# Thermochemical Sulphate Reduction Can Improve Carbonate Petroleum Reservoir Quality

3 Lei Jiang<sup>1,2,3,4\*</sup>, Richard H Worden<sup>5</sup>, Changbing Yang<sup>3</sup>

<sup>1</sup>Key Laboratory of Petroleum Resources Research, Institute of Geology and
Geophysics, Chinese Academy of Sciences, Beijing 100029, China (email:
<u>lei.jiang@mail.iggcas.ac.cn</u>)

- <sup>7</sup> <sup>2</sup>Institutions of Earth Science, Chinese Academy of Sciences, Beijing 100029, China
- <sup>3</sup>Bureau of Economic Geology, Jackson School of Geosciences, The University of
- 9 Texas at Austin, Austin, TX 78713, USA

<sup>4</sup>Department of Geology and Geophysics, Yale University, New Haven, CT 06511,
USA

- <sup>5</sup>Department of Earth, Ocean and Ecological Sciences, University of Liverpool, 4
  Brownlow Street, Liverpool, L69 3GP, UK
- 14
- 15 Abstract
- 16

Interest in the creation of secondary pore space in petroleum reservoirs has increased 17 18 because of a need to understand deeper and more complex reservoirs. The creation of new secondary porosity that enhances overall reservoir quality in deeply buried 19 carbonate reservoirs is controversial and some recent studies have concluded it is not 20 21 an important phenomenon. Here we present petrography, geochemistry, fluid inclusion data, and fluid-rock interaction reaction modeling results from Triassic 22 Feixianguan Formation, Sichuan Basin, China, core samples and explore the relative 23 importance of secondary porosity due to thermochemical sulphate reduction (TSR) 24

25 during deep burial diagenesis. We find that new secondary pores result from the dissolution of anhydrite and possibly from dissolution of the matrix dolomite. 26 Assuming porosity before TSR was 16 % and the percentage of anhydrite was 6 %, 27 28 modelling shows that, due to TSR, 1.6 % additional porosity was created that led to permeability increasing from 110 mD (range 72 to 168 mD within a 95% confidence 29 interval) to 264 mD (range 162 to 432 mD within a 95 % confidence interval). 30 Secondary porosity results from the density differences between reactant anhydrite 31 and product calcite, the addition of new water during TSR, and the generation of 32 33 acidity during the reaction of new H<sub>2</sub>S with the siderite component in pre-existing dolomite in the reservoir. Fluid pressure was high during TSR, and approached 34 35 lithostatic pressure in some samples; this transient overpressure may have led to the 36 maintenance of porosity due to the inhibition of compactional processes. An additional 1.6 % porosity is significant for reserve calculations, especially considering 37 that it occurs in conjunction with elevated permeability that results in faster flow rates 38 39 to the production wells.

40

41 Keywords: Mesogenetic dissolution, thermochemical sulphate reduction, carbonate
42 reservoir, fluid inclusion, fluid pressure.

#### 43 Introduction

Thermochemical sulphate reduction (TSR) is the abiological oxidation of
hydrocarbons by sulphate (due to the dissolution of anhydrite, celestite and barite in
sedimentary basins) at elevated temperatures, typically higher than 110 °C (Jiang et al.,
2015c; Machel, 2001; Worden et al., 1995). Significant alteration of petroleum,
generation of reduced forms of sulphur (S and H<sub>2</sub>S), and oxidized forms of carbon

49	(carbonate minerals and CO <sub>2</sub> ) are typically the results of TSR. TSR can also generate
50	water, metal sulphides, organosulphur compounds, and bitumen (Bildstein et al., 2001;
51	Cai et al., 2003; Jiang et al., 2015c; Machel, 1987; Machel et al., 1995; Worden et al.,
52	2000; Worden et al., 1995).

53 A general reaction summarizing TSR:

sulphate+ petroleum 
$$\rightarrow$$
 calcite + H<sub>2</sub>S  $\pm$  H<sub>2</sub>O  $\pm$  CO<sub>2</sub>  $\pm$  S  $\pm$  altered petroleum (1)

Simple stoichiometric TSR reactions in many sedimentary basins (e.g. Feixianguan
Formation, Sichuan Basin; Khuff Formation, Abu Dhabi; Upper Devonian and
Mississippian strata, Alberta, Canada) between anhydrite and the two simplest
hydrocarbons:

59 
$$CaSO_4 + CH_4 \rightarrow CaCO_3 + H_2S + H_2O$$
 (2)

$$60 \quad 2CaSO_4 + C_2H_6 \rightarrow 2CaCO_3 + H_2S + 2H_2O + S \tag{3}$$

61 
$$7CaSO_4 + 4C_2H_6 \rightarrow 7CaCO_3 + 7H_2S + 5H_2O + CO_2$$
 (4)

62 Most TSR-related studies have focused on stable isotope geochemistry and the geochemistry of petroleum, sulphate, and the products of TSR (e.g. CO<sub>2</sub>, H<sub>2</sub>S, S), and 63 64 few focus on the ability of TSR to form new pore spaces in deeply buried rocks and thus improve carbonate reservoirs quality. For example, it is proposed that deep burial 65 diagenesis in a closed system (including TSR) is not able to enhance porosity and 66 permeability in carbonate reservoirs (Ehrenberg et al., 2012; Heydari, 1997; Machel 67 68 and Buschkuehle, 2008). In the Upper Jurassic Smackover Formation, Mississippi, TSR appears to have resulted in large amount of calcite precipitation with significant 69 porosity-loss (Heydari, 1997). Carbonate reservoirs in the Southesk-Cairn Carbonate 70

71 Complex (SCCC), Alberta Basin, indicate the overall change of porosity and permeability during TSR is small. Similarly, based on a literature review and 72 modelling, Ehrenberg et al. (2012) suggested that mesogenetic dissolution, producing 73 74 a net increase in secondary porosity, is not likely during deep burial diagenesis. By contrast, detailed petrographic, geochemical, and modelling work (TSR impact on the 75 carbonate reservoir quality) from the Western Canada Sedimentary Basin (WCSB) 76 determined that while 75% of the porosity created by dissolution of anhydrite during 77 TSR was lost due to calcite precipitation, net porosity increased 1 to 2% (Hutcheon 78 79 and Krouse, 1994; Hutcheon et al., 1995).

There is evidence for extensive TSR in the Feixianguan Formation, based on: (1) 80 81 coexistence of TSR calcite, elemental sulphur, pyrite, sulphur-enriched bitumen, and anhydrite (Jiang et al., 2014a); (2) broad overlap of sulphur isotopes of TSR-82 generated pyrite, elemental sulphur, sulphur in bitumen and H<sub>2</sub>S, with values 83 approaching  $\delta^{34}$ S value of the pre-existing, early diagenetic anhydrite that is 84 equivalent to coeval seawater sulphur isotopes (Cai et al., 2010; Hao et al., 2008; Zhu 85 et al., 2005); (3) relatively high fluid inclusion homogenization temperatures found in 86 TSR calcite (from ~110 °C to 220 °C) (Jiang et al., 2014a, 2015c), and; (4) gas 87 geochemical characteristics including: relatively high H<sub>2</sub>S concentration (up to 58%) 88 that replaced the CH<sub>4</sub> in gas reservoirs, the extremely high gas dryness  $(C_1/\Sigma C_n)$ 89 90 (>0.95), positive relationship of gas souring index [GIS =  $H_2S/(H_2S + \sum Cn)$ ] (Worden et al., 1995) and TSR extent parameter  $[CO_2/(CO_2 + \sum C_nH_{2n+2})]$  (Krouse et al., 1988), 91 and the  $\delta^{13}$ C values of methane and ethane with a different extent of TSR (Cai et al., 92 2013; Hao et al., 2008; Li et al., 2005; Liu et al., 2013; Liu et al., 2014). 93

94 The Feixianguan Formation in the Sichuan Basin, China, offers an ideal place to study the impact of TSR on deeply buried carbonate reservoir quality, because it was a 95 relatively closed diagenetic environment during TSR (Jiang et al., 2015a; Jiang et al., 96 97 2015c). There is little evidence for diagenetic fluids (e.g., meteoric water, hydrothermal fluids, deep basinal fluids) influencing the carbonate reservoir during 98 99 TSR. While a number of studies have focused on TSR in the Feixianguan Formation, only two have investigated the impact of TSR on porosity: Cai et al. (2014) suggested 100 101 TSR was responsible for dissolution of dolomite and anhydrite with an overall 102 positive effect on dolostone reservoir quality; by contrast, Hao et al. (2015) suggested TSR diagenesis was dominated by calcite cementation that reduced reservoir quality 103 104 in the Feixianguan Formation. This study will focus on resolving this conflict and 105 establishing the impact of TSR on carbonate reservoirs using the Feixianguan 106 Formation, by addressing the following questions:

107 1. Did new secondary pores develop in the Feixianguan Formation during TSR?

108 2. If secondary pores did form during TSR, what are their characteristics?

109 3. What are the possible mechanisms for enhanced reservoir quality during TSR?

To answer these questions, we focus on representative wells which have experienced
TSR by applying conventional core description techniques, point counting,
transmitted-light petrography, scanning electron microscope (SEM) petrography,
fluid-inclusion analysis and pressure modelling, and diagenesis modelling.

## 114 Geological setting

The diamond-shaped, intracratonic, 230,000 km<sup>2</sup> Sichuan Basin is located in the east
of Sichuan Province, southwest China (Fig. 1A). The Sichuan Basin is tectonically-

bounded by the Longmenshan fold belt in the northwest, the Micangshan uplift in the
north, the Dabashan fold belt in the northeast, the Hubei-Hunan-Guizhou fold belt in
the southeast, and by the Emeishan-Liangshan fold belt in the southwest (Fig. 1A).

