Limited thermochemical sulfate reduction in hot, anhydritic, sour gas carbonate reservoirs: the Upper Jurassic Arab Formation, United Arab Emirates

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Abstract

Limited thermochemical sulfate reduction (TSR) in hot (130–160°C) and anhydrite-rich sour gas reservoir carbonates of the Arab Formation (Upper Jurassic) is manifested by rare calcitization of anhydrite with slightly lower $\delta^{13}C_{VPDB}$ values (-3.2 to -0.1‰) than calcite precipitated in equilibrium with Late Jurassic seawater. Fluid inclusion microthermometry of calcite that has replaced anhydrite indicates that TSR occurred between 130°C and 160°C. The lack of evidence for extensive TSR, despite the suitable current temperatures and abundant sulfates in the gas reservoir, coupled with the more common TSR-related calcite in the flanks (water zone) than crest (gas zone), indicate that: (1) gas emplacement while the reservoir was buried at shallower depth slowed down or inhibited TSR in the crest even when it subsequently reached depths where extensive TSR would occur, and (2) H₂S (up to 38 vol.%) has migrated from the underlying Permo-Triassic and/or Jurassic sulfate-carbonate deposits. This study demonstrates that constraining the timing of hydrocarbon emplacement within the context of burial-thermal history is crucial for a better understanding of the origin of H₂S in hot, anhydrite-rich, sour gas reservoirs.

Key words: Thermochemical sulfate reduction, TSR, diagenesis, carbonate, anhydrite

1. Introduction

Thermochemical sulfate reduction (TSR) is a common reaction in sulfate-bearing carbonate reservoirs at temperatures of around 110–140°C resulting in the production of H₂S by interaction of hydrocarbons and sulfate (Krouse et al., 1988; Worden et al., 2000; Machel, 2001; Worden et al., 2004). The reaction can be envisaged as: Sulfate + hydrocarbons \rightarrow calcite + H₂S (\pm water \pm CO₂ \pm S \pm altered hydrocarbons). The required minimum temperatures and other parameters (e.g. hydrocarbon composition, wettability) controlling the extent of TSR across hydrocarbon fields are still debated topics (Worden et al., 1996; 1998; Machel, 1998; Worden et al., 2004). In addition to the formation of hazardous H₂S, TSR results also in a decrease in reserve volume (Worden and Smalley, 1996; Heydari, 1997; Machel, 2001) and may also improve reservoir quality (Hill, 1995; Ma et al., 2007; Morad et al., 2012; Jiang et al., 2018). Therefore, constraining the origin and distribution of TSR in carbonate reservoirs has important implications for hydrocarbon exploration and production.

The studied anhydrite-rich Upper Jurassic Arab Formation is a major reservoir in several oil and sour gas fields in Abu Dhabi, United Arab Emirates (UAE; Fig. 1A, B). Locally, the H₂S reaches up to 38 vol.% of produced reservoir fluids (Nederlof et al., 2016) imposing a challenge for hydrocarbon production. The extent of TSR and origin of H₂S are not yet fully explored in these reservoirs (Morad et al., 2012; Nederlof et al., 2016). This study uses integrated analyses (petrography, isotope, fluid inclusion microthermometry and Raman spectroscopy) of diagenetic minerals in the Arab Formation reservoir from four wells in order to constrain: (1) timing, extent, conditions and distribution of TSR; and (2) why TSR is limited while the H₂S concentrations are high.

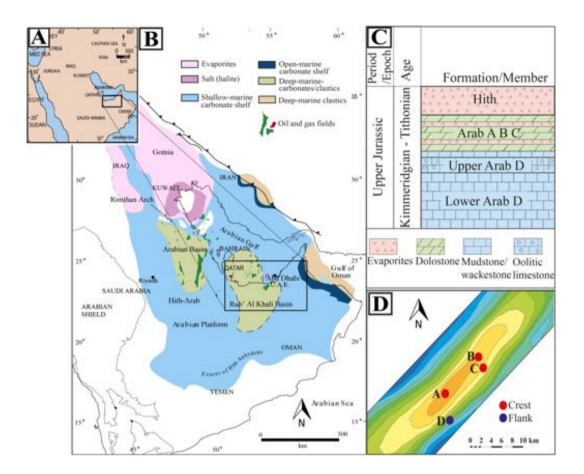


Figure 1. (A) Small-scale map of the Arabian Peninsula. (B) Late Jurassic depositional environments of the Arab and Hith formations (modified after Ziegler, 2001) showing the location of the studied field and several other oil and gas fields onshore and offshore UAE. (C) Simplified stratigraphy of the Arab Formation onshore Abu Dhabi (Morad et al., 2018). (D) Map of the studied anticline showing the sampled wells in the crest (A, B and C) and flank (D).

2. Geological setting

The shallow-water marine platform deposits of the Arab Formation (Kimmeridgian to Tithonian) in Abu Dhabi hydrocarbon fields are composed of carbonates and intercalated anhydrite deposits (Fig. 1C; Morad et al., 2012; Marchionda et al., 2018; Morad et al., 2018).

The Arab Formation in the studied field can be subdivided, from bottom to top, into Arab D (lower and upper D) and C-B-A members (Fig. 1C). Limestones of the lower Arab D Member in the studied field are dominated by dark grey, bioturbated massive mudstones and wackestones that are intercalated with floatstones, which are rich in skeletal fragments (bivalves, gastropods, foraminifera, algae) and mud intraclasts (Marchionda et al., 2018). The mudstones and wackestones were deposited in an outer ramp environment and the floatstones represent storm events (Marchionda et al., 2018). The upper Arab D is composed dominantly of cross or parallel laminated or massive grainstones and packstones (Lawrence et al., 2015; Marchionda et al., 2018). The allochems in the limestones include ooids, peloids, and bioclasts. Deposition of these limestones presumably occurred in shoal and mid ramp environments (Lawrence et al., 2015; Marchionda et al., 2018). The Arab A-B-C members comprise dominantly dolostones and intercalated anhydrites at the top and lime mudstones and wackestones at the base. In the studied field (Fig. 1), cores from Arab A-B-C members were limited to lagoon environment in the anticline flank well. Deposition of anhydrites and associated mudstones (now dolomitized) is interpreted to have occurred in hypersaline lagoon and sabkha settings (i.e. supratidal and lagoon environments; Alsharhan, 1989). The dolostones, which are strongly bioturbated, have mainly mudstones and wackestones precursors. The limestones in the Arab A-B-C members are composed mainly of algal laminated and bioturbated mudstones, wackestones and packstones. These limestones have possibly been deposited in a low energy subtidal lagoon and/or in an intertidal flat (Marchionda et al., 2018). The overlying evaporite dominated Hith Formation (Fig. 1C) is the main seal in the area (Alsharhan and Scott, 2000).

