This new approach is illustrated via hydrothermal dolomite samples from the 31 Variscan foreland fold-and-thrust belt in northern Spain. Clumped isotope analyses yield 32 crystallization temperatures (107-168 °C) which are higher compared to homogenization temperatures in corresponding samples (95-145 °C). The calculated pressure values suggest that 33 34 dolomitizing fluids were overpressured during formation of zebra dolomite textures, whereas lower pressures are obtained for dolomite cement from breccia textures. This new approach to 35 36 geobarometry opens up the possibility of estimating the pressure of carbonate crystallization and 37 has potential applications in diagenesis, basin analysis, ore geology and tectonics.

38

39 INTRODUCTION

40 Pressure is an important parameter to be considered when investigating a wide range of 41 geological processes. Geobarometry can yield information on diagenesis, basin evolution, 42 hydrocarbon generation and boiling of ore-forming fluids, among others. To date, only a limited 43 number of geobarometers exist, only a few of which are applicable to sedimentary and diagenetic 44 settings. Most of these geobarometers are, at least partly, based on the use of fluid inclusion (FI) 45 data (Roedder and Bodnar, 1980). One way to estimate pressure conditions during crystal growth 46 is by combining information from two immiscible fluids which are trapped simultaneously (e.g. 47 petroleum and aqueous fluids; Munz, 2001). This approach is hampered by the need for multiple

48 cogenetic FI generations, which are not frequently encountered. If only one generation of 49 inclusions is present, information on its homogenization temperature and composition can be 50 used for geobarometry, but only in combination with an independent geothermometer which can 51 provide the temperature of formation of the host crystal (e.g. chlorite thermometry; Schroyen and 52 Muchez, 2000). An important assumption is that both FIs and mineral phases used as 53 independent geothermometers are spatially and temporally associated, which is not easy to prove 54 (Roedder and Bodnar, 1980).

55 This study presents a new approach to geobarometry by combining aqueous FI 56 microthermometry with the technique of carbonate clumped isotope (CI) paleothermometry (Δ_{47} ; 57 Ghosh *et al.*, 2006). Homogenization temperatures (T_h) obtained from hydrothermal dolomite 58 samples are compared to crystallization temperatures $(T\Delta_{47})$ obtained by measuring the abundance of ¹³C-¹⁸O bonds in the mineral lattice. On the assumption that both FI and CI 59 60 signatures are pristine and have not been reset, the offset (ΔT) between both temperature sets can 61 be explained by the fact that T_h represents the minimum temperature of FI entrapment while 62 $T\Delta_{47}$ represents the true temperature of crystallization of the host dolomite mineral (figure 1). ΔT 63 is influenced by the pressure at the time of inclusion entrapment, referred to as the pressure 64 correction (Goldstein, 2001). Hence, it can give a direct estimate of pressure at the time of 65 dolomite crystallization, without the need for cogenetic FI generations or other types of geothermometers, since both temperatures are obtained from the same carbonate mineral phase. 66

67

68 **GEOLOGICAL SETTING**

The Cantabrian Zone (CZ) represents the Variscan foreland fold-and-thrust belt (FFTB) on the
Iberian Peninsula (Julivert, 1971). Carbonate production prevailed in the foreland basin during

the Mississippian and Early Pennsylvanian, following an important marine transgression. One of
the most common Carboniferous limestone formations in the CZ is the Serpukhovian Barcaliente
Formation (Fm.), composed of thinly bedded dark carbonate mudstones (Wagner *et al.*, 1971).

74 The Paleozoic successions of the CZ were folded and thrusted into an east-verging FFTB during 75 Moscovian times (Alonso et al., 2009). The original N-S oriented mountain range evolved into a 76 thick-skinned secondary orocline during the Late Carboniferous (ca. 310-297 Ma; Gutiérrez-77 Alonso et al., 2015). Following oroclinal buckling, the thickened lithosphere underneath the 78 center of the orocline (corresponding to the CZ) became unstable, detached and sunk away 79 resulting in upwelling of hot asthenospheric material (ca. 295-285 Ma; Valverde-Vaquero et al., 80 1999). The upwelling created near-surface uplift and extension as well as an increase in crustal 81 heat flow (Weil et al., 2013).