The Lower Triassic Feixianguan Formation  $(T_1f)$  carbonates occupy high energy 120 facies belts dominated by oolitic carbonates (Fig. 1B) (Ma et al., 2008a; Zhao et al., 121 122 2005). During deposition of the Feixianguan Formation, open to semi-restricted 123 carbonate platforms, in an arid climate, with fluctuating sea level, resulted in deposition of oolitic shoals and multiple gypsiferous layers on the margins of 124 125 Kaijiang-Liangping Bay. Towards the end of Feixianguan deposition, aridity increased and the platforms became more restricted, which resulted in multiple 126 gypsum beds interlayered with thin-bedded micritic limestones (Fig. 2). These upper 127 evaporitic layers provide the regional seal for the underlying carbonate reservoirs 128 129 (Zhao et al., 2005).

Seawater evaporation from lagoons, on the restricted platform to platform margin, is 130 reported to have caused the initial dolomitization in the Feixianguan Formation (Jiang 131 132 et al., 2013; Jiang et al., 2014b; Zhao et al., 2005). The most significant dolomitization phase then occurred in shallow burial environments by reflux of 133 mesohaline water or seawater dolomitization (Jiang et al., 2014b). The latest 134 dolomitization event occurred during burial, at temperatures ranging from 80°C to 135 140°C, due to invasion of high salinity water, most likely from the overlying 136 Jialingjiang Formation (Jiang et al., 2014b). The Feixianguan Formation on the NE 137 138 side of the Kaijiang-Liangping Bay is more dolomitized compared to the SW side, perhaps because of local differences in aridity during deposition (Jiang et al., 2014b; 139 Zhao et al., 2005). 140

141 Burial histories in the Sichuan Basin were dominated by rapid burial in the Triassic and early Jurassic to depths of ~7500 m and temperatures of 220°C followed by uplift 142 prior to the Cretaceous Yanshan movement. This history has led to early Triassic gas 143 reservoirs with temperatures between 100 and 140°C at the present day (Ma et al., 144 2008a). Thermochemical sulphate reduction (TSR) occurred during appropriate 145 elevated temperature conditions, where there was sufficient supply of anhydrite and 146 147 petroleum. The Feixianguan Formation carbonate has variable H<sub>2</sub>S concentrations generally between 10% and 20%, but up to 60% in some reservoirs (Cai et al., 2013; 148 Hao et al., 2008; Li et al., 2005; Liu et al., 2014). 149

#### 150 Methods

More than 100 core samples were collected from the lower unit of the Triassic 151 Feixianguan Formation from the Puguang, Luojiazhai, Dukouhe, Maoba and Jingzhu 152 carbonate reservoir gas fields. These fields were chosen since they have variable 153 degrees of gas sourness. Polished sections, and conventional thin sections that were 154 stained with Alizarin Red S to distinguish calcite and dolomite, were prepared from 155 156 all samples. Selected samples were examined by scanning electron microscope (SEM) in backscattered electron imaging mode (BSEM), and elemental analysis by energy 157 158 dispersive X-ray spectroscopy (EDS). Point counting of the stained thin sections (500 points for each sample) was used to determine the mineral and pore-type proportions. 159

Fluid inclusion homogenization temperatures ( $T_h$ ) and last ice melting temperatures ( $T_m$ ) were measured from fluid inclusion assemblages (FIAs) containing two-phase aqueous inclusions in five doubly polished, detached (50 to 60 µm thick) wafers. The use of FIAs to determine temperatures of mineral growth, as opposed to single inclusions, provides confidence that the  $T_h$  data are credible and minimizes the effects of artefacts, such as thermal re-equilibration (Goldstein, 2012; Goldstein and
Reynolds, 1994). Fluid inclusion microthermometry was conducted using a Zeiss
Axioskop 40A Pol light microscope with a Linkam THM600/TS90 heating and
cooling stage. Last ice melting temperatures were converted to salinity using standard
equations (Bodnar, 2003; Oakes et al., 1990).

170 Coexistence of two-phase aqueous and single-phase hydrocarbon inclusions indicates 171 that fluid inclusions were trapped in the immiscible two-phase field, and measured 172 aqueous inclusion homogenization temperatures thus represent trapping temperatures 173 (Goldstein and Reynolds, 1994). Raman analysis was undertaken for individual hydrocarbon fluid inclusions using LabRAM ARAMIS equipment. The position of 174 each measured Raman line was determined, after baseline correction, using 175 parameters for Gaussian/Lorentzian peak fitting (Lin et al., 2007). The 2851.38 and 176 2972.44 cm<sup>-1</sup> Ne lines (relative to the 514.529 nm Rayleigh line of the Ar ion laser, in 177 178 air) were used for calibration. Concentrations of methane and trapping pressures were calculated following the equations of state for the CH<sub>4</sub> system and the H<sub>2</sub>O-NaCl-CH<sub>4</sub> 179 system (Duan and Mao, 2006; Duan, 1992), using the approach of Becker et al. 180 181 (2010).

The aim of geochemical modelling in this study is to simulate the overall change in mineral assemblage volume (including dolomite, calcite, and anhydrite) by considering the reaction described in equation (2) during TSR. A similar modelling strategy to the 3D reactive transport presented by Fu et al. (2016) was applied in this study. In contrast to Fu's model, our geochemical model focuses on geochemical reactions without considering groundwater flow and solute transport during the TSR. This simplification renders fast computation. The US Geological Survey's computer 189 program PHREEQC and the database thermodynamic wateq4f.dat were used to simulate equilibrium reactions for aqueous species (Parkhurst and Appelo, 2013). 190 Temperatures, pressures, and fluid compositions obtained from the fluid study were 191 192 used as input to the geochemical model (details in the supplemental input file). In addition, the kinetic rates of TSR were obtained from the model presented by Fu et al. 193 194 (2016). Mineral proportions were derived from point counting of the thin-section samples. The modelling results for volume changes in mineral compositions were 195 further used to estimate change in porosity following the similar method presented by 196 197 Yang et al. (2008) and (Xu et al., 2010).

Permeability modelling was achieved by obtained the equation of relationship between porosity and permeability, which can be obtained from the measured core plug petrophysical data. The changes of permeability during TSR were calculated by inputting the modelled porosity change during TSR to the calculated porositypermeability relationship equation, using 95 % confidence limits to present ranges for the derived permeability values.

#### 204 **Results**

#### 205 Thermochemical sulphate reduction diagenetic minerals

Dolomite crystal sizes typically range from 50 to 200 µm in Feixianguan Formation sucrosic dolostone reservoirs (Fig. 3). Here we illustrate that late diagenetic TSR calcite locally fills pores but is heterogeneously distributed in these dolostone reservoirs, with some parts of the reservoir filled with calcite whereas other parts contain open pores (Fig. 3A, C; Fig. 4B, D; Fig. 5C). Pore characteristics have been strongly modified by diagenetic processes, with some pores occupied by diagenetic minerals and some enlarged by carbonate dissolution (Fig. 3B, D; Figs. 4 and 5). 213 Calcite produced by TSR represents the most volumetric mineral during TSR (in contrast to elemental sulphur), with average point-counted volumes of  $1.6 \pm 3.4$  % 214 (N=44) and a maximum volume of up to 17 % (Table 1). Two types of TSR calcite 215 216 have been identified: TSR calcite that contains oil or bitumen inclusions is here defined as oil-stage TSR calcite, and TSR calcite with no evidence of oil or bitumen 217 inclusions is defined as gas-stage TSR calcite. These calcites may either have 218 219 precipitated from fluids in open pores or replaced the former anhydrite cement within the reservoir (Jiang et al., 2014a). 220

Euhedral pyrite is locally present in pores either as replacement of dolomite or in 221 222 pores that are also typically associated with elemental sulphur (Fig. 4A). The average point counted volume of pyrite is less than  $0.4 \pm 0.9$  % (N=44) (Table 1). Elemental 223 sulphur in dolostone reservoir is best shown using BSEM analysis and commonly 224 225 occurs with pyrite and TSR calcite (Figs. 3, 4, and 5). The total point counted volume of sulphur is less than 0.1% (N=44) although it is locally enriched at the edges of 226 227 dissolution-enlarged pores. Elemental sulphur is appears similar to bitumen in the 228 optical microscope (opaque and irregular in form). Bitumen is commonly present as sheet- and sphere-shape masses, with length or diameter in a range from 10 to 100 µm 229 230 (Figs. 3 and 5), and represents  $4.4 \pm 5.4$  % (N=44) rock volume (Table 1).

231 **Dissolution-enlarged pores** 

Pore spaces in the Feixianguan Formation are dominated by dissolution-enlarged pores, which are commonly associated with TSR calcite, elemental sulphur, pyrite and bitumen (Figs. 3 and 45). There are two main occurrences of dissolution-enlarged pores in the Feixianguan dolostone reservoirs. The first type is selective dissolution pores. These are represented either by solution-enhanced vugs, where moldic pores

have been enlarged, or by complete dissolution of anhydrite cement; these dissolution pores are relatively large (up to 2 mm) (Fig. 4; Fig. 5A, B). The second type is characterised by partial or complete dissolution of coarse crystalline dolomite (Fig. 5D). Elemental sulphur commonly occurs on the edges of dissolution pores but there is notably little TSR calcite present (Figs. 3 and 5). The point counted (meso) porosity for these good dolostone reservoirs is in the range from 0% to 33 %, average at 9.7 ± 7.5 % (N=44) (Table 1).

## 244 Fluid inclusion microthermometry

Calcite produced by TSR contains primary, two-phase aqueous inclusions. Fluid
inclusion homogenization temperature (T<sub>h</sub>) variations within fluid inclusion
assemblages (FIAs) from TSR calcite are generally less than about 10 °C (Fig. 6).
Salinity values, determined from last ice melting temperatures, vary from less than 5
wt % NaCl to nearly 25 wt % NaCl for TSR calcites.

