How important is carbonate dissolution in buried sandstones: evidences from petrography, porosity, experiments, and geochemical calculations

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Received: 15 June 2018 / Published online: 30 July 2019 © The Author(s) 2019

Abstract
Burial dissolution of feldspar and carbonate minerals has been proposed to generate large volumes of secondary pores in subsurface reservoirs. Secondary porosity due to feldspar dissolution is ubiquitous in buried sandstones; however, extensive burial dissolution of carbonate minerals in subsurface sandstones is still debatable. In this paper, we first present four types of typical selective dissolution assemblages of feldspars and carbonate minerals developed in different sandstones. Under the constraints of porosity data, water–rock experiments, geochemical calculations of aggressive fluids, diagenetic mass transfer, and a review of publications on mineral dissolution in sandstone reservoirs, we argue that the hypothesis for the creation of significant volumes of secondary porosity by mesodiagenetic carbonate dissolution in subsurface sandstones is in conflict with the limited volume of aggressive fluids in rocks. In addition, no transfer mechanism supports removal of the dissolution products due to the small water volume in the subsurface reservoirs and the low mass concentration gradients in the pore water. Convincing petrographic evidence supports the view that the extensive dissolution of carbonate cements in sandstone rocks is usually associated with a high flux of deep hot fluids provided via fault systems or with meteoric freshwater during the eodiagenesis and telodiagenesis stages. The presumption of extensive mesogenetic dissolution of carbonate cements producing a significant net increase in secondary porosity should be used with careful consideration of the geological background in prediction of sandstone quality.

Keywords Mesodiagenetic · Carbonate dissolution · Petrography · Geochemical · Buried sandstones

1 Introduction

The term secondary porosity refers to pore space resulting from the post-depositional dissolution of detrital grains or cements (Taylor et al. 2010). Ten genetic mechanisms have been proposed for the generation of aggressive fluids capable of dissolving minerals in sandstones, which are meteoric water penetration (Emery et al. 1990), mixing corrosion (Edmunds et al. 1982; Plummer 1975), acidic fluids generated from CO2 produced by the thermal maturation of organic matter (Schmidt and McDonald 1979a; Surdam et al. 1989; Surdam and Boese 1984), carboxylic acids generated during the thermal maturation of organic matter (Surdam et al. 1989; Surdam and Boese 1984), acidic fluids generated by clay mineral reactions (Giles and Marshall 1986), acid fluids generated by thermogenic sulfate reduction (TSR) and bacterial sulfate reduction (BSR) (Machel 2001; Machel et al. 1995), deep hot fluids (Taylor 1996), acidic...
fluids generated by silicate hydrolysis (Hutcheon and Abercrombie 1990), acidic fluids generated by silicate–carbonate interactions (Smith and Ehrenberg 1989), aggressive fluids due to cooling of formation fluids (Giles and De Boer 1989), and hot alkaline brines (Pye 1985).

The idea that the sandstone porosity can be significantly increased via burial dissolution of minerals (e.g., carbonate cements, feldspars) at depths of approximately 3 km by CO₂ and organic acids originating from kerogen maturation in source rocks was proposed in 1970s to 1980s (Schmidt and McDonald 1979a; Surdam et al. 1989; Surdam and Boese 1984) (Fig. 1a). Based on petrographic identification and interpretation, this idea has been prominent in the literature on sandstone diagenesis for about 40 years (Bjørlykke and Jahren 2012; Boggs 2011; Burley 1986; Dutton and Wil¬lis 1998; Higgs et al. 2010; Khidir and Catuneanu 2010; Kordi et al. 2011; Schmidt and McDonald 1979a; Shan¬mugam 1984; Xi et al. 2016; Wilkinson et al. 1997; Yuan et al. 2015a, b, c). At the same time, however, the advent of the deep burial dissolution proposals has caused intense debates (Fig. 1b). The opposing views are centered on the apparent lack of viable geochemical mechanisms by which dissolution and mass transfer could occur in the subsurface rocks (Bjørkum et al. 1998; Bjørlykke 1984; Bjørlykke and Jahren 2012; Li et al. 2017; Taylor et al. 2010; Yuan et al. 2013a, b, 2015a, b, c). Recently, rock diagenesis and the significance of secondary pores generated by the burial dissolution of feld¬spars and carbonate minerals have been reviewed within the constraints of petrography, porosity data, and the openness versus closeness of geochemical systems (Bjørlykke 2014; Bjørlykke and Jahren 2012; Ehrenberg et al. 2012; Taylor et al. 2010; Yuan et al. 2013a, b). These reappraisals showed that burial-induced carbonate dissolution in sandstones and carbonates is commonly insignificant. This conclusion is not new (Ehrenberg et al. 2012); however, the retrospective nature of these new presentations is striking because the subjective idea that up to 20% secondary porosity can be formed by burial dissolution of minerals still persists in some very recent publications (Khidir and Catuneanu 2010; Kordi et al. 2011) and textbooks (Boggs 2011). Particularly, this idea