82 As a result, hypersaline marine-derived brines infiltrated the subsurface along reactivated 83 Variscan fractures in the Bodón Unit, an important thrust unit in the CZ (figure 2; Julivert, 1971; 84 Gasparrini et al., 2006a). These brines were modified through water-rock interactions and the 85 resulting hydrothermal fluids dolomitized Paleozoic limestones through thermal convection 86 (Gasparrini et al., 2006a; Honlet et al., in press). Inclusions within the dolomite crystals are two-87 phase aqueous with consistent liquid to vapor ratios. FI microthermometry on void-filling 88 dolomite yielded T_h values between 110 to 150°C (Gasparrini et al., 2006b). Salinity calculations 89 based on microthermometry coupled with cryo-Raman spectroscopy revealed salinities around 90 26 eq. wt% CaCl₂ (Gasparrini et al., 2006b). Raman spectroscopic measurements on the gas 91 bubbles did not detect gases such as CO₂, CH₄ or N₂ (Gasparrini, 2003). Hydrothermal fluid 92 circulation was the hottest event affecting the Paleozoic rocks of the CZ, based on apatite fission-93 track (AFT) thermochronology (Carrière, 2006).

94 METHODS

Our dataset consists of 7 new samples and 2 samples provided by Gasparrini *et al.* (2006b; table
1). All samples were collected from the Barcaliente Fm. exposed in the Bodón Unit, from
locations (figure 2) that correspond to those of Gasparrini *et al.* (2006b).

98 Samples and thin sections were investigated with plane light microscopy (Leica DM LP 99 microscope) and cathodoluminescence microscopy (Techosyn Model 8200 Mark II instrument 100 mounted on a Nikon microscope).

101 In order to obtain a precise comparison between T_h and $T\Delta_{47}$, new FI measurements were 102 performed on all samples, including those provided by Gasparrini et al. (2006b). Inclusions 103 suitable for microthermometric analysis were selected based on a petrographic study of both thin 104 sections and double polished thick sections (\pm 100 µm thick; cold-polishing technique; cf. 105 Muchez et al., 1994). Measurements were carried out on a Linkam MDS-600 heating/cooling 106 stage mounted on an Olympus BX51 microscope. The stage was calibrated by synthetic Syn FlincTm FI standards. Isochores were calculated for each sample using salinity data from 107 108 Gasparrini et al. (2006b) and the equation of state of Zhang and Frantz (1987).

109 Clumped isotope analyses were performed in the Qatar Stable Isotope Lab at Imperial College 110 London. The method used for the analyses is based on Dennis and Schrag (2010) and described 111 for the Imperial College laboratory in Dale et al. (2014). Powder samples were drilled from rock 112 slabs using a dental drilling device. On average 3 to 5 replicate measurements were performed 113 for each sample using the same manual CO₂ purification line and mass spectrometer. Aliquots of 114 5 to 6 mg of dolomite powder were digested online in a phosphoric acid bath held at 90°C for 20 115 minutes to allow complete reaction. The liberated CO₂ gas was subsequently purified and 116 measured on a Thermo Fisher MAT 253 isotope ratio mass spectrometer. Data processing was

117 performed using the free software Easotope (John and Bowen, 2016). Non-linearity corrections 118 were done based on heated gases (Huntington et al., 2009), and the projection of the data into the 119 absolute reference frame of Dennis et al. (2011) was performed using heated gases and inter-120 laboratory standards Carrara marble, ETH2 and ETH3 (Meckler et al., 2014; average SE 121 between 0.0025 and 0.0054‰). All data are presented in the absolute reference scale (CDES) 122 using the permil (‰) unit. Reliability of the measurements was assessed based on the Δ_{48} offset 123 and 49 parameter (cf. Huntington *et al.*, 2009). Δ_{47} values were converted into temperatures 124 using the Kluge *et al.* (2015) calibration, reprocessed using the "Brand" parameters for ¹⁷O 125 correction (Daëron et al., 2016). The Kluge et al. (2015) calibration, which is based on calcite, 126 has been shown to be indistinguishable from several other calibrations (Petersen *et al.*, 2017), 127 including calibrations based on dolomite (Bonifacie et al., 2017). Temperatures are reported at 1 128 standard error (SE) precision and at 95% confidence level (CL).