The studied field consists of a gently dipping and elongated anticline (Fig. 1D) that belongs to a series of structural traps that are characterized by NE–SW alignments of fold axes. The anticline was presumably a result of the oblique movement that occurred along a basement fault that was reactivated during the obduction of Semail ophiolites and later collision of Arabia with Iranian Plate (Koyi et al., 2015). The field is situated on the NE part of the Arabian Plate, SW of Abu Dhabi. The measured bottom hole temperatures in the crest and flank of the anticline are around 140°C and 160°C, respectively, indicating that the present day geothermal gradient is around 38°C/km.

The Diyab and Hanifa formations have been proposed as source rocks for the Upper Jurassic reservoirs in the UAE (Alsharhan and Scott, 2000). The Diyab Formation (early Late Jurassic) reached the phase of maximum oil generation in onshore areas of Abu Dhabi during the Early Eocene, whereas the organic-rich Hanifa Formation (early Upper Jurassic) entered the oil generation window during Middle Cretaceous (Ayoub and En Nadi, 2000).

A few studies have previously addressed the diagenesis of the Arab Formation in Abu Dhabi fields (Morad et al., 2012; Nader et al., 2013; Hollis et al., 2017; Morad et al., 2018). However, no studies have focused on the diagenesis related to TSR. The diagenesis in the Arab Formation includes dolomitization of limestones (sabkha and reflux models; Morad et al., 2012) and associated precipitation of anhydrite, calcite associated with dissolution of micritized allochems and stylolitization of the host limestones, burial diagenetic precipitation of sulfates and cementation (mainly saddle dolomite, fluorite and calcite) related to hot basinal brines (Morad et al., 2018).

3. Material and methods

Three hundred and fifty samples were collected from a total of 420 m (1377 ft) studied cores covering the Arab Formation from four wells in the crest and flank of the anticline (Fig. 1D). Transmitted-light, cathodoluminescence (CL), fluorescence, and backscattered electron microscopy were performed on polished thin sections, which were vacuum impregnated with

blue epoxy and stained with alizarin red S and potassium ferricyanide in order to distinguish dolomite from calcite and determine qualitatively their iron content. The abundance and textural types of anhydrite and calcite were quantified by point counting on 500 points with JMicroVision (Roduit, 2005) on fifty selected scanned thin sections. A Technosyn MK2 system and an OPEA system (OPEAFrance) were used for the CL analyses, operating at 10–15 kV gun potential and 300–500 μ A. Gold-palladium covered broken surfaces and carbon coated thin sections were examined with a scanning electron microscope (SEM; FEI Quanta 200 with an acceleration voltage of 30kV) equipped with an Energy Dispersive X-ray Spectrometer.

A computer-automated micro-mill mounted on a binocular microscope and a dentaldrill were used to micro-sample carbonate and sulfate minerals for oxygen, carbon, sulfur and strontium isotope analyses. Contamination by host rock and other phases cannot be ruled out in some cases. Carbon and oxygen isotopes were analyzed for sixty-four samples. The evolved gas (CO₂) was analyzed using a Delta-plus mass spectrometer (University of Windsor, Canada). The isotopic data are presented in δ -notation and reported in per mil (‰) relative to the V-PDB standard. The precision of carbon and oxygen isotope values is better than 0.05%. The ratio of ⁸⁷Sr/⁸⁶Sr isotopes was obtained for two calcite and five sulfate samples by using an automated Finnigan 261 mass spectrometer equipped with nine Faraday collectors (Ruhr-Universität Bochum, Germany). Correction for isotopic fractionation during the analyses was made by normalization to 86 Sr/ 88 Sr = 0.1194. The mean standard error of mass spectrometer performance was ± 0.00003 for standard NBS-987. Sulfur isotope ratios were analyzed for twenty-six sulfate samples and the δ^{34} S values are reported in % relative to CDT. Solid samples were analyzed for sulfur on a Costech Elemental Analyzer (CHNS-O ECS 4010-Italy) coupled to an Isochrom Continuous Flow Stable Isotope Ratio Mass Spectrometer (GVI/Micromass-UK, 1995; University of Waterloo, Canada). Sulfate samples

are corrected by using BaSO₄ standards NBS-127, EIL-42 and EIL-41 (PbSO₄). The error for such clean BaSO₄ standard material is $\pm 0.3\%$ for sulfur.

Aqueous fluid inclusions (FI) in diagenetic calcite and anhydrite were analyzed in seven unstained wafers, which were prepared without inducing substantial sample heating. Special emphasis was made on identifying fluid inclusion assemblages (FIA), i.e. the most finely discriminated groups of petrographically associated fluid inclusions (Goldstein and Reynolds, 1994). CL and BSE imaging proved to be instrumental in determining the origin of the fluid inclusions, by revealing possible relationships to zoning patterns, solid inclusions, or microfractures. Microthermometry, including homogenization temperatures (T_h), and the first (apparent eutectic temperature (T_e) and final ice melting temperatures $(T_m(ice))$, was performed using a Linkam MDS 600 heating-freezing stage coupled with a Nikon LV100 Eclipse combined with a UV-light device (IFPEN, France). The thermocouple was calibrated with synthetic fluid inclusions (H₂O, H₂O-NaCl and CO₂). The Linksys 32 software was used to run the operations for fluid inclusion microthermometry. The total salinity is expressed as wt% NaCl eq. and was calculated from T_m(ice) by using the Bodnar (1993) equation. Heating was performed prior to freezing in order to prevent fluid inclusion reequilibration. These measurements were obtained in the order of increasing T_h (with a precision of 0.5°C) using a calibrated Linkam THMSG 600 stage and low heating rates. Pressure correction was not applied to the measured homogenization temperatures (T_h) and the reported T_h should, if nothing else is stated, be regarded as minimum trapping temperatures.

In two selected samples containing nodular calcite, additional detailed FI petrography was performed on unstained wafers using an Olympus BX51 microscope equipped with UV epi-illumination, and a CITL MK5 cathodoluminescence device operating at 15 kV and 200 μ A. After graphite coating, the polished sections were examined by backscattered electron (BSE) mode using a JEOL JXA-8900 electron microprobe (EMP) at the Complutense

University, Madrid, Spain. The EMP was also used to analyze the composition of mineral inclusions using wavelength-dispersive spectrometry.