| Kerogen evolution stage | Primary Φ, % | Secondary Φ, % | Porosity Φ, % | Reservoir spaces evolution |
|-------------------------|--------------|---------------|--------------|---------------------------|
| Immature stage:        |              |               |              |                           |
| Mainly mechanical       | 40           | 20            | 40           |                           |
| reduction of primary    |              |               | 20           |                           |
| porosity               |              |               |              |                           |
| Semi-mature stage:     |              |               |              |                           |
| Mainly chemical         | 40           | 20            | 40           |                           |
| reduction of primary    |              |               | 20           |                           |
| porosity               |              |               |              |                           |
| Mature stage:          |              |               |              |                           |
| Primary porosity at     | 0            | 0             | 0            |                           |
| irreducible levels,     |              |               |              |                           |
| Secondary porosity may  | 0            | 0             | 0            |                           |
| exist                   |              |               |              |                           |
| Super mature stage:    |              |               |              |                           |
| Primary and secondary   | 0            | 0             | 0            |                           |
| porosity at irreducible |              |               |              |                           |

Fig. 1 Textural stages of mesodiagenesis of sandstone porosity and the petrographer’s dilemma of secondary porosity (after Schmidt and McDonald 1979a, b; Giles 1987)
is still prominent with regard to the origin of anomalously high porosity in the deeply buried sandstones in China (Bai et al. 2013; Si and Zhang 2008; Tang et al. 2013; Wang et al. 2013; Yuan et al. 2012; Zhu and Zhang 2009; Zhu et al. 2007).

According to laboratory water–rock interaction experiments, carbonate minerals can be dissolved more easily and to be dissolved much faster than feldspar minerals in open geochemical systems under steady-state conditions far from equilibrium (Bertier et al. 2006; Chen et al. 2008; Liu et al. 2012; Weibel et al. 2011; Yang et al. 1995), and the carbonate minerals seem likely to be the most important minerals for the development of secondary pores in buried sandstones (Giles and Marshall 1986; Schmidt and McDonald 1979a). However, things may be different in closed subsurface sandstone geochemical systems. Based on our studies, we identified four types of selective dissolution assemblages of feldspar and carbonate minerals in different sandstone rocks (Fig. 2), which may have some significant implications for this debate (Fig. 1) (Yuan et al. 2015a, b, c). At the same time, some recent papers presented the dissolution of silicate minerals with no dissolution of carbonate minerals in the Kimmeridge Clay mudstones (Macquaker et al. 2014) and in the Eocene sandstones in the Bohai Bay Basin (Yuan et al. 2015a, b, c). Also, Turchyn and DePaolo (2011) suggested that the dissolution of carbonate minerals in mudstones can be significantly suppressed by the presence of silicate minerals, and the dissolution rate is much smaller even when compared with the already-slow rates typical of carbonate-rich sediments (Turchyn and DePaolo 2011).

Stimulated by these recent reviews and the selective dissolution phenomena of feldspars and carbonate minerals in buried subsurface sandstones, the objectives of this article are to: (1) provide detailed petrographic evidence of selective dissolution assemblages of feldspars and carbonate minerals in buried sandstones; (2) discuss the significance of burial carbonate dissolution in buried sandstones with the constraints of porosity-depth data, water–rock experiments, and geochemical calculations; and (3) review the literature on the dissolution of carbonate minerals in buried sandstones with petrographic and geochemical constraints.