129

130 **RESULTS**

131 Dolomite samples display zebra or breccia textures. Zebra textures (figure 3A) have been 132 formerly interpreted as alternations of dark gray replacive and white dolomite cement filling 133 voids developed by focused flow of overpressured basinal fluids (Nielsen et al., 1998; 134 Vandeginste et al., 2005). The dolomite veins making up zebra sets in the samples used for this 135 study consist of thin rims made up of idiotopic white dolomite crystals (figure 3B). They show a 136 mottled appearance under the cathodoluminescence microscope, except for a last growth zone 137 exhibiting a dull luminescence (figure 3C). Breccia occurs as pockets (figure 3D) composed of 138 angular fragments of replacive dolomite in a matrix of dolomite and calcite cement (figure 3E). 139 They are mostly clast-supported, with thick dolomite cement rims which show a luminescence

140 similar to the rims observed in zebra veins (figure 3F). Individual dolomite crystals are larger 141 compared to those making up the zebra veins, though they show stepwise crystal terminations. 142 The dull rims in both zebra and breccia textures show signs of dedolomitization, and were 143 avoided during sampling prior to measurements. Two breccia samples (samples 202 and 144 CaBD2B) are coarse crystalline and show an orange weathering patina (figure 3G), which 145 suggests a strong enrichment in Fe. They are made up of replacive dolomite with large crystal 146 sizes (around 250 µm; figure 3H) and abundant development of saddle dolomite crystals (up to 147 2000 µm; figure 3I). Both replacive and void-filling dolomite in these samples are non-148 luminescent.

FIs were carefully selected to avoid analysis of secondary inclusions. Primary inclusions, with sizes generally between 4 and 10 μ m, were selected from petrographically associated assemblages (FIA; Goldstein and Reynolds, 1994) and from growth zones within individual crystals. T_h modes broadly correspond to those obtained by Gasparrini *et al.* (2006b) and range between 95 and 145°C (figure 4; table 1). Frequency histograms of T_h data for individual samples show unimodal distributions.

155 Clumped isotope Δ_{47} temperatures obtained from dolomite crystals range between 107 and 168 156 °C (figure 4; table 1). For 8 out of 9 samples investigated, the T Δ_{47} are higher compared to T_h-157 values. External precisions range between ±1 and ±8°C (±1SE). The error margins at 95% CL 158 are on average 3 to 4 times higher compared to SE, but could be reduced with additional 159 measurements.

160

161 **DISCUSSION**

162 Fluid inclusions *versus* clumped isotopes

163 Several sources of error can influence both FI and CI datasets and can result in significant 164 variations. Fluid inclusions should be primary and not affected by post-entrapment processes 165 such as thermal reequilibration or crystal deformation (Goldstein and Reynolds, 1994). The 166 carbonate crystals should not be affected by solid-state diffusion of O and C atoms, which could 167 induce an overestimation of the clumped isotope signature (Came et al., 2017). Solid-state 168 diffusion is a molecular-scale diffusion process whereby atoms can migrate through a crystal 169 when it is exposed to high temperatures (Dennis and Schrag, 2010; Passey and Henkes, 2012). 170 Temperatures necessary for diffusion likely approach 300°C for dolomite (MacDonald et al., 171 2017; Lloyd et al., 2017). Hydrothermal dolomitization was the hottest event affecting the 172 Paleozoic rocks in the CZ (Carrière, 2006) excluding the possibility of solid-state diffusion.

173 A secondary cause which could account for part of the temperature variation between clumped 174 isotopes and FIs is the difference in the sample volumes for both techniques. Powder samples 175 used for clumped isotope analyses were drilled from dolomite crystals visible on rock slabs and 176 are inevitably derived from a suite of crystals (Defliese and Lohmann, 2015). FI measurements, 177 on the other hand, allow obtaining T_h-values for individual crystals and growth zones. 178 Hydrothermal systems are known to be episodically very dynamic (Eichhubl and Boles, 2000), 179 which may account for rapid variations in temperature. FI data obtained from individual crystals 180 might therefore differ from clumped isotope data which give an "averaged" temperature. 181 Analytical improvements resulting in increasingly small sample sizes will allow more precise 182 clumped isotope measurements (Hu et al., 2014; Müller et al., 2017).