Representative fluid inclusions were analyzed in order to detect the types of gas by using a HORIBA Jobin–Yvon LabRAM HR confocal Raman microspectrometer (Eötvös University, Hungary). For the measurements we applied 532 nm emission of a frequency-doubled Nd:YAG laser with a grating of 1800 grooves mm-1. The maximum laser power incident on the sample was c. 40 mW. The detected phases were identified by their characteristic Raman peaks. A 100× microscope objective was used to focus the laser onto the sample and to collect the Raman signal. Spectra were gathered 2–10× accumulations and 2–150 s acquisition time (all depending on the maximum intensity). Raw data were processed using LabSpec v5.25.15 software designed for Jobin-Yvon Horiba LabRam instruments.

Porosity and permeability data were obtained by a contractor for 882 core plugs (diameter of 3.8 cm) from the four wells. Prior to measurements, the core plugs were examined for microfractures, cleaned by oil extractor and dried in a vacuum oven at 60°C for 24 hours. A helium porosimeter was used for porosity measurements and the permeability was measured by helium permeameter by applying 0.689 to 2.76 MPa.

4. Results

4.1 Sulfate fabric and crystal habit

Sulfate minerals with various crystal habits and fabric occur in four depositional facies: (i) Anhydrite nodules and beds are most common in the supratidal/sabkha and intertidal facies (up to 100% anhydrite). These anhydrites are composed of parallel and sub-parallel lath-shaped and felted crystals that are separated by or float in fine-crystalline

dolomite (50–500 μ m long and 10–20 μ m wide; Fig. 2A). The anhydrite nodules are, in some cases, composed of coarser crystals (typically 50–500 μ m long and 50–300 μ m wide; Fig. 2A). These types of anhydrites were analyzed only in the crest of the anticline because cores from these depositional facies were not available in the flank. However, high bulk density log response indicates that anhydrite is present in the supratidal facies of the flank as well. (ii) Anhydrite associated with dolostone beds (0-50% anhydrite) and limestones (0-30% anhydrite) are common in the lagoon facies. Poikilotopic anhydrite engulfs dolomite crystals in the dolostones (Fig. 2B). Anhydrite and minor amounts of celestine occur as lath-shaped crystals (300–3500 μ m) that replace the lagoon limestones (1–2 mm; Fig. 2C). Celestine but no anhydrite was observed in the flank (Fig. 3B). However, cores from the lagoon facies were mainly available from the crest. (iii) Rare anhydrite (0–2%; 1–2 mm) that enclose allochems in host grainstones (Fig. 2D) in the shoal facies on the crest. (iv) Anhydrite and celestine (0–25%) nodules and crystals that replaced host limestone (Fig. 2E, F) were observed in the outer ramp facies in cores from the crest. These anhydrite nodules consist of medium to large sized tabular crystals (20–500 µm long, 50–400 µm wide; Fig. 2E).

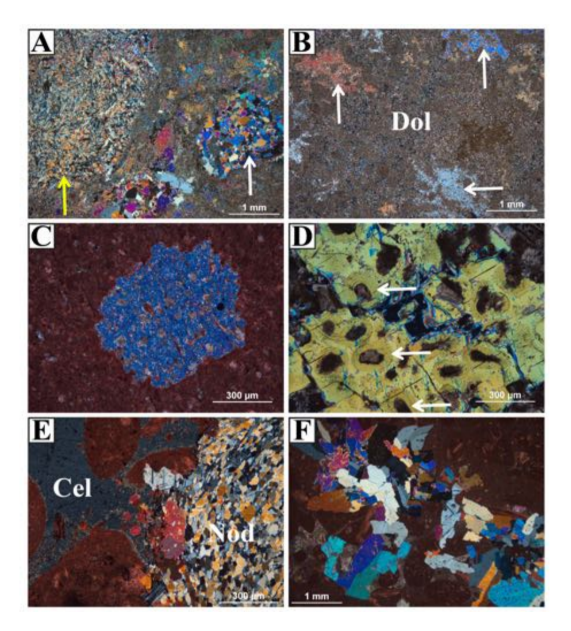


Figure 2. Photomicrographs (XPL) showing anhydrite and celestine in the Arab Formation. (A) Anhydrite nodules in supratidal dolomicrite. Some nodules are composed of fine crystals (yellow arrow) while others by coarse crystals (white arrow). (B) Poikilotopic anhydrite cements (arrows) engulfing dolomite crystals (Dol) in lagoon facies. (C) Replacive coarse anhydrite crystal are often scattered in the lagoon mudstone facies. (D) Anhydrite crystals that have cemented intergranular pores and replaced ooids (arrows) in a shoal grainstone; note the presence of relict concentric lamellae of the replaced ooids. (E) Celestine cement (Cel) and anhydrite nodule (Nod) in outer ramp mudstone. (F) Replacive anhydrite crystals in outer ramp wackestone.

Aggregates of fine to blocky (5 μ m up to 2 mm) calcite crystals are observed within anhydrite and, less commonly, celestine nodules and cement (Fig. 3, 4). Calcite within sulfate crystals and nodules are more common at the edges than in the centers of these sulfate phases (Fig. 3C, G, 4C, D). The anhydrite is locally partly dissolved when observed in close relationship with calcite (Fig. 4E). Nodular, tabular and blocky calcites are more common in the outer ramp than in the supratidal and lagoon facies (Fig. 3, 4). Minor amounts of such calcite are observed in the shoal facies on the flank of the anticline (Fig. 3F). Tiny remnants of anhydrite within calcite cement were observed in the outer ramp facies in the flank (Fig. 3H; < 1 % of the rock volume), while a few celestine crystals associated without any or only small amounts of calcite were observed in the lagoon facies in the flank (Fig. 3B).