2 Evidence from the reservoirs

2.1 Petrography

Feldspar grains and carbonate cements are common minerals in subsurface sandstones. As both the feldspar and carbonate minerals can be dissolved by the acids (e.g., CO₂ and organic acids) originating from thermal maturation of organic matter, the potential to generate secondary pores in sandstone reservoirs through the dissolution of the feldspar and carbonate mineral has been discussed a great deal. Petrographic evidence has been used to demonstrate the presence of secondary porosity in sandstones (Bjørlykke and Jahren 2012; Giles and Marshall 1986). The porosity related to framework grain dissolution (e.g., feldspars) can be recognized and statistically quantified (Taylor et al. 2010). Though extensive burial dissolution of carbonate cements has been suggested by many researchers (Schmidt and McDonald 1979a; Surdam et al. 1989; Surdam and Boese 1984), intergranular pores without carbonate cements should not be interpreted as secondary porosity unless considerable petrographic evidences of its former presence can be established (Taylor et al. 2010). Experiments under steady-state conditions far from equilibrium illustrate that the dissolution rates of carbonate minerals are much faster than the rate of feldspars. In the natural sandstone rocks, however, things are likely to be more complex. Based on our studies, we identified four types of typical selective dissolution assemblages of feldspar and carbonate minerals in sandstone rocks.

(1) Type I: Little feldspar dissolution vs. extensive carbonate precipitation

In buried sandstones and sandstone outcrops, carbonate-cemented concretions are very common (Dos Anjos et al. 2000; Dutton 2008; Gluyas and Coleman 1992; Saigal and Bjørlykke 1987; Wang et al. 2016; Yuan et al. 2015a, b, c). Petrography and relevant stable isotope data usually suggest that the carbonate cements in such concretions formed soon after sediment deposition and prior to the occurrence of the key dissolution period in the rocks (Dutton 2008; Gluyas and Coleman 1992). In such concretions, large amounts of carbonate cements precipitated and preserved both the depositional fabric and the composition of the sand grains with little if any grain replacement. The early carbonate cements occupied almost all primary intergranular pores (Fig. 2a, b) and formed flow barriers (Saigal and Bjørlykke 1987), which led to little dissolution of both the feldspars and the carbonate cements in such concretions during the later burial (Fig. 2a, b). In buried sandstones, the development of such concretions usually occurs near the sandstone–mudstone interface and the thickness of these concretions ranges from centimeters to several meters (Dutton 2008; Gluyas and Coleman 1992; Mcbride and Milliken 2006).

(2) Type II: Little feldspar dissolution vs. extensive carbonate dissolution

In buried sandstones, the phenomenon of little feldspar dissolution versus extensive carbonate dissolution is rare and few publications have ever reported on it. However, one paper reported on the extensive dissolution of early calcite cements (Fig. 2c, c’) in Quaternary beach deposits.
by meteoric water during periods of falling of sea levels (Cavazza et al. 2009). The microphotograph suggests little dissolution of the associated silicate minerals (Fig. 2c). The associated silicate minerals were dissolved much less extensively than the calcite cements, probably due to the short geological time period and the low temperature in the meteoric diagenetic environment (Cavazza et al. 2009). These observations are consistent with the laboratory experiments under steady-state conditions far from equilibrium in which calcite can be dissolved more easily than silicate minerals (Chen et al. 2008; Liu et al. 2012; Weibel et al. 2011).
(3) Type III: Extensive feldspar dissolution and extensive carbonate dissolution

In buried sandstones, the dissolution of feldspar and carbonate minerals has been suggested as common occurrence by many authors (Schmidt and McDonald 1979a, 1979b; Surdam et al. 1989; Surdam and Boese 1984). Little convincing petrographic evidence, however, has been reported to support the coexistence of extensive feldspar dissolution and extensive carbonate dissolution in buried sandstones. One typical example was provided by Taylor (1990, 1996), who presented a striking and convincing microphotograph to show the dissolution of carbonate cements and detrital carbonate grains at Picaroon field (Fig. 2f) (Taylor 1990, 1996; Taylor et al. 2010). In the microphotographs, we can also identify the dissolution of feldspar grains (Fig. 2f). Another example we have identified is the Es3 sandstones from Well Yan 16 in the Mingfeng area, Dongying Sag. In the thin sections from Well Yan16, we observed the typical dissolution of feldspar grains and ferroan calcite cements in the sandstones of the middle Es3 