Measured homogenization temperatures (T<sub>h</sub>) for FIAs (and isolated fluid inclusions) 250 for oil-dominated TSR calcite range from approximately 110 °C to 200 °C (Fig. 6). In 251 contrast, gas-dominated TSR calcite has a T<sub>h</sub> range from 135 °C to 210 °C (Fig. 6). 252 253 The temperatures obtained from the inclusions in TSR calcite represent minimum trapping temperatures (Goldstein and Reynolds, 1994). Oil-stage TSR calcite has 254 decreasing salinity with increasing temperature, while, in contrast, gas-stage TSR has 255 256 decreasing salinity with decreasing temperature (Fig. 6). The T<sub>h</sub>-salinity distribution defines a progressive evolution during burial and heating, initially in the presence of 257 oil with ever-falling salinity, and then in the presence of gas once the oil underwent 258 cracking at maximum burial, followed by uplift and cooling in the presence of the 259 evolved gas charge (Fig. 6). 260

#### 261 Methane concentrations of fluid inclusions and trapping pressures

Aqueous fluid inclusions in TSR calcite in the Feixianguan Formation have bulk 262 methane concentrations over a wide range from  $\sim 1500$  to 11000 ppm, corresponding 263 to trapping pressures of ~ 25 to 165 MPa (Fig. 7). By reference to the large-scale 264 thermal cycle revealed by the oil-stage TSR calcite and then the gas-stage TSR calcite 265 (Fig. 6), it is possible to infer the pressure evolution of the Feixianguan Formation 266 267 during TSR. The fluid inclusions show variations in pressure with temperature with a clear subdivision of the oil-stage and gas-stage TSR calcites. The increasing 268 temperature for the oil-stage calcite reveals increasing fluid pressure with time (and 269 270 with heating) as the Feixianguan Formation was buried to ~7000 m (temperature of  $\sim$ 220°C). The switch to gas-stage calcite at elevated temperature shows that uplift to 271 the current burial depth of ~3000 to 5000 m (temperatures of ~ 120 to 140 °C) was 272 accompanied by decreasing fluid pressure with time (and with cooling). The highest 273 fluid pressures approach the simulated trapping pressures (153-160MPa) for high 274 density methane inclusions reported elsewhere for the Feixianguan Formation (Liu et 275 al., 2009). Fluid pressures approach, and exceed in some cases, the modelled 276 277 lithostatic pressure gradient suggesting that at least some of the fractures present in 278 these rocks maybe due to excess fluid pressure.

## 279 Thermochemical sulphate reduction modelling result

Prior to TSR, calcite and anhydrite are here interpreted to have been abundant
minerals in the dolostone Feixianguan Formation reservoirs (representing 1 and 6
wt. %, respectively) with porosity of 16% (Table 2), based on the point count data.
The gas phase in these reservoirs is dominated by CH<sub>4</sub> (95 vol. %), CO<sub>2</sub> (2 vol. %),
and N<sub>2</sub> (1.5 vol. %). The initial composition of pore water before the onset of TSR in

the geochemical model has been calculated by equilibrating of seawater through evaporation to the salinity obtained from fluid inclusion data. Temperature in the geochemical model was varied from 120 to 200 °C, based on the fluid inclusion evidence (Fig. 6).

Relative changes in mineral volumes for anhydrite, calcite, and dolomite at the steady 289 state, based on the PHREEQC modelling results, are listed in Table 2. Anhydrite is 290 consumed due to TSR, resulting in decreasing solid rock volume by 312.4 cm<sup>3</sup> (6.00 % 291 292 of the total rock volume), whereas carbonate minerals precipitated and increased the solid rock volume by 212.8 cm<sup>3</sup> due to calcite (4.05 % of the total rock volume) and 293 1.3 cm<sup>3</sup> rock volume due to dolomite (0 % of the total rock volume). Hence there was 294 a 1.95 % increase in relative porosity equating to an absolute post-TSR porosity 295 increase of 1.6 %. 296

297 Sensitivity modelling of initial porosity of 10 % before TSR was also conducted; the298 results show that porosity increased by 1.7 % due to TSR (Table 2).

## 299 **Permeability modelling result**

An equation of the relationship between porosity and permeability was calculated based on the core plug petrophysical data:  $Y = 0.0166e^{0.5498X}$ ,  $R^2 = 0.6489$  (where Y and X stands for mD-permeability and percentage porosity, respectively).

The modelled permeability changes during TSR in the most proper porosity change in

Case 1 (16 to 17.6 % porosity-increase due to TSR) is from 110 mD (before TSR,

with a range 72 to 168 mD within a 95% confidence interval) to 264 mD (after TSR

with a range 162 to 432 mD within a 95% confidence interval). For case 2 (10 to 11.7 %

307 porosity-increase due to TSR) is from 4.1 mD (before TSR) to 10.3 mD (after TSR)308 (Table 3).

309 **Discussion** 

## 310 Conditions during TSR calcite growth

311 Comparison of the homogenization temperatures and salinities reveals a distinct separation that also relates to the type of host TSR calcite (Fig. 6). Oil was present in 312 the reservoir before gas became the dominant petroleum phase, so initial calcite 313 314 growth started in the presence of both oil and high salinity waters (25 wt % NaCl). Formation water salinity decreased to about 9 wt % NaCl during increasing 315 temperature as oil-induced TSR progressed. After the oil charge evolved to a gas 316 charge at elevated temperature, uplift occurred accompanied by decreasing 317 temperature; the salinity of the formation water continued to fall (to 5 wt % NaCl) at 318 319 this stage. The increasing and then decreasing temperatures in TSR calcites from the Feixianguan Formation are due to burial and uplift rather than invasion and 320 321 subsequent cooling of hydrothermal fluids, (Jiang et al., 2015a; Jiang et al., 2014a).

The measured homogenisation temperature data from TSR calcite (Fig. 6) have been 322 related to the derived trapping pressures (Fig. 7B). A modelled lithostatic pressure 323 324 gradient of 25.4 MPa/km was calculated for a depth-averaged rock density of 2.59 g/cm<sup>3</sup>, using wireline log data from the Sichuan Basin. A modelled hydrostatic 325 pressure gradient was calculated as 9.9 MPa/km, based on a water density of 1.02 326 327  $g/cm^3$ . Depths for the lithostatic and hydrostatic pressure gradients have been converted into temperature assuming a geothermal gradient of 24 °C/km (Liu et al., 328 2016; Qiu et al., 2008). Broadly speaking, the calculated fluid inclusion trapping 329 pressures at the time of TSR calcite growth represent pore-fluid pressures that were 330

significantly above the hydrostatic pressure gradient (Fig. 7B). Many of the pore-fluid
pressure measurements are near lithostatic pressures; some of the low pressuretemperature points lie at, or just below, the hydrostatic gradient (Fig. 7B).

## 334 Impact of thermochemical sulphate reduction on reservoir quality

According to petrographic observations reported here, dissolution-enhanced pore 335 spaces tend to be spatially-close to TSR-related diagenetic minerals (pyrite, elemental 336 sulphur, calcite) (Figs. 3 and 45). The close spatial association suggests that this 337 secondary porosity, with point counted porosity values up to 32 % in one thin-section, 338 339 formed by TSR. Our modelling work (Table 2) suggests that a porosity gain of 1.6 % 340 is a consequence of TSR in the Feixianguan Formation, mainly due to the dissolution of anhydrite cement and/or nodules (Jiang et al., 2014b). This is different from the 341 TSR modelling by Fu et al. (2016), within which the sulphate for TSR was derived 342 343 from anhydrite seal, and porosity should be decreased by TSR-calcite precipitation in their model. Note that Hao et al. (2015) suggested, instead, that calcite cementation, 344 rather than anhydrite and carbonate dissolution, dominated TSR diagenesis in the 345 Feixianguan Formation, leading to porosity-loss during TSR. Our petrographic 346 observations lead to a different conclusion. Porosity related to TSR dissolution has 347 348 likely been underestimated by Hao et al. (2015) because they did not provide a detailed paragenetic sequence and also failed to discriminate TSR calcite from other 349 types of diagenetic calcite. The diagenetic environments, reactants and products of 350 351 TSR, and the gas geochemistry characteristics of the Feixianguan Formation are similar to those in the Western Canada Sedimentary Basin (Cai et al., 2014; Hao et al., 352 2008; Hutcheon et al., 1995; Jiang et al., 2015c; Zhu et al., 2005). Significantly, TSR 353 modelling of the Western Canada Sedimentary Basin, by Hutcheon et al. (1995), also 354

355 showed that TSR probably led to an increase in porosity by 1 to 2 %, concurring with 356 model outputs from the Feixianguan Formation. The permeability modelling result 357 suggests that a net porosity increased from 1.6 % to 1.7 % by TSR, which doubled the 358 reservoir permeability (Table 3), thus resulting in significantly enhancement of 359 reservoir deliverability.

360 Elemental sulphur and calcite produced by TSR, as well as the TSR by-product 361 pyrite, , typically occur at the edge of dissolution enlarged pores (Figs. 3 and 5). 362 Elemental sulphur is routinely present in appreciable quantities in sour gas and oil 363 reservoirs in the Arabian Gulf region and United Arab Emirates (Abou-Kassem, 2000), as well as in the Smackover Formation in southern Mississippi (Kuo, 1972). 364 The freezing point of elemental sulphur, at atmospheric pressure, is 119 °C. Elemental 365 sulphur and H<sub>2</sub>S are miscible at high pressure allowing dissolution of H<sub>2</sub>S into the 366 liquid sulphur phase, and vice versa. Hence, it is likely that elemental sulphur is 367 368 present as a liquid (or is dissolved in H<sub>2</sub>S-rich gas) under reservoir temperatures and pressures conditions during and after TSR (Meyer, 1976). Elemental sulphur in the 369 Feixianguan Formation may have been present either at the contact between local 370 371 (pore-scale) oil-water or gas-water contacts with the host dolomites in the dissolution enlarged pores, possibly inhibiting calcite growth on surface of dolomite crystals (Fig. 372 3D; Fig. 5). 373

374 Pyrite was formed during TSR either via equation (5) or via equation (6) in the
375 Feixianguan Formation (Fig. 4A) (Jiang et al., 2014a; Liu et al., 2013).