Figure 3. Photomicrographs comparing the intimate association of sulfate and calcite in the various depositional facies from the crest and flank. (A) Anhydrite nodules (Anh) separated by dolomicrite (Dol) in supratidal/sabkha facies (crest; PPL). (B) Same photomicrographs as in (A) with XPL. Cores from the supratidal/sabkha facies from the flank were not available in this study. However, bulk density logs suggest that anhydrite is present here too. (C) Minor calcite (Cal) is observed along the outer rim of anhydrite cement (Anh) in a partly dolomitized (Dol) mudstone from the lagoon facies (crest; PPL). (D) Minor calcite (white arrow) engulfing celestine crystals (Cel) in the lagoon facies (flank; PPL). (E) Anhydrite (Anh) enclosing calcite cement and allochems in a shoal grainstone (crest; PPL). (F) Calcite cement (Cal) and saddle dolomite (SD) in the shoal facies (flank; PPL). (G) Calcite (Cal) that

is enclosing anhydrite cement (Anh) in the outer ramp facies (crest; PPL). (H) Tabular calcite in the outer ramp facies. Small remnants of anhydrite (arrows) are observed within this calcite. Saddle dolomite (SD) is observed in close association with the tabular calcite (flank; PPL).

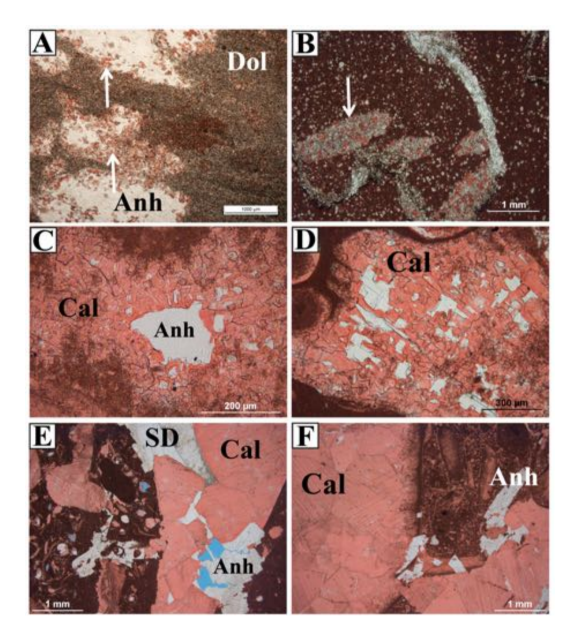


Figure 4. Photomicrographs (PPL) of calcite (Cal; stained red by alizarin red) in close relationship with anhydrite on the crest of the anticline. (A) Calcite concentrated mainly along the edges of the anhydrite nodules (arrow) in a partly dolomitized mudstone (outer ramp facies). (B) Partly dolomitized calcite crystals in the lagoon facies. (C) and (D) Calcite

engulfing anhydrite cements (Anh). Note that calcite is more common at the edges of the anhydrite. (E) and (F) Coarse crystalline calcite engulfing anhydrite cement (Anh); note the partial dissolution of the anhydrite (E; blue color). Saddle dolomite (SD) is often associated with the calcite.

The margins of anhydrite nodules locally contain tabular calcite crystals (Fig. 5A). The calcite in anhydrite nodules is non-fluorescent, dull brownish mottled to homogeneous CL (Fig. 5B), and contains abundant, relatively large (\sim 10–100 µm) presumably primary biphase aqueous inclusions (Fig. 5B, C), which are associated with dolomite and rare anhydrite inclusions (Fig. 5D, E). Saddle dolomite is locally engulfed by the calcite, especially along the margins of calcite nodules. Small (1–10 µm wide) discrete pyrite crystals are observed in close association with nodular and tabular calcite.

It was not possible to compare the presence/abundance of nodular and tabular shaped in the supratidal and lagoon facies between crest and flank because cores from these facies were not available or limited in the flank.

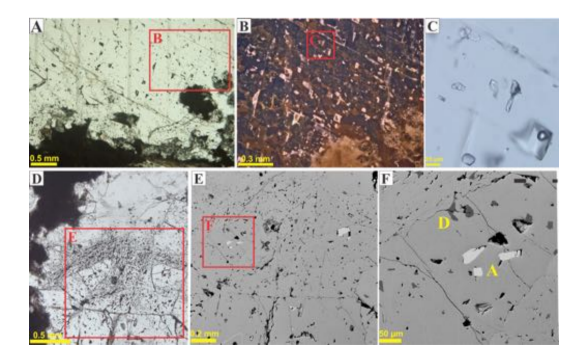


Figure 5. Petrography of fluid inclusions in nodular calcite. (A) Photomicrograph (PPL) of the margin of a large calcite nodule showing relatively abundant aqueous inclusions (dark spots). (B) CL image of (A), showing the markedly mottled characteristics and the abundant aqueous inclusions (bright spots). (C) Photomicrographs (PPL) of the biphase aqueous inclusions in the calcite in (B). (D) to (F): Photomicrograph (PPL; D) and BSE images (E, F) of calcite showing abundant corroded remnants of dolomite (D) and anhydrite (A). The rectangles in the photomicrographs and BSE images represent the next figure.

4.3 Isotope geochemistry of carbonates and sulfates

Calcite nodules with various amounts of anhydrite in the outer ramp facies have $\delta^{13}C_{VPDB}$ values ranging from -3.2 to -0.1‰ (av. -1.3‰) and $\delta^{18}O_{VPDB}$ values from -7.7 to -4.0‰ (av. -4.9‰; Fig. 6). Two Sr-isotope analyses of such calcite nodules have values of 0.707034 and 0.707057. Bulk limestone samples display $\delta^{13}C_{VPDB}$ values from -3.0 to +3.6‰ (av. +1.9‰) and $\delta^{18}O_{VPDB}$ values from -6.6 to -2.9‰ (av. -4.9‰; Fig. 6)

The $\delta^{34}S_{CDT}$ values of anhydrite nodules, beds and cements in the supratidal and lagoon facies show similar values (+17.4 to +20.2‰; av. +18.7‰). The $\delta^{34}S_{CDT}$ values obtained for sulfate cements (anhydrite and celestine) in the outer ramp facies range from +23.1 to +24.7‰ (av. +23.8‰). Sulfates in the supratidal/lagoon and outer ramp facies reveal similar ⁸⁷Sr/⁸⁶Sr ratios between 0.70694 and 0.70698 (av. 0.70696) and between 0.70693 and 0.70701 (av. 0.70696), respectively.