183 Geobarometry

The CI dataset yields crystallization temperatures between 107 and 168 $^{\circ}$ C, which can be plotted on the isochores constructed for each sample based on their T_h mode and composition of their

186 FIs (figure 5). Corresponding pressures range between 0 and 72 MPa. Void-filling dolomite 187 cement from breccia samples is characterized by low pressures (0-8 MPa), while dolomite 188 crystals in zebra veins formed at higher pressures, between 29 to 72 MPa. The lower pressures 189 recorded by breccia cements could be related to an abrupt decrease in fluid pressure that results 190 from the creation of open space following hydraulic brecciation (Knipe, 1993). The massive 191 nature of the cements and their imperfect crystal morphologies (e.g. stepwise crystal 192 terminations) support a model of fast precipitation in response to a sudden decrease in fluid 193 pressure. The higher overpressures recorded for dolomite crystals forming zebra veins support 194 the idea that zebra textures form during a pre-seismic buildup of suprahydrostatic to 195 supralithostatic pressures (Nielsen et al., 1998; Muchez and Sintubin, 2002). These 196 overpressures are also in agreement with the relatively new view on zebra textures introduced by 197 Merino et al. (2006), stating that zebra textures are self-organized, pushing aside replacive 198 dolomite through pressure-dissolution. Dolomite rhombs displace the host constituents due to 199 local induced stress generated by crystal growth. This induced stress can reach 47 MPa for 200 dolomite crystals growing at 100°C (Merino et al., 2006). The breccia samples (CaBD2B and 201 202) composed of coarse saddle crystals lacking luminescence are characterized by higher 202 pressures compared to pristine breccia (figure 5). These two samples might originate from a 203 separate episode of faulting and expulsion of dolomitizing fluids resulting in coarse crystalline 204 dolomitization or recrystallization of the initial dolomite rocks. The observed temperatures and 205 pressures are thus not representative of the initial crystallization process, but of a later pulse of 206 dolomitization or recrystallization.

207 The dataset used in this study illustrates the potential power of coupled T_h -T Δ_{47} geobarometry in 208 carbonate minerals, suggesting a high pressure for zebra dolomite formation in the study area.

Additional measurements on dolomite exposures in different geodynamic settings would allow
 generalization towards regional processes.

211

212 CONCLUSIONS

213 Hydrothermal dolomite samples from the Cantabrian Zone (northern Spain) have been used to 214 test a new geobarometry approach by combining fluid inclusion and clumped isotope 215 thermometry. This approach allows an estimation of the pressure during crystallization of 216 carbonate minerals, without the need for cogenetic generations of FIs or the use of independent 217 geothermometers. Clumped isotope temperatures (107-168 °C) of dolomite crystals from zebra 218 and breccia textures are higher compared to fluid inclusion homogenization temperatures from 219 the same samples (95-145 °C). The offset between these values mainly depends on the pressure 220 during FI entrapment. Zebra dolomite likely formed during pre-seismic pressure buildup (29-72 221 MPa), while dolomite cement in breccia samples precipitated in response to a sudden decrease in 222 fluid pressures upon fault rupture. Geobarometry based on fluid inclusion and clumped isotope 223 thermometry is not only a promising tool for carbonate diagenesis studies, but could prove useful 224 for a broad range of geological applications.

225

226 ACKNOWLEDGEMENTS

R. Honlet is supported by Research Foundation - Flanders. C. John acknowledges support from
the Qatar Carbonate and Carbon Storage Research Center for the clumped isotopes laboratory,
funded by Qatar Petroleum, Shell, and the Qatar Science and Technology Park. E. Bemer
(IFPEN) is thanked for supporting the involvement of M. Gasparrini.

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- 365

366 FIGURE CAPTIONS

Figure 1. Water phase diagram illustrating the principle of geobarometry based on fluid inclusion T_h and clumped isotope $T\Delta_{47}$ measurements. With known fluid composition and with $T\Delta_{47}$ representing the trapping temperature of fluid inclusions (T_t), the pressure of carbonate crystallization can be obtained from ΔT .

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Figure 2. Structural sketch map of the Bodón Unit adapted from Alonso *et al.* (2009) and Muñoz
Quijano (2015). The map shows the locations of samples analyzed in this study. Inset shows
location of the Bodón Unit on a map of the Iberian Peninsula.