 $376 \quad \mathrm{Fe}^{2+} + 2\mathrm{H}_2\mathrm{S} \to \mathrm{Fe}\mathrm{S}_{2^+} + 4\mathrm{H}^+ \tag{5}$ 

 $377 \quad \text{FeCO}_3 + 2\text{H}_2\text{S} \rightarrow \text{FeS}_2 + \text{CO}_2 + \text{H}_2\text{O} + \text{H}_2 \tag{6}$ 

378 
$$CaMg(CO_3)_2 + 4H^+ \rightarrow Ca^{2+} + Mg^{2+} + 2H_2O + 2CO_2$$
 (7)

379 For equation (6), we are not suggesting that pure siderite (FeCO<sub>3</sub>) was present in the 380 rock, rather that this represents the ferroan-component in dolomite, with iron likely 381 being originally sourced from anoxic dolomitizing fluids. Point counted volumes of 382 pyrite in the Feixianguan Formation range from 0 to 4 % (average of 0.4 %). Hence, 383 at least near to the site of pyrite precipitation-- equation (5), the acidity of diagenetic 384 fluids must have been increased by the release of H<sup>+</sup>. As a consequence, carbonate 385 dissolution is likely to have occurred, via equation (7), during and after pyrite 386 precipitation in the Feixianguan Formation as a result of acid-creating equation (5) or via equation (6). Our modelling results show that TSR processes have not changed the 387 volume of dolomite in these reservoirs (Table 2), although dolomite dissolution 388 occurred after TSR. As mentioned above, this is consistent with isotopically-heavy 389 CO<sub>2</sub> found in the Feixianguan Formation. Calcite produced by TSR has not been 390 391 observed in direct association with pyrite and elemental sulphur in the Feixianguan 392 Formation suggests that the slow rate of diffusion exceeds the rate of TSR. This may also suggest that the components required to make TSR calcite may have been 393 394 transported away from the immediate reaction site and TSR in porous carbonates may be able to proceed more efficiently than TSR in a finely crystalline dolomite matrix 395 (with low porosity and low permeability) (Jiang et al., 2014a; Worden et al., 2000). 396 397 Calcite precipitation rates are slower than pyrite precipitation rates (Fu et al., 2016), 398 and therefore it is possible that TSR calcite has been transported into other parts of the 399 Feixianguan dolostone reservoirs via diffusion, fractures and/or faults formed by local 400 tectonic movements, whereas pyrite growth may have occurred close to the original TSR site (Ma et al., 2008b). Although precipitation of 0.4% TSR-derived pyrite may 401 402 have reduced porosity, the data presented here also shows that advanced carbonate

dissolution occurred as a result of pyrite precipitation. Hence, it is possible that the
overall effects of TSR-related pyrite precipitation and carbonate dissolution on these
dolostone reservoirs are positive in terms of reservoir quality.

#### 406 TSR impact on reservoir fluid composition and pressure

According to balanced TSR equations (1-4), fluid phase H<sub>2</sub>S, CO<sub>2</sub>, elemental sulphur 407 and H<sub>2</sub>O are all produced during TSR. Hence, both pore-fluid pressure and fluid 408 composition were significantly altered by TSR. Trace elements, rare earth elements, 409 410 and strontium isotopic data from TSR calcite demonstrate that TSR diagenesis most likely represents a relatively closed system in the Feixianguan Formation (Jiang et al., 411 412 2015a). Previously published oxygen isotope data from TSR calcite in the 413 Feixianguan Formation suggest that TSR-calcite precipitated in isotopic and thermal equilibrium with the host rock, TSR water and sulphate minerals, and the negative 414 shift of carbon isotopes in this type of calcite is indicative of carbon partially sourced 415 from hydrocarbons due to TSR and partial adoption of the matrix dolomite  $\delta^{13}$ C 416 417 signal (Cai et al., 2014; Huang et al., 2012; Jiang et al., 2015c) similar to the explanation of mixed  $\delta^{13}$ C values in TSR calcite for other formations (Worden and 418 Smalley, 1996). 419

Fluid inclusion data show that about four times the volume of fresh water, compared to the initial residual formation water, was generated by TSR and added to the fluids in the Feixianguan Formation during TSR (Fig. 6) (Jiang et al., 2015c). Similar findings have also been reported for the Permian Khuff Formation from Abu Dhabi (Worden et al., 1996), and in the Devonian fields from the Western Canada Sedimentary Basin (Yang et al., 2001). Generation of low salinity water by TSR possibly resulted in the formation water being transiently undersaturated with respect 427 to calcite and dolomite, which would also facilitating the dissolution of carbonate428 minerals in the Feixianguan Formation.

429 The contact relationships of the ooids demonstrate that there is lack of compaction in these dolostone reservoirs, precluding the possibility that disequilibrium compaction 430 431 contributed to the overpressure during progressive burial (Heydari, 2000). Therefore, 432 the elevated fluid pressure (near lithostatic pressure) at maximum burial and the continuation of these high pressures during early uplift stage were probably related to 433 434 the addition of gas either by oil cracking or TSR. It is possible that systematic 435 increase in fluid overpressure during burial (Fig. 7B) was related to the generation of hydrocarbon gases due to secondary cracking of the primary oil that was originally 436 derived from Upper Permian marine source rocks (Cai et al., 2010; Hao et al., 2008). 437 Thermochemical sulphate reduction seems to have occurred simultaneously with oil 438 cracking and had a significant impact on both gas composition and isotopes (Cai et al., 439 440 2013; Hao et al., 2008). An increase of fluid pressure due to the production of  $H_2S$ and  $CO_2$  in these dolostone reservoirs (equation 4) has been proven to be a result of 441 TSR (Liu et al., 2006). However, fluid pressure progressively decreased to near-442 443 hydrostatic pressure (~56 MPa) during further uplift (Fig. 7A)(Liu et al., 2009). Fluid pressure increasing during burial followed by a decreasing trend during uplifting 444 demonstrates that oil cracking may have been completed from the maximum burial to 445 early uplift stage, and continuation of exhumation and the generation of fractures may 446 447 have released some fluid pressure in these dolostone reservoirs.

We propose a model in which oil cracking due to progressive burial and increasing
temperature, associated with TSR, led to increasing pore-fluid pressure in the
Feixianguan Formation. During uplift, after oil cracking and bitumen formation, TSR

451 continued to occur and produced H<sub>2</sub>S and CO<sub>2</sub>, maintaining some of the fluid pressure 452 in the reservoir. In addition, pressure may have been episodically released from the system during uplift, resulting in methane exsolving from pore water into free-gas 453 454 phase, maintaining methane saturation in the aqueous phase (Becker et al., 2010). Overpressure during TSR diagenesis may have resulted in the forcing of pore-fluids 455 out of the TSR site. It is possible that a complex cycling of fluids on a reservoir scale 456 457 is involved during TSR, resulting in some zones of reservoir develop higher porosity whereas others are occupied by calcite cements. 458

### 459 Implications for deeply buried carbonate reservoir exploration

In carbonate reservoirs, open system diagenesis, that can lead to enhanced porosity, 460 461 has been proposed by numerous authors. In some carbonate systems, secondary porosity has been interpreted to have formed by near-surface karstification (Loucks, 462 1999), due to early dissolution by meteoric water and/or dolomitization during water 463 leaching (Dickson and Kenter, 2014; Jiang et al., 2016; Lucia et al., 1994; Zhu et al., 464 2006), by mesogenetic dissolution (Kenter et al., 2006; Mazzullo and Harris, 1992), 465 466 as well as by hydrothermal karstification and dissolution (Biehl et al., 2016a; Davies and Smith, 2006; Jiang et al., 2015b; Packard et al., 2001; Saller and Dickson, 2011; 467 Smith, 2006). In contrast, in a closed or semi-closed burial diagenetic system, where 468 469 water/rock ratios is low, the formation waters are commonly interpreted to be saturated with respect to calcite (Bjørlykke and Jahren, 2012; Ehrenberg et al., 2012). 470

471 Based on detailed petrographic and geochemical data, as well as permeability,
472 pressure and geochemical modelling, we here conclude that porosity and permeability
473 has been increased by TSR in the Feixianguan Formation even though this rock unit is
474 probably part of a relatively closed system. The current CO<sub>2</sub> concentration measured

475 in the dolostone reservoir appears to correlate with H<sub>2</sub>S concentrations, and carbon isotopic composition of the contemporary CO<sub>2</sub> gas is <sup>13</sup>C-enriched and does not 476 reflect a <sup>12</sup>C-rich CO<sub>2</sub> that would be expected from oxidation of hydrocarbon by 477 sulphate (Cai et al., 2014; Hao et al., 2015; Huang et al., 2012). This observation is 478 consistent with significant dissolution occurring and <sup>13</sup>C-rich CO<sub>2</sub> added to these 479 reservoirs. Carbonate dissolution during deep burial environments may be related to 480 TSR by generation of water, supported by this study, TSR modelling (Fu et al., 2016), 481 and generation of acidity during the reaction of  $H_2S$  with siderite component in 482 483 dolomite. In addition, elemental sulphur generated by TSR may have maintained porosity from calcite precipitation. It is likely that TSR is capable of increasing 484 485 reservoir heterogeneity and maintaining porosity by the inhibition of compactional 486 processes due to the high fluid pressure conditions. Our study shows that TSR has enhanced the reservoir porosity by 1.6 % (Table 2) and doubled the permeability in 487 the Feixianguan Formation dolostone (Table 3), which has clear and significant 488 489 implications for petroleum exploration in deep sedimentary basins that experienced TSR (Biehl et al., 2016a; Biehl et al., 2016b; Bjørlykke and Jahren, 2012; Cai et al., 490 2014; Ehrenberg et al., 2012; Heydari, 1997; Jiang et al., 2015c; Machel and 491 Buschkuehle, 2008; Mazzullo and Harris, 1991; Worden et al., 1996; Yang et al., 492 2001). 493

## 494 Conclusions

1. Deeply buried Feixianguan Formation dolostone reservoirs from the Sichuan Basin
(mainly between 3000 and 6000 m) contain dissolution enlarged pore spaces. These
dissolution pores have close genetic links to thermochemical sulphate reduction
(TSR), suggesting that TSR was responsible for the enhancement of reservoir quality.

This is in agreement with geochemical model results that demonstrate an overallporosity increase of 1.6% and doubled permeability.

501 2. Elemental sulphur occurred as a liquid (or was dissolved in gas) present at the
502 contact between petroleum and water or directly in contact with the host dolomites in
503 the dissolution pores, inhibiting the new secondary pores from undergoing calcite
504 precipitation.