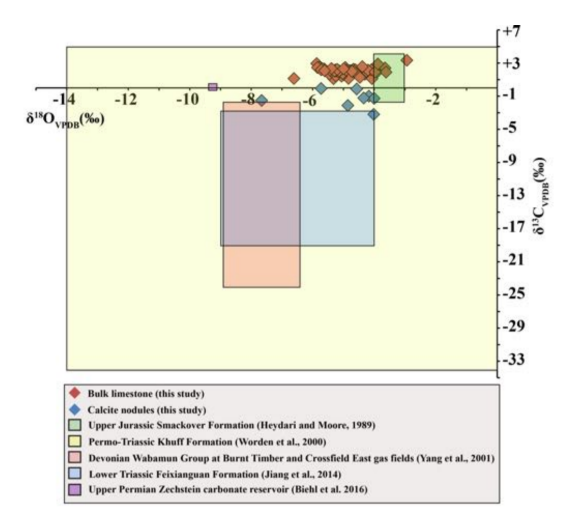


Figure 6. Carbon *versus* oxygen isotope cross plot showing that nodular calcite has lower $\delta^{13}C_{VPDB}$ values than the bulk limestone samples from the Arab Formation. Isotope compositions of calcite associated with TSR from other studies are included for purpose of comparison. The calcites in this study have slightly depleted $\delta^{13}C_{VPDB}$ values compared to bulk limestone samples.

4.4 Fluid inclusion petrography, microthermometry and Raman spectroscopy

Fluid inclusion (FI) petrography, microthermometry and Raman spectroscopy data were obtained from nodular and tabular calcite in the flanks, whereas FI in similar calcites from crest wells are rare and have thus not been analyzed. Observation of the FI under UVlight did not reveal any hydrocarbons containing aromatic compounds. Nodular and tabular calcite in the flank of the structure contain abundant aqueous and possibly primary FI. The calcite contains abundant clusters of aqueous biphase FI, particularly near the nodule margins (Fig. 5). The FI show negative crystal shapes near the margins (Fig. 5F), whereas more irregular shapes are observed towards the central parts of the calcite nodules (Fig 5D, E).

The overall T_h range is 123°C to > 200°C (N=170). However, most T_h measurements fall between 135 and 157°C (Fig. 7A). In individual fluid inclusion assemblages (FIA), the T_h measurements are mostly within a 10-15°C range, attesting that they did not experience significant post-entrapment modifications (e.g. neckingdown, stretching or leakage due to thermal reequilibration). Two FIA yielded relatively inconsistent T_h (140 to > 200°C, and 125 to 165°C). However, in these FIA most inclusions homogenize in a narrow range (140–155°C and 137-151°C, respectively). The higher temperature tails recorded in these two FIA (160-165°C and 163 to > 200°C) are formed by FI showing highly variable liquid to vapor ratios. The high temperature limit (200°C) is arbitrary, as some of the FI retained bubbles at this temperature. These bubbles show consistent volume changes during heating-cooling cycles, indicating that they have not resulted from leakage. The two FI that homogenize at temperatures below 130°C can belong to a different calcite generation or due to some error during measurements. The homogenization temperatures of the FI in the central parts of the calcite nodules are, in general lower, than near the margins. In aqueous and possibly secondary FI located along trails cutting through calcite nodules, the overall Th range is 138-160°C (N=75). However, in individual FIA the T_h data is typically highly consistent. For instance, all the FI measured along a single trail (N=38) yielded T_h ranging from 148 to 153°C. The salinity of the aqueous FI in the calcite nodules ranges from 20.3 to 22.5 wt% NaCl eq. (Fig. 7B); the apparent eutectic temperatures range from around -50 to -28 °C.

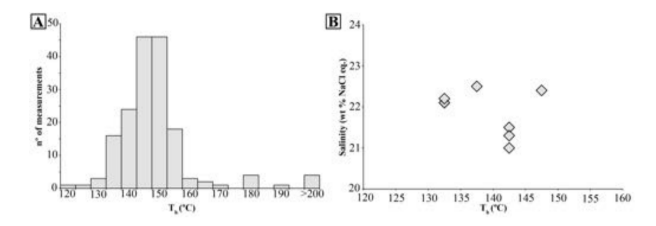


Figure 7. Homogenization temperatures (T_h) and salinity data of aqueous and presumably primary fluid inclusions in nodular and tabular calcite obtained from the Arab Formation. (A) Histogram for all measured aqueous and presumably primary fluid inclusions. Most T_h values range between 135 and 155°C. (B) Cross-plot showing that there is no correlation between T_h and salinity.

Raman spectroscopy of fluid inclusions in nodular calcite reveals the presence of methane and carbon dioxide in the gas phase. Hydrogen sulfide, methane and minor ethane and propane were detected in fluid inclusions in pore-filling calcite and in calcite replacing saddle dolomite, which are not associated with anhydrite.

4.5 Gas composition

Gas samples from well C (Fig. 1D) are composed mainly of methane, hydrogen sulfide and minor amounts of ethane and propane (Table 1). The H₂S content range between 19.55 and 23.74% and is slightly higher in the lower Arab D Member (av. 23.08%) than in upper Arab D (av. 22.15%) and Arab A-B-C members (av. 21.37%). The dryness coefficient $(C_1/(C_1-C_3))$ of the gas, which indicates the relationship between the amounts of methane (C₁), ethane (C₂) and propane (C₃), varies between around 87 and 92. The dryness coefficient

is slightly higher in lower (av. 91.00%) and upper (av. 91.09%) Arab D than in Arab A-B-C (av. 88.77%) members. The water (vapor) content in the crest of the anticline (i.e. gas zone) is around 1.5%.