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377 Figure 3. (A) Outcrop photograph of zebra textures developed in well-bedded dolomitized 378 mudstones. (B) Zebra texture displayed on polished and stained hand specimen. Calcite is 379 stained in red. (C) Cathodoluminescence image of dolomite crystals making up the zebra texture. 380 Orange calcite cement is visible in the upper part of the image. (D) Outcrop photograph of a 381 breccia pocket in wel-bedded dolomitized mudstones. Bedding surfaces are visible in the upper 382 part of the photograph (indicated with black dashed lines). Hammer for scale (black arrow). (E) 383 Breccia texture displayed on polished and stained hand specimen. Calcite is stained in red. (F) 384 Cathodoluminescence image of dolomite cement in breccia. Note the large crystal size and 385 stepwise crystal terminations (white arrows). (G) Outcrop photograph of bedding surface of 386 coarse crystalline and Fe-rich dolomite. (H) Breccia texture displayed on polished hand 387 specimen. (I) Photomicrograph (crossed polarized light) of saddle dolomite crystals in coarse 388 crystalline and Fe-rich dolomite.

Figure 4. Graph showing relationship between the modes of fluid inclusion homogenization temperatures (T_h) and clumped isotope (Δ_{47}) temperatures. T_h-values range between 95 and 145°C. Error bars correspond to Q1 – Q3 ranges (25th – 75th percentiles; see table 1). Δ_{47} temperatures range between 107 and 168°C. Error bars correspond to ±1SE (see table 1). The black dashed line represents a 1:1 fit between both temperatures, while the blue dashed lines represent offsets in Δ_{47} temperature (A: +10°C; B: +20°C; C: +30°C). 'ReXX' refers to recrystallized dolomite.

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Figure 5. Pressure-temperature phase diagram of saline water. T_h modes obtained for each sample are plotted on the liquid-vapor curve (dashed line), and corresponding isochores have been drawn. $T\Delta_{47}$ values are reported along the isochores of the corresponding samples.

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412 **FIGURES:**

FIGURE 1:





- **FIGURE 2:**



FIGURE 3:





TABLE CAPTION:

470 Table 1. Results from fluid inclusion microthermometry and clumped isotope paleothermometry. 471 $T\Delta_{47}$ has been calculated using Kluge *et al.* (2015) reprocessed with new parameters for ¹⁷O 472 correction (Daëron *et al.*, 2016). Pressure values have been calculated with the equation of state 473 from Zhang and Frantz (1987). 474

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476 **TABLE:**

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		Th												
		(°C)	T _h (°C)	T _h (°C)	#	$\delta^{13}C$	δ ¹⁸ 0	Δ_{47}	Δ_{47}	$T \Delta_{47}$	±1SE	±1SD	±1CL	Р
Sample	Texture	mode	range	Q1 - Q3	inclusions	(‰VPDB)	(‰VPDB)	(‰CDES)	average	(°C)*	(°C)	(°C)	(°C)	(MPa)
001	Zebra	133	108-151	122-135	56	4.14	-10.17	0.466	0.462	149	8	14	34	31
								0.474						
								0.447						
028	Breccia	128	102-141	124-129	33	4.35	-9.58	0.495	0.489	125	3	6	14	0
								0.491						
								0.481						
166	Breccia	130	102-154	125-139	103	4.53	-8.24	0.482	0.48	133	4	8	14	7
								0.483						
								0.487						
								0.467						
173	Breccia	123	104-138	117-126	45	3.62	-9.52	0.495	0.489	126	7	16	20	9
								0.520						
								0.467						
								0.479						
								0.483						
202	Breccia	105	79-139	95-111	127	3.3	-6.06	0.489	0.494	121	2	3	7	37
								0.495						
								0.497						
205	Zebra	110	99-125	107-115	54	3.52	-11.3	0.494	0.489	125	4	6	54	34
								0.484						
243	Zebra	95	86-128	70-167	50	3.38	-8.34	0.515	0.513	107	1	1	3	29
								0.514						
								0.511						
45 [↑]	Zebra	100	88-119	98-106	52	3.64	-8.55	0.481	0.482	131	5	9	22	72
								0.471						
								0.493						
$CaBD2B^{\dagger}$	Breccia	145	129-160	138-148	41	4.23	-9.76	0.438	0.444	168	7	9	93	50
		-				-		0.450				-		

TABLE 1. TEMPERATURE AND CALCULATED PRESSURE VALUES

*Calculated using Kluge et al. (2015), reprocessed with new parameters for ¹⁷O correction (Daëron et al., 2016)

[†]Samples from Gasparrini *et al.* (2006a)