505 3. Creation of fresh water under deep burial environments may cause dissolution of
506 carbonate minerals because the formation water may become transiently
507 undersaturated with respect to calcite and dolomite.

4. Overpressure caused complex cycling of fluids within the reservoir during TSR,
resulting in some reservoir zones being occupied by calcite cements and others
developing higher porosity, thus increasing reservoir heterogeneity.

5. Further dissolution of carbonate probably occurred because of the release of H<sup>+</sup> due to pyrite precipitation, with Fe sourced from a ferroan carbonate component in dolomite. This is supported by the positive  $\delta^{13}$ C values of present-day CO<sub>2</sub> in these reservoirs.

6. This is the first documented case of how TSR can improve carbonate reservoir
quality under a relatively closed diagenetic system. This phenomenon seems not to
have been fully appreciated in other sedimentary basins that experienced TSR.

#### 518 ACKNOWLEDGEMENTS:

This work has been financially supported by the Natural Science Foundation of China
(Grant No. 41402132), the National Science and Technology Majoy Project (Grant
No. 2017ZX05008-004), and scholarships under the China Postdoctoral Science
Foundation award for International Postdoctoral Exchange Fellowship Program

523 (Grant No. 20150035) and the Chinese Scholarship Council (CSC) (Grant No. 201704910007).

525

#### 526 **References**

- Abou-Kassem, J.H. (2000) Experimental and numerical modeling of sulfur plugging in
   carbonate reservoirs. Journal of Petroleum Science and Engineering 26, 91-103.
- Becker, S., Eichhubl, P., Laubach, S., Reed, R., Lander, R. and Bodnar, R. (2010) A 48 my
  history of fracture opening, temperature, and fluid pressure: Cretaceous Travis Peak
  Formation, East Texas basin. Geological Society of America Bulletin 122, 1081-1093.
- Biehl, B.C., Reuning, L., Schoenherr, J., Lüders, V. and Kukla, P.A. (2016a) Impacts of
  hydrothermal dolomitization and thermochemical sulfate reduction on secondary
  porosity creation in deeply buried carbonates: A case study from the Lower Saxony
  Basin, northwest Germany. AAPG Bulletin 100, 597-621.
- Biehl, B.C., Reuning, L., Schoenherr, J., Lewin, A., Leupold, M. and Kukla, P.A. (2016b) Do
   CO<sub>2</sub>-charged fluids contribute to secondary porosity creation in deeply buried
   carbonates? Marine and Petroleum Geology 76, 176-186.
- Bildstein, O., Worden, R.H. and Brosse, E. (2001) Assessment of anhydrite dissolution as the
  rate-limiting step during thermochemical sulfate reduction. Chemical Geology 176,
  173-189.
- 542 Bjørlykke, K. and Jahren, J. (2012) Open or closed geochemical systems during diagenesis in
  543 sedimentary basins: Constraints on mass transfer during diagenesis and the prediction
  544 of porosity in sandstone and carbonate reservoirs. American Association of Petroleum
  545 Geologists, Bulletin 96, 2193-2214.
- Bodnar, R.J. (2003) Reequilibration of fluid inclusions. Fluid inclusions: Analysis and
   interpretation 32, 213-230.
- Cai, C.F., He, W.X., Jiang, L., Li, K.K., Xiang, L. and Jia, L.Q. (2014) Petrological and
  geochemical constraints on porosity difference between Lower Triassic sour-and
  sweet-gas carbonate reservoirs in the Sichuan Basin. Marine and Petroleum Geology 56,
  34-50.
- Cai, C.F., Li, K.K., Zhu, Y.M., Xiang, L. and Jiang, L. (2010) TSR origin of sulfur in
  Permian and Triassic reservoir bitumen, East Sichuan Basin, China. Organic
  Geochemistry.
- Cai, C.F., Worden, R.H., Bottrell, S.H., Wang, L.S. and Yang, C.C. (2003) Thermochemical
  sulphate reduction and the generation of hydrogen sulphide and thiols (mercaptans) in
  Triassic carbonate reservoirs from the Sichuan Basin, China. Chemical Geology 202,
  39-57.
- Cai, C.F., Zhang, C.M., He, H. and Tang, Y.J. (2013) Carbon isotope fractionation during
  methane-dominated TSR in East Sichuan Basin gasfields, China: A review. Marine and
  Petroleum Geology 48, 100-110.
- Davies, G.R. and Smith, L.B. (2006) Structurally controlled hydrothermal dolomite reservoir
   facies: An overview. American Association of Petroleum Geologists, Bulletin 90, 1641.
- 564 Dickson, J.A.D. and Kenter, J.A.M. (2014) Diagenetic Evolution of Selected Parasequences
   565 Across A Carbonate Platform: Late Paleozoic, Tengiz Reservoir, Kazakhstan. Journal
   566 of Sedimentary Research 84, 664-693.
- 567 Duan, Z. and Mao, S. (2006) A thermodynamic model for calculating methane solubility,
  568 density and gas phase composition of methane-bearing aqueous fluids from 273 to
  569 523K and from 1 to 2000bar. Geochimica et Cosmochimica Acta 70, 3369-3386.
- Duan, Z.H. (1992) An equation of state for the C&-CO\*-Hz0 system: II. Mixtures from 50 to
  1000 C and 0 to 1000 bar.

- 572 Ehrenberg, S.N., Walderhaug, O. and Bjørlykke, K. (2012) Carbonate porosity creation by
  573 mesogenetic dissolution: Reality or illusion? American Association of Petroleum
  574 Geologists, Bulletin 96, 217-233.
- Fu, Y.J., van Berk, W. and Schulz, H.-M. (2016) Hydrogen sulfide formation, fate, and
  behavior in anhydrite-sealed carbonate gas reservoirs: A three-dimensional reactive
  mass transport modeling approach. American Association of Petroleum Geologists,
  Bulletin 100, 843-865.
- Goldstein, R.H. (2012) Fluid inclusion geothermometry in sedimentary systems: from
   paleoclimate to hydrothermal. SEPM special publication, Thermal History Analysis of
   Sedimentary Basins, 31 103, 45-63.
- Goldstein, R.H. and Reynolds, T.J. (1994) Systematics of fluid inclusions in diagenetic
   minerals: SEPM Short Course Notes, 31. 199.
- Hao, F., Guo, T.L., Zhu, Y.M., Cai, X.Y., Zou, H.Y. and Li, P.P. (2008) Evidence for
  multiple stages of oil cracking and thermochemical sulfate reduction in the Puguang
  gas field, Sichuan Basin, China. American Association of Petroleum Geologists,
  Bulletin 92, 611.
- Hao, F., Zhang, X.F., Wang, C.W., Li, P.P., Guo, T.L., Zou, H.Y., Zhu, Y.M., Liu, J.Z. and
  Cai, Z.X. (2015) The fate of CO2 derived from thermochemical sulfate reduction (TSR)
  and effect of TSR on carbonate porosity and permeability, Sichuan Basin, China.
  Earth-Science Reviews 141, 154-177.
- Heydari, E. (1997) The role of burial diagenesis in hydrocarbon destruction and H2S
  accumulation, Upper Jurassic Smackover Formation, Black Creek Field, Mississippi.
  American Association of Petroleum Geologists, Bulletin 81, 26-45.
- Heydari, E. (2000) Porosity loss, fluid flow, and mass transfer in limestone reservoirs:
  Application to the upper Jurassic Smackover formation, Mississippi. American
  Association of Petroleum Geologists, Bulletin 84, 100-118.
- Huang, S.J., Huang, K.K., Lü, J. and Lan, Y.F. (2012) Carbon isotopic composition of Early
  Triassic marine carbonates, Eastern Sichuan Basin, China. Science China Earth
  Sciences 55, 2026-2038.
- Hutcheon, I. and Krouse, H.R. (1994) Thermochemical sulfate reduction, evidence from the
  field, theory and experiment. Abstracts of Papers of the American Chemical Society
  208, 105-GEOC.
- Hutcheon, I., Krouse, H.R. and Abercrombie, H.J. (1995) Controls on the origin and
  distribution of elemental sulfur, H<sub>2</sub>S, and CO<sub>2</sub> in paleozoic hydrocarbon reservoirs in
  Western Canada, in: Vairavamurthy, M.A., Schoonen, M.A.A. (Eds.), Geochemical
  Transformations of Sedimentary Sulfur, pp. 426-438.
- Jiang, L., Cai, C., Worden, R., Li, K.K. and Xiang, L. (2013) Reflux dolomitization of the
  Upper Permian Changxing Formation and the Lower Triassic Feixianguan Formation,
  NE Sichuan Basin, China. Geofluids.
- Jiang, L., Cai, C.F., Worden, R.H., Crowley, S.F., Jia, L.Q., Zhang, K. and Duncan, I.J. (2016)
  Multiphase dolomitization of deeply buried Cambrian petroleum reservoirs, Tarim
  Basin, north-west China. Sedimentology 63, 2130-2157.
- Jiang, L., Cai, C.F., Worden, R.H., Li, K.K., Xiang, L., Chu, X.L., Shen, A.J. and Li, W.J.
  (2015a) Rare earth element and yttrium (REY) geochemistry in carbonate reservoirs during deep burial diagenesis: Implications for REY mobility during thermochemical sulfate reduction. Chemical Geology 415, 87-101.
- Jiang, L., Pan, W.Q., Cai, C.F., Jia, L.Q., Pan, L.Y., Wang, T.K., Li, H.X., Chen, S.L. and
  Chen, Y. (2015b) Fluid mixing induced by hydrothermal activity in the Ordovician
  carbonates in Tarim Basin, China. Geofluids.
- Jiang, L., Worden, R.H. and Cai, C.F. (2014a) Thermochemical sulfate reduction and fluid
  evolution of the Lower Triassic Feixianguan Formation sour gas reservoirs, northeast
  Sichuan Basin, China. American Association of Petroleum Geologists, Bulletin 98,
  947-973.
- Jiang, L., Worden, R.H. and Cai, C.F. (2015c) Generation of isotopically and compositionally
   distinct water during thermochemical sulfate reduction (TSR) in carbonate reservoirs:

- Triassic Feixianguan Formation, Sichuan Basin, China. Geochimica et Cosmochimica
  Acta 165, 249-262.
- Jiang, L., Worden, R.H., Cai, C.F., Li, K.K., Xiang, L., Cai, L.L. and He, X.Y. (2014b)
  Dolomitization of Gas Reservoirs: The Upper Permian Changxing and Lower Triassic
  Feixianguan Formations, Northeast Sichuan Basin, China. Journal of Sedimentary
  Research 84, 792-815.
- Kenter, J.A.M., Harris, P.M., Collins, J.F., Weber, L.J., Kuanysheva, G. and Fischer, D.J.
  (2006) Late Visean to Bashkirian platform cyclicity in the central Tengiz buildup,
  Precaspian Basin, Kazakhstan: Depositional evolution and reservoir development, in P.
  M. Harris and L. J. Weber, eds., Giant hydrocarbon reservoirs of the world: From rocks
  to reservoir characterization and modeling: AAPG Memoir 88, SEPM Special
  Publication, 7-54.
- Krouse, H.R., Viau, C.A., Eliuk, L.S., Ueda, A. and Halas, S. (1988) Chemical and isotopic
  evidence of thermochemical sulfate reduction by light hydrocarbon gases in deep
  carbonate reservoirs. Nature 333, 415-419.
- Kuo, C.H. (1972) On the production of hydrogen sulfide-sulfur mixtures from deep formations. Journal of Petroleum Technology 24, 1,142-141,146.
- Li, J., Xie, Z., Dai, J., Zhang, S., Zhu, G. and Liu, Z. (2005) Geochemistry and origin of sour
  gas accumulations in the northeastern Sichuan Basin, SW China. Organic
  Geochemistry 36, 1703-1716.
- Lin, F., Bodnar, R. and Becker, S. (2007) Experimental determination of the Raman CH4
  symmetric stretching (v1) band position from 1–650bar and 0.3–22° C: Application to
  fluid inclusion studies. Geochimica et Cosmochimica Acta 71, 3746-3756.
- Liu, D.H., Dai, J.X., Xiao, X.M., Tian, H., Yang, C., Hu, A.P., Mi, J. and Song, Z.G. (2009)
  High density methane inclusions in Puguang Gasfield: Discovery and a TP genetic
  study. Chinese Science Bulletin 54, 4714-4723.
- Liu, D.H., Xiao, X.M., Xiong, Y.Q., Geng, A.S., Tian, H., Peng, P., Shen, J.G. and Wang,
  Y.P. (2006) Origin of natural sulphur-bearing immiscible inclusions and H<sub>2</sub>S in oolite
  gas reservoir, Eastern Sichuan. Science in China Series D-Earth Sciences 49, 242-257.
- Liu, Q.Y., Worden, R.H., Jin, Z.J., Liu, W.H., Li, J., Gao, B., Zhang, D.W., Hu, A.P. and
  Yang, C. (2013) TSR versus non-TSR processes and their impact on gas geochemistry
  and carbon stable isotopes in Carboniferous, Permian and Lower Triassic marine
  carbonate gas reservoirs in the Eastern Sichuan Basin, China. Geochimica et
  Cosmochimica Acta 100, 96-115.
- Liu, Q.Y., Worden, R.H., Jin, Z.J., Liu, W.H., Li, J., Gao, B., Zhang, D.W., Hu, A.P. and
  Yang, C. (2014) Thermochemical sulphate reduction (TSR) versus maturation and their
  effects on hydrogen stable isotopes of very dry alkane gases. Geochimica et
  Cosmochimica Acta 137, 208-220.
- Liu, Y.F., Qiu, N.S., Xie, Z.Y., Yao, Q.Y. and Zhu, C.Q. (2016) Overpressure compartments
  in the central paleo-uplift, Sichuan Basin, southwest China. AAPG Bulletin 100, 867888.
- Loucks, R.G. (1999) Paleocave carbonate reservoirs; origins, burial-depth modifications,
  spatial complexity, and reservoir implications. American Association of Petroleum
  Geologists, Bulletin 83, 1795-1834.
- Lucia, F.J., Major, R.P., Purser, B., Tucker, M. and Zenger, D. (1994) Porosity evolution
  through hypersaline reflux dolomitization. Dolomites: A Volume in Honour of
  Dolomieu 21.
- Ma, Y.S., Guo, T.L., Zhao, X.F. and Cai, X.Y. (2008a) The formation mechanism of highquality dolomite reservoir in the deep of Puguang Gas Field. Science in China Series
  D-Earth Sciences 51, 53-64.
- Ma, Y.S., Zhang, S., Guo, T.L., Zhu, G., Cai, X.Y. and Li, M. (2008b) Petroleum geology of
  the Puguang sour gas field in the Sichuan Basin, SW China. Marine and Petroleum
  Geology 25, 357-370.
- Machel, H.G. (1987) Saddle dolomite as a by-product of chemical compaction and
   thermochemical sulfate reduction. Geology 15, 936-940.

- Machel, H.G. (2001) Bacterial and thermochemical sulfate reduction in diagenetic settings—
   old and new insights. Sedimentary Geology 140, 143-175.
- Machel, H.G. and Buschkuehle, B.E. (2008) Diagenesis of the Devonian Southesk-Cairn
  Carbonate Complex, Alberta, Canada: Marine Cementation, Burial Dolomitization,
  Thermochemical Sulfate Reduction, Anhydritization, and Squeegee Fluid Flow.
  Journal of Sedimentary Research 78, 366.
- Machel, H.G., Krouse, H.R., Riciputi, L.R. and Cole, D.R. (1995) Devonian Nisku sour gas
  play, Canada: A unique natural laboratory for study of thermochemical sulfate
  reduction, in: Vairavamurthy, M.A., Schoonen, M.A.A. (Eds.), Geochemical
  Transformations of Sedimentary Sulfur, pp. 439-454.
- Mazzullo, S.J. and Harris, P.M. (1991) An overview of dissolution porosity development in
  the deep-burial environment, with examples from carbonate reservoirs in the Permian
  Basin. Permian Basin Plays CTomorrow; s Technology Today: West Texas
  Geological Society, Publ, 125-138.
- Mazzullo, S.J. and Harris, P.M. (1992) Mesogenetic dissolution: its role in porosity
   development in carbonate reservoirs (1). AAPG bulletin 76, 607-620.
- Meyer, B. (1976) Elemental sulfur. Chemical Reviews 76, 367-388.
- 699 Oakes, C.S., Bodnar, R.J. and Simonson, J.M. (1990) The system NaCl CaCl 2 H 2 O: I.
  700 The ice liquidus at 1 atm total pressure. Geochimica et Cosmochimica Acta 54, 603701 610.
- Packard, J.J., Al-Aasm, I., Samson, I., Berger, Z. and Davies, J. (2001) A Devonian
  hydrothermal chert reservoir: the 225 bcf Parkland field, British Columbia, Canada.
  American Association of Petroleum Geologists, Bulletin 85, 51-84.
- Parkhurst, D.L. and Appelo, C.A.J. (2013) Description of input and examples for PHREEQC
  version 3—A computer program for speciation, batch-reaction, one-dimensional
  transport, and inverse geochemical calculations. US Geological Survey Techniques and
  Methods 6, 497.
- Qiu, N.S., Qin, J.Z., McInnes, B., Wang, J. and Zhen, L.J. (2008) Tectonothermal evolution
  of the northeastern Sichuan Basin: Constraints from apatite and zircon (U-Th)/He ages
  and vitrinite reflectance data. Geol J Chin Univ 14, 223-230.
- Saller, A.H. and Dickson, J.A.T.D. (2011) Partial dolomitization of a Pennsylvanian
  limestone buildup by hydrothermal fluids and its effect on reservoir quality and
  performance. American Association of Petroleum Geologists, Bulletin 95, 1745-1762.
- Smith, L.B. (2006) Origin and reservoir characteristics of Upper Ordovician Trenton–Black
   River hydrothermal dolomite reservoirs in New York. American Association of
   Petroleum Geologists, Bulletin 90, 1691-1718.
- Worden, R.H. and Smalley, P.C. (1996) H<sub>2</sub>S-producing reactions in deep carbonate gas reservoirs: Khuff Formation, Abu Dhabi. Chemical geology 133, 157-171.
- Worden, R.H., Smalley, P.C. and Cross, M.M. (2000) The influence of rock fabric and mineralogy on thermochemical sulfate reduction: Khuff Formation, Abu Dhabi. Journal of Sedimentary Research 70.
- Worden, R.H., Smalley, P.C. and Oxtoby, N.H. (1995) Gas souring by thermochemical
  sulfate reduction at 140 degrees C. American Association of Petroleum Geologists
  Bulletin 79, 854-863.
- Worden, R.H., Smalley, P.C. and Oxtoby, N.H. (1996) The effects of thermochemical sulfate
   reduction upon formation water salinity and oxygen isotopes in carbonate gas
   reservoirs. Geochimica et Cosmochimica Acta 60, 3925-3931.
- Xu, T.F., Kharaka, Y.K., Doughty, C., Freifeld, B.M. and Daley, T.M. (2010) Reactive
   transport modeling to study changes in water chemistry induced by CO<sub>2</sub> injection at the
   Frio-I Brine Pilot. Chemical Geology 271, 153-164.
- Yang, C., Hutcheon, I. and Krouse, H.R. (2001) Fluid inclusion and stable isotopic studies of
  thermochemical sulphate reduction from Burnt Timber and Crossfield East gas fields in
  Alberta, Canada. Bulletin of Canadian Petroleum Geology 49, 149-164.

735	Yang, C., Sa	imper.	, J. and Mon	tenegro, L. (20	008) A coup	oled	non	-isother	mal reactive	trai	isport
736	model	for	long-term	geochemical	evolution	of	а	HLW	repository	in	clay.
737	Enviro	onmen	ntal Geology	53, 1627-1638	8.						