Member	Depositional facies	Gas chemical composition (%)				Dryness coefficient (%)	
		C ₁	C ₂	C ₃	H_2S	$C_1/(C_1-C_3)$	
A-B-C	Supratidal/lagoon	59.70	6.04	2.78	19.71	87.13	
A-B-C	Supratidal/lagoon	60.20	6.02	2.66	19.55	87.40	
A-B-C	Supratidal/lagoon	60.00	6.06	2.74	19.71	87.21	
A-B-C	Supratidal/lagoon	60.00	6.05	2.76	19.69	87.20	
A-B-C	Supratidal/lagoon	59.75	6.04	2.78	19.72	87.14	
A-B-C	Supratidal/lagoon	59.48	6.04	2.83	19.85	87.02	
A-B-C	Supratidal/lagoon	58.90	4.38	1.93	22.98	90.32	
A-B-C	Supratidal/lagoon	58.60	4.38	1.93	23.11	90.28	
A-B-C	Supratidal/lagoon	58.99	4.38	1.87	23.07	90.42	
A-B-C	Supratidal/lagoon	59.10	4.37	1.89	22.94	90.42	
A-B-C	Supratidal/lagoon	58.87	4.38	1.91	23.06	90.35	
A-B-C	Supratidal/lagoon	58.83	4.38	1.92	23.10	90.33	
Upper D	Shoal	60.37	4.23	1.67	22.11	91.10	
Upper D	Shoal	60.55	4.23	1.64	21.95	91.16	
Upper D	Shoal	61.24	4.23	1.59	21.34	91.32	
Upper D	Shoal	60.21	4.24	1.65	22.28	91.09	
Upper D	Shoal	59.66	4.25	1.69	22.61	90.95	
Upper D	Shoal	61.88	4.17	1.49	20.95	91.62	
Upper D	Shoal	59.11	4.27	1.76	23.06	90.74	
Upper D	Shoal	59.22	4.26	1.78	22.92	90.74	
Lower D	Outer ramp	61.38	4.07	1.45	21.31	91.75	
Lower D	Outer ramp	59.14	4.19	1.71	23.13	90.93	
Lower D	Outer ramp	60.57	4.15	1.51	22.13	91.45	
Lower D	Outer ramp	58.81	4.20	1.70	23.43	90.88	
Lower D	Outer ramp	58.27	4.20	1.74	23.66	90.75	
Lower D	Outer ramp	58.35	4.20	1.74	23.59	90.76	
Lower D	Outer ramp	58.11	4.20	1.75	23.65	90.71	
Lower D	Outer ramp	58.11	4.21	1.70	23.74	90.77	

Table 1. Gas composition and the dryness coefficient of gases in various depositional facies

 and formation members from well C.

Porosity and permeability values of carbonates in the various depositional facies and corresponding lithologies and limestone textures in the crest and flank are shown in Table 2. Porosity and permeability are higher in carbonates from the crest than from the flank. The highest porosity and permeability values in the Arab A-B-C members (i.e. supratidal and lagoon facies) are encountered in dolostones that are poor in anhydrite cement. The highest porosity and permeability values in upper D (i.e. shoal facies) are in grainstones with high intergranular macro- and microporosity. The lower D (i.e. outer ramp facies) is characterized by overall low porosity and permeability values are attributed to microfractures.

Table 2. Porosity and permeability values of samples from the various depositional facies and formation members in the crest and flank of the anticline.

	Depositional facies	Lithology	Porosity (%)	Average	Permeability (mD)	Average
Crest						
Arab A-B-C	Supratidal	Anhydrite/dolomicrite	0.1–6.6	0.9	0–11.6	0.5
Arab A-B-C	Lagoon	Dolostones	0.2–28.8	13.5	0–43.9	5.1
Arab A-B-C	Lagoon	Lime mud-/wackestone	0.2–16.3	2.7	0–19.1	0.8
Upper D	Shoal	Lime pack-/grainstone	1.8-28.8	18.5	0–254.0	3.9
Lower D	Outer ramp	Lime mud-/wacke-	1.1–19.1	6.7	0-82.0	1.0
		/floatstone				
Flank						
Arab A-B-C	Supratidal	No data	No data	No data	No data	No data
Arab A-B-C	Lagoon	Lime mud-/wackestone	0-1.3	0.6	0–40.1	1.9
Upper D	Shoal	Lime pack-/grainstone	0.4–3.2	1.7	0–1.3	0.2
Lower D	Outer ramp	Lime mud-/wacke-	0.2-12.7	1.3	0–54.1	2.2
		/floatstone				

5. Discussion

Integrating the results from petrography, stable isotopes and fluid inclusion microthermometry and Raman spectroscopy into the burial-thermal history allowed us to constrain the reasons behind the limited evidence of TSR, origin of abundant H_2S and relative timing of sour gas emplacement in the studied Upper Jurassic Arab Formation reservoir.

5.1 Evidence and conditions of thermochemical sulfate reduction

Hydrogen sulfide (H₂S) is abundant in Arab Formation gas reservoirs in Abu Dhabi, particularly in the onshore fields, which are buried deeper than in offshore fields (Grötsch et al., 2003). H₂S in carbonate reservoirs can be produced by both bacterial sulfate reduction (BSR) and thermochemical sulfate reduction (TSR; Machel, 2001). The following lines of evidence suggest that limited TSR, i.e. the reaction between hydrocarbons (methane, ethane and propane) and anhydrite, can account for the formation of limited amounts of H₂S and calcite in the studied reservoir: (i) the presence of petrographic evidence for replacement of anhydrite and minor celestine by calcite (Fig. 3, 4). This evidence is indicated by calcite nodules and crystals that contain tiny remnants of anhydrite and resemble anhydrite precursors (e.g. nodules and tabular crystals). (ii) The lower $\delta^{13}C_{VPDB}$ values (-3.2 to -0.1‰) of calcite that has replaced anhydrite compared to those of the host limestones (av. +1.9‰), which have $\delta^{13}C_{\text{VPDB}}$ values of calcites precipitated in equilibrium with Late Jurassic seawater $(\delta^{13}C_{VPDB} = -0.1 \text{ to } +2.5\%)$; Veizer et al., 1999). Wide ranges of $\delta^{13}C_{VPDB}$ values have been reported in the literature for calcite derived from TSR. While most values for such calcite are far more negative than those obtained in this study (Fig. 6; Worden et al., 2000; Yang et al., 2001; Jiang et al., 2014), denoting derivation of dissolved carbon from the oxidation of

hydrocarbons, Heydari and Moore (1989) have reported similar to considerably higher $\delta^{13}C_{VPDB}$ values (-1.7‰ to +4.2‰) in the Jurassic Smackover Formation (USA). These authors suggested that such high values were because the $\delta^{13}C$ was mainly derived from host rock. The marginally depleted $\delta^{13}C_{VPDB}$ values of such calcite corroborate the limited TSR in the studied Arab Formation, i.e. most dissolved carbon was derived from the host limestone and not from the oxidization of methane. (ii) The relatively high precipitation temperatures inferred form FI homogenization for the calcite that has replaced anhydrite ($\sim 130-160^{\circ}$ C) are around or above the minimum temperatures required for TSR (i.e. >110 to 140°C; Worden et al., 1995; Machel, 1998; Worden et al., 2004). Knowing that the vapor phase in the analyzed fluid inclusions in this study contains some methane (as detected by Raman spectroscopy), the pressure corrections should be minimal (Goldstein and Reynolds, 1994), i.e. the T_h values are probably very close to true entrapment temperatures. (iv) The low dryness coefficient $(C_1/(C_1-C_3) = ~ 87-92$; Table 1) of gas samples obtained from the various depositional facies is consistent with relatively limited TSR in the reservoir because ethane (C_2) and propane (C_3) are preferentially removed during TSR (Krouse et al., 1988). (v) The presence of pyrite and sphalerite in the studied formation (Morad et al., 2018), as well as CO₂ and H₂S detected by Raman spectroscopy in fluid inclusions, which are all typical byproducts of TSR (e.g. Machel, 2001).

5.2 Temperature, fluid composition and timing of TSR

The minimum temperature required for TSR is controversial and has been suggested to vary from around 110 to 140°C (Krouse et al., 1988; Heydari and Moore, 1989; Machel, 1998; Worden et al., 2000, Yang et al., 2001; Worden et al., 2004). The fluid inclusion microthermometry of calcite that has replaced anhydrite in the studied reservoir suggests that TSR occurred mostly between 135° C and 155° C. These temperatures are comparable to bottom hole temperatures in the crest and flanks, respectively. The homogenization temperatures (T_h) coupled with the modeled burial-tectonic history (Al Darmaki et al., 2014; Fig. 8) suggest that TSR commenced at around 2.7 km depth around 35–50 Ma ago (Early to Late Eocene). This interpretation is made derived assuming that TSR starts at around 130°C and a mean geothermal gradient of 38°C/km (Fig. 8, 9).

By using the main T_h range (135–155°C), the $\delta^{18}O_{VPDB}$ values (-7.7 to -4.0‰) of TSR calcite and the calcite-water oxygen isotope fractionation equation of Friedman and O'Neil (1977), a $\delta^{18}O_{SMOW}$ signature of +9.2 to +14.5‰ can be deduced for the water from which calcite was formed. Such $\delta^{18}O_{SMOW}$ values, which suggest that TSR was mediated by geochemically evolved basinal brines (cf. Zeeh et al., 1995; Calvo et al., 2011; Dickson and Kenter, 2014), have been reported for TSR calcites elsewhere by Yang et al. (2001; +7.8 to +11.7‰) and Machel and Buschkuehle (2008; +4 to +13‰). Moreover, the salinity data (20.3 and 22.5 wt% NaCl eq.) obtained from fluid inclusion microthermometry of calcite are typical for basinal brines. Based on the apparent eutectic temperatures (-50 to -28°C) of the fluid inclusions of TSR calcite, the fluids can be characterized as NaCl-CaCl₂-MgCl₂ dominated (cf. Goldstein and Reynolds, 1994).

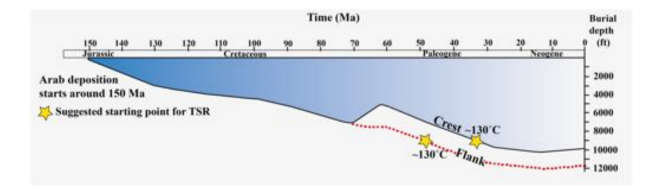


Figure 8. Burial history curve of the Arab Formation for the studied crest and flank wells spanning from deposition to present time (modified from Al Darmaki et al., 2014). A small

uplift and development of anticline of the field during Late Cretaceous were associated with the obduction of Oman ophiolites. TSR is suggested to start at around 130°C, meaning that TSR starts earlier on the flanks and then on the crest.

5.3 Origin of hydrogen sulfide – evidence of limited TSR

The presence of up to 38 vol.% H_2S in the studied field (Nederlof et al., 2016) suggests derivation by extensive TSR. However, the unexpected presence of only limited amounts of calcitized anhydrite indicates that TSR is relatively limited in the studied sour gas reservoir. Hence, the high amounts of H₂S have presumably been mostly derived from elsewhere in the basin. Based on ${}^{34}S_{CDT}$ values of H₂S in the lower parts if the formation (i.e. lower Arab D), Nederlof et al. (2016) suggested that the H₂S was derived from TSR in the underlying early Upper Jurassic Hanifa Formation, whereas H₂S in the upper parts of the formation (i.e. Arab A, B, C) was formed in-situ. These authors have based their interpretation on the similar ${}^{34}S_{CDT}$ values of the anhydrite and H₂S, i.e. assuming there will be only minor or no sulfur isotopic fractionation during TSR (Yang et al., 2001). However, the ³⁴S_{CDT} values obtained in the present study for anhydrites in the Arab D, which were not analyzed by Nederlof et al. (2016), are similar to the ³⁴S_{CDT} values of the H₂S in the Arab D (Nederlof et al., 2016). Consequently, at least some of the H₂S in the Arab D was also derived from *in-situ* TSR. Moreover, lower amounts of H₂S in the upper parts of the formation compared to the lower parts (Nederlof et al., 2016) supports our interpretation that TSR was relatively more extensive in the lower parts (Fig. 3). Nevertheless, the limited amounts of sulfates in the lower parts of the formation indicate that all the bulk of the abundant H₂S could not be derived from TSR within the Arab Formation. Hence, migration of H₂S from underlying formations (Grötsch et al., 2003) is a more plausible explanation for the high

amounts of H_2S in the Arab Formation. The underlying Permo-Triassic Khuff Formation (Worden et al., 2000) and/or early Upper Jurassic Hanifa (Nederlof et al., 2016) are possible external sources of the H_2S .

The reasons behind the limited extent of TSR despite the presence of ideal conditions in the studied succession, i.e. sufficiently high temperatures and presence of abundant sulfates, are not fully understood. One reason could be gas emplacement during shallow burial before temperatures increased high enough to ensure extensive TSR. Numerous studies have concluded that high hydrocarbon saturation in carbonate reservoirs might retard or even stop diagenesis including extensive TSR (Machel et al., 1995; Machel, 1998) because minerals (in this case anhydrite) have nil solubility in gas (Fezel and Schatzinger, 1985; Heasley et al., 2000; Cox et al., 2010; Paganoni et al., 2016; Morad et al., 2018). Thus, while TSR was presumably retarded in the crest (i.e. gas zone) due to gas emplacement, it continued during gas migration through the water-saturated flanks. This scenario is depicted in figure 9. Migration of hydrocarbons in the flank is supported by the presence of methane, ethane and propane in fluid inclusions in calcite cements. Moreover, the somewhat more extensive TSR in the lower than in the upper parts of the studied formation might be because: (i) the anhydrite nodules in the upper parts contain finer crystals than those in the lower parts. Worden et al. (2000) noted that finer crystals experience more TSR than coarser crystals in the Permo-Triassic Khuff Formation. However, we did not observe any clear correlation between crystal size and TSR in this study; or (ii) gas emplacement is expected to start in the upper parts of the succession and continue downwards (Fig. 9). This is the more plausible scenario because gradual gas emplacement would lead to earlier retardation of TSR in the upper parts compared to the lower parts of the formation. Nevertheless, anhydrite has a greater surface affinity for water than non-polar alkanes (e.g. methane, ethane, propane), which suggests that alkane-anhydrite systems are always water-wet rather than gas-wet, even after gas emplacement (Worden et al., 2000). Hence, even if the overall diagenesis was retarded substantially in the crest after gas emplacement, anhydrite dissolution could have continued in the presence of irreducible water. However, the low total water content (1.5%) in the reservoir suggests that the anhydrite is presumably gas wet (Fig. 9).