- Zhao, W., Luo, P., Chen, G., Cao, H. and Zhang, B. (2005) Origin and reservoir rock
  characteristics of dolostones in the Early Triassic Feixianguan Formation, NE Sichuan
  Basin, China: Significance for future gas exploration. Journal of Petroleum Geology 28,
  83-100.
- Zhu, G.Y., Zhang, S.C., Liang, Y.B., Dai, J.X. and Li, J. (2005) Isotopic evidence of TSR
  origin for natural gas bearing high H<sub>2</sub>S contents within the Feixianguan Formation of
  the northeastern Sichuan Basin, southwestern China. Science in China Series D-Earth
  Sciences 48, 1960-1971.
- Zhu, G.Y., Zhang, S.C., Liang, Y.B., Ma, Y.S., Guo, T.L. and Zhou, G.Y. (2006) Distribution
  of high H<sub>2</sub>S-bearing natural gas and evidence of TSR origin in the Sichuan basin. Acta
  Geologica Sinica 80, 1208-1218(In Chinese).
- 749

#### 750 **Table Caption**

- 751
- 752 Table 1. Point counting data showing percentages of each component in TSR-
- 753 prevailed dolo-grainstone reservoirs in the Feixiangian Formation
- 754
- Table 2. Mineral volume and porosity change by TSR modeling, porosity before TSR.
- Table 3. Permeability change by TSR modeling, the equation of relationship between porosity and permeability ( $y = 0.0166e^{0.5498x}$ ,  $R^2 = 0.6489$ ) was obtained from core plug measurement from the Luojia-2 well.
- 760
- 761 Figure Caption
- 762

Figure 1. (A) Location and main structural elements of the Sichuan Basin, modified
from Hao et al. (2008). (B) Paleogeography and locations of the sampled gas fields in
the Feixianguan Formation dolostone reservoirs.

766

Figure 2. Representative wells showing stratigraphic and porosity correlation of the ooid-enriched limestone (that has not experienced TSR) and dolostone (that experienced TSR) in the Feixianguan Formation. Dolostone that experienced TSR appears to have the better reservoir quality; modified from (Ma et al., 2008a).

771

Figure 3. Photomicrographs show TSR-related calcite and elemental sulphur indissolution pores in the Feixianguan Formation. A) EDS image of calcite (white

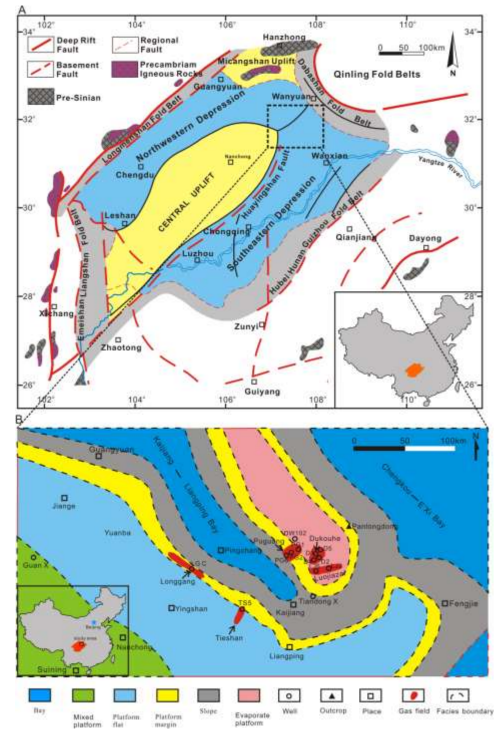
774 colour in the pore space) with a small volume of elemental sulphur (green colour in 775 the pore space) filling in dolomite host rock (red colour), from well D2, depth 4308 m. B) EDS image of elemental sulphur (green colour in the pore space) present around 776 the dissolution enlarged pore spaces in dolomite host rock (red colour), from well LJ2, 777 778 depth 3256.5 m; C) EDS image showing big elemental sulphur aggressive (green colour) and calcite cement (write colour) in dissolution enlarged pore, from well D2, 779 depth 4308 m; D) BSEM image of elemental sulphur (red arrow) occurs as shield of 780 dissolution enlarged pore, from well Du 4, depth 4793 m. 781

782 Figure 4. BESM images showing dissolution enlarged pore spaces and TSR-related calcite and pyrite in the Feixianguan Formation. A) Dissolution enlarged pore with 783 784 size up to few millimetres locally filling with small volume of pyrite, well LJ 2, depth 3256.5 m. B) Dolo-grainstone enriched in dissolution enlarged pores with few calcite 785 786 cements, well LJ 2, depth 3256.5 m. C) Dolostone reservoir containing open dissolution-enlarged pore spaces and a lack of cementation, well LJ 3-58. D) Intensive 787 788 calcite cementation is observed in the left side of the image, whereas abundant dissolution enlarged pores present in the right side, from well Du 4, depth 4793 m. 789

Figure 5: BSEM images showing the protection of pore spaces by elemental sulphur 790 and the dissolution of late stage dolomite in the Feixianguan Formation. A) Elemental 791 sulphur (red arrow) occurs as a shield around the dolomite cements and host rock in 792 an open pore, from well LJ 2, depth 3232.2 m. B) Elemental sulphur (red arrow) grow 793 794 on the edge of an open pore, from well Po 2. C) Calcite precipitated in pore spaces 795 that are lacking in elemental sulphur, pore spaces appear to be well-developed where elemental sulphur is present, from well Du 4, depth 4793 m. D) Late stage, pore-796 filling dolomite cement showing evidence of dissolution demonstrates that 797 mesogenetic dissolution occurred during deep burial diagenetic environments, from 798 799 well LJ 3-58.

Figure 6: Comparison of salinity and temperature from fluids inclusions in of oil-stage and gas-stage TSR calcite. Salinity of the formation water decreased from approximately 25 wt. % to less than 10 wt. % during oil-stage TSR, and formation salinity continue to decrease from about 10 wt. % down to approximately 5 wt. % during gas-stage TSR. Figure 7: (A) Trapping pressures of fluid inclusions in TSR calcite. (B) Trapping
temperature plotted against calculated trapping fluid pressures of fluids inclusions in
TSR calcite with the data split into samples that had oil-stage TSR (oil inclusions
present) and ones that gas-stage TSR (no oil inclusions present).

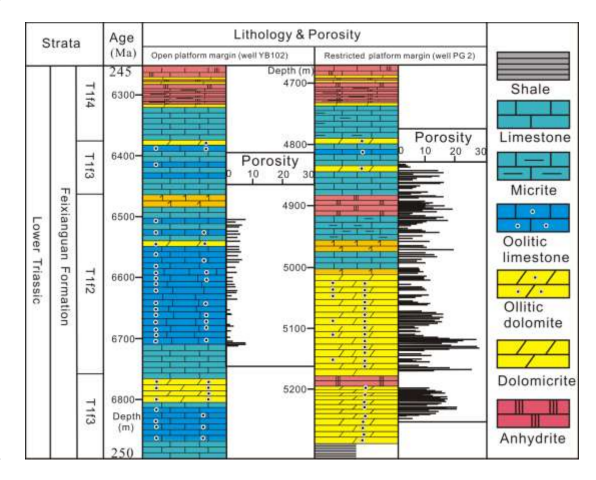
- 810 Figure 1. (A) Location and main structural elements of the Sichuan Basin, modified
- 811 from Hao et al. (2008). (B) Paleogeography and locations of the sampled gas fields in
- 812 the Feixianguan Formation dolostone reservoirs.



814

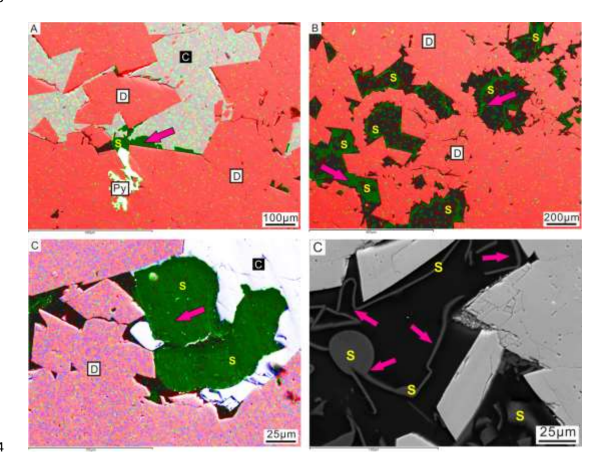
813

Figure 2. Representative wells showing stratigraphic and porosity correlation of the
ooid-enriched limestone (that has not experienced TSR) and dolostone (that
experienced TSR) in the Feixianguan Formation. Dolostone that experienced TSR
appears to have the better reservoir quality; modified from (Ma et al., 2008a).



823 Figure 3. Photomicrographs show TSR-related calcite and elemental sulphur in dissolution pores in the Feixianguan Formation. A) EDS image of calcite (white 824 825 colour in the pore space) with a small volume of elemental sulphur (green colour in 826 the pore space) filling in dolomite host rock (red colour), from well D2, depth 4308 m. 827 B) EDS image of elemental sulphur (green colour in the pore space) present around the dissolution enlarged pore spaces in dolomite host rock (red colour), from well LJ2, 828 829 depth 3256.5 m; C) EDS image showing big elemental sulphur aggressive (green colour) and calcite cement (write colour) in dissolution enlarged pore, from well D2, 830 831 depth 4308 m; D) BSEM image of elemental sulphur (red arrow) occurs as shield of dissolution enlarged pore, from well Du 4, depth 4793 m. 832

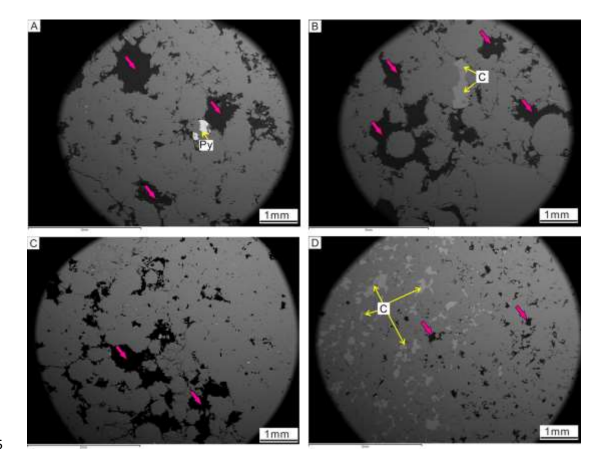
833



834

836 Figure 4. BESM images showing dissolution enlarged pore spaces and TSR-related calcite and pyrite in the Feixianguan Formation. A) Dissolution enlarged pore with 837 838 size up to few millimetres locally filling with small volume of pyrite, well LJ 2, depth 3256.5 m. B) Dolo-grainstone enriched in dissolution enlarged pores with few calcite 839 cements, well LJ 2, depth 3256.5 m. C) Dolostone reservoir containing open 840 841 dissolution-enlarged pore spaces and a lack of cementation, well LJ 3-58. D) Intensive 842 calcite cementation is observed in the left side of the image, whereas abundant dissolution enlarged pores present in the right side, from well Du 4, depth 4793 m. 843