Figure 9. Schematic conceptual model of the extent of TSR in the Arab Formation across the anticline of the field. (A) Formation of the anticline and gas migration occurred in Late Cretaceous during the obduction of the Oman ophiolites. TSR started in the flanks at around 130°C in Early Eocene during gas migration when the reservoir temperatures were too low to cause extensive TSR in the crest. (B) TSR continued in the flanks and commenced in the lower parts (i.e. outer ramp) of the crest but not in the upper parts (i.e. supratidal and lagoon) because of early gas emplacement in Early to Late Eocene. (C) TSR continued in the water-saturated flanks, whereas stopped or slowed down significantly in the crest because of gas emplacement during the Late Eocene.

5.4 Interplay between TSR and porosity and permeability

It has been argued in the literature that there is interplay between porosity and permeability of the host carbonate reservoir and the rates of TSR. High porosity and permeability, which implies increase in rates of fluid flux and water/rock ratios, could increase the rates of anhydrite dissolution and migration of hydrocarbons, and hence the rate of TSR (Worden et al., 2000; Machel, 2001). Despite this, we found that the lower porosity and permeability in the mud dominated carbonates of outer ramp facies exhibit evidence of relatively more extensive TSR compared to the higher porosity and permeability dolostones in

the lagoon facies in the crest. Therefore, the rates of anhydrite dissolution and gas migration to the reaction site are presumably not controlling the extent of TSR in the Arab Formation. In the flank, however, the overall significantly lower porosity and permeability, which has been suggested to be because calcite cementation associated with stylolitization continued after gas emplacement in the crest (Morad et al., 2018), could hinder gas migration to the upper parts of the formation and, thus, prevent TSR. TSR might be common in the outer ramp facies in the flank because these rocks contain stylolites, which act as conduits for flux of gas and other fluids. This postulation is supported by: (i) the common presence of saddle dolomite and fluorite cements along stylolites and related fractures (Morad et al., 2018), which are more frequent and better developed in the flanks than in the crest. It has been suggested in the literature that stylolites and fractures act as conduits for flux of hot basinal fluids (Davies and Smith, 2006; Mansurbeg et al., 2016; Paganoni et al., 2016). (ii) The presence of CH₄ bearing fluid inclusions in these cements, indicating that gases required for TSR was present. Such types of cements and fluid inclusions are common in the lower parts while absent in the upper parts of the formation, which witness difference in rates of fluid flow.

 H_2S is an aggressive gas that can cause mineral dissolution. Accordingly, TSR has been suggested to improve reservoir quality (e.g. Hill, 1995; Ma et al., 2007; Morad et al., 2012; Jiang et al., 2018). However, we did not observe any evidence indicating that TSR affected the reservoir quality in the studied succession, which is in agreement with other studies (e.g. Machel et al., 1995; Heydari, 1997; Machel, 2001; Hao et al., 2015). A possible scenario is that the methane, ethane and propane were emplaced in the reservoir before H_2S , and hence inhibited any further diagenesis including dissolution. Instead, the better reservoir properties of the limestones on the crest compared to those on the flanks was attributed by Morad et al. (2018) to preservation of porosity and permeability due to gas emplacement rather than creation by mineral dissolution. Morad et al. (2018) attributed the destruction of reservoir quality in the flanks to continuous chemical compaction and cementation.

6. Conclusions

The main findings of this integrated petrographic, isotope, fluid inclusion microthermometry and Raman spectroscopy study of thermochemical sulfate reduction (TSR) in the Jurassic hot, anhydrite-rich carbonate reservoirs include:

- Despite the presence of abundant H_2S in the sour gas reservoir, there is limited evidence of TSR, which is manifested by rare calcitization of anhydrite. Carbon isotope composition of calcite replacing anhydrite ($\delta^{13}C_{VPDB} = -3.2$ to -0.1%) indicates that dissolved carbon was derived mainly from the host Jurassic carbonates and only partly from oxidation of hydrocarbons.
- The limited TSR despite the high temperatures reached by the reservoir and presence of abundant anhydrite deposits in the crest (i.e. gas zone) is attributed to emplacement of gas before burial of the reservoir to depths that promote TSR. In other words, early gas emplacement retarded diagenesis including TSR.
- Consequently, most of the abundant H₂S present in the sour gas reservoir has migrated from underlying anhydrite-rich early Upper Jurassic and/or Permo-Triassic carbonate deposits that are undergoing TSR.
- Fluid inclusion microthermometry of calcitized anhydrite indicates that limited TSR occurred at temperatures between 140°C and 155°C in the presence of geochemically evolved (δ¹⁸O_{SMOW} = +9.2 to +14.5‰), highly saline (20.3 and 22.5 wt% NaCl eq.), NaCl-CaCl₂-MgCl₂ dominated brines.
- There was no impact of TSR on mineral dissolution and thus not on reservoir quality

presumably because gas (methane ethane and propane) migration and emplacement in the reservoir occurred before the migration of H₂S. Hence, gas emplacement inhibited any further diagenesis including any possible dissolution associated with TSR.

- The more extensive TSR in the lower than upper parts of the crest is attributed to gradual gas saturation of the reservoir starting in the upper and then in the lower part of the anticline.
- This study demonstrates that the presence of abundant H₂S in anhydrite-rich, hot carbonate sour gas reservoir does not automatically imply extensive TSR in-situ. Derivation of H₂S from other parts of the basin where abundant H₂S is formed by TSR should have important implication for origin and timing of fluid (including hydrocarbons) migration and hence for successful exploration strategies in sedimentary basins.

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