844



845

847 Figure 5: BSEM images showing the protection of pore spaces by elemental sulphur and the dissolution of late stage dolomite in the Feixianguan Formation. A) Elemental 848 849 sulphur (red arrow) occurs as a shield around the dolomite cements and host rock in 850 an open pore, from well LJ 2, depth 3232.2 m. B) Elemental sulphur (red arrow) grow 851 on the edge of an open pore, from well Po 2. C) Calcite precipitated in pore spaces 852 that are lacking in elemental sulphur, pore spaces appear to be well-developed where 853 elemental sulphur is present, from well Du 4, depth 4793 m. D) Late stage, porefilling dolomite cement showing evidence of dissolution demonstrates that 854 mesogenetic dissolution occurred during deep burial diagenetic environments, from 855 well LJ 3-58. 856

857

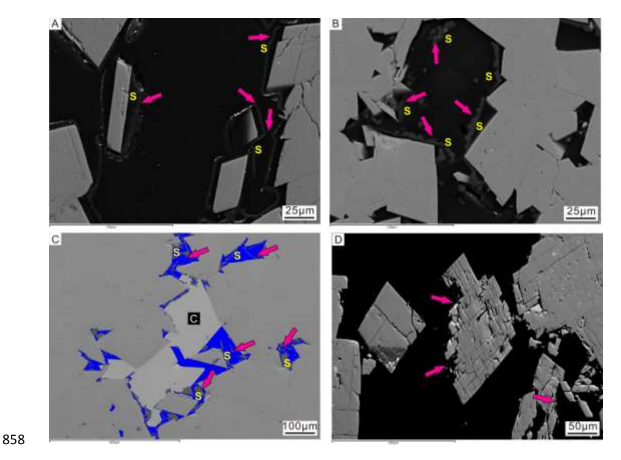


Figure 6: Comparison of salinity and temperature from fluids inclusions in of oil-stage and gas-stage TSR calcite. Salinity of the formation water decreased from approximately 25 wt. % to less than 10 wt. % during oil-stage TSR, and formation salinity continue to decrease from about 10 wt. % down to approximately 5 wt. % during gas-stage TSR.

865

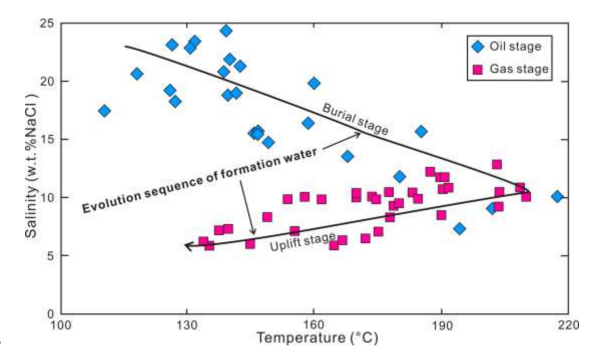
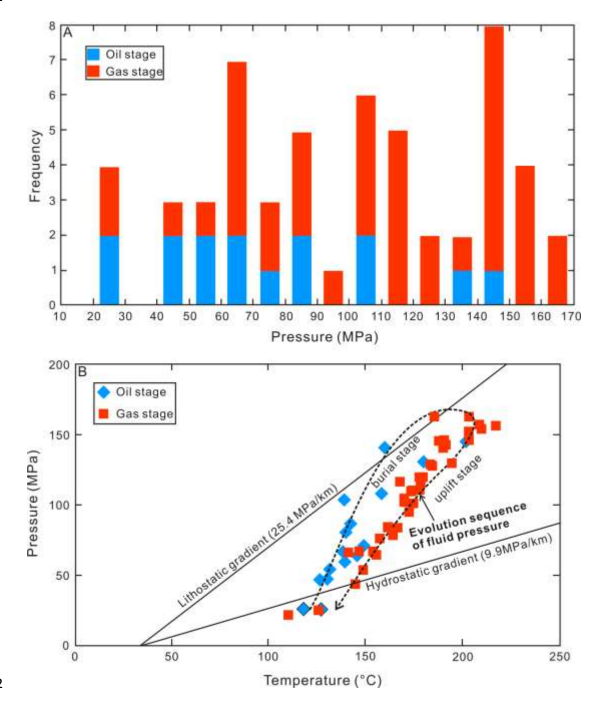


Figure 7: (A) Trapping pressures of fluid inclusions in TSR calcite. (B) Trapping
temperature plotted against calculated trapping fluid pressures of fluids inclusions in
TSR calcite with the data split into samples that had oil-stage TSR (oil inclusions
present) and ones that gas-stage TSR (no oil inclusions present).



874	Table 1.

	Depth	Grai	Matri	Ca	rbona	te cem	ents (	(%)	Porosit	Bitume	Pyrit	Quart
No.	(m)	n (%)	x (%)	C1	C2	C3	D	D	y (%)	n (%)	e (%)	z (%)
LJ2-1	3198	67	6	0	0	2	1 7	$\frac{2}{2}$	6	9	0	1
LJ2-18	3267	70	0	0	0	0	15	8	5	2	0	0
LJ2-25	3256	65	0	0	0	0	15	5	8	5	2	0
LJ2-33	3232	75	0	0	0	0	0	15	8	2	0	0
LJ2-37	3233	56	0	0	4	0	17	0	20	3	0	0
LJ2-26	3256	71	0	0	0	0	0	17	8	1	3	0
LJ3-58		69	0	0	2	0	10	5	13	0	1	0
LJ3-27		66	0	0	0	0	18	7	3	6	0	0
LJ2		71	0	0	2	0	8	15	2	1	1	0
LJ2-23		67	0	0	0	0	25	0	3	5	0	0
D1		58	0	0	3	0	17	4	9	8	0	1
D2-25	4309	62	0	0	5	0	0	20	10	1	2	0
D2-7		57	0	0	0	0	0	18	16	2	2	5
D4-4	4236	78	0	0	4	0	5	0	13	0	0	0
D5	4793	55	0	0	3	0	11	6	12	9	0	4
DW10	4901	71	9	17	3	0	0	0	0	0	0	0
2 PLD-3	outcro	80	0	0	0	6	0	0	0	12	2	0
PLD-4	p outcro p	72	0	0	0	14	0	0	0	14	0	0
LJ6-7	р 3936	65	0	0	0	0	12	8	8	7	0	0
PG1a		70	0	0	0	0	12	0	17	1	0	0
PG1b		72	0	0	0	0	5	5	17	1	0	0
PG2-	5020	63	0	0	0	0	10	0	27	0	0	0
24 PG2- 21	4987	60	0	0	0	0	0	14	25	1	0	0
PG2-	4978	83	0	0	0	0	12	0	3	2	0	0
20a PG2-	5043	75	0	0	0	0	18	0	5	2	0	0
27 PG2-	5076	72	0	0	0	0	5	20	0	3	0	0
31 PG2- 32	5085	74	0	0	0	0	7	14	3	2	0	0
PG2- 41	5196	70	0	0	0	0	13	5	10	2	0	0
PG2- 39	5166	65	0	0	0	0	10	8	17	0	0	0
PG2- 22a	4980	64	0	0	0	0	0	20	14	2	0	0
PG2- 26	4937	75	0	0	0	0	13	0	12	0	0	0
PG2-	4934	80	0	0	6	0	0	7	7	0	0	0
21 PG2- 22b	4935	64	0	0	17	0	0	15	0	4	0	0

PG2-5	4776	67	0	0	14	0	0	5	13	0	0	1
PG2- 20b	4982	38	0	0	0	0	12	17	33	0	0	0
PG2- 30	5066	87	0	0	0	0	0	0	12	1	0	0
PG6-a		60	0	0	0	0	18	7	10	5	0	0
PG6-b		68	0	0	0	0	0	0	2	26	4	0
PG6-c	5142	75	0	0	0	0	0	18	5	2	0	0
TS 5-9		66	0	0	3	0	14	0	6	11	0	0
TS 5- 11		56	0	0	0	0	14	0	18	12	0	0
TS 5- 12		49	0	0	1	0	24	0	10	16	0	0
TS 5- 13		53	18	0	2	0	14	0	2	11	0	0
LGC	5933	65	0	0	3	0	17	0	13	2	0	0
Average	value	67.0	0.8	0. 4	1. 6	0. 5	8. 6	6. 5	9.7	4.4	0.4	0.3

875 -- data unavailable; C1: Pre-TSR calcite; C2: TSR calcite; C3: Post-TSR calcite; D1: Early reflux

dolomite; D2: late burial dolomite

# Table 2.

	Minerals	Initial volume before TSR (cm <sup>3</sup> )	Final volume after TSR (cm <sup>3</sup> )	Relative porosity change (%)	Net porosity change (%)
Case 1	Anhydrite	505	0.0	-6.00	-5.0
16% porosity	Calcite	84	428	4.1	3.4
	Dolomite	7811	7811	0.0	0.0
	Total	8400	8239	-1.92	-1.6
Case 2	Anhydrite	541	0.0	-6.00	-5.4
10%	Calcite	90	459	4.1	3.7
porosity	Dolomite	8369	8369	0.0	0.0
	Total	9000	8830	-2.06	-1.7

## Table 3.

	Initial Porosity before TSR (%)	Final Porosity after TSR (%)	Initial Permeability before TSR (mD)	Final Permeability before TSR (mD)
Case 1	16	17.6	109.7	264.3
Case 2	10	11.7	4.1	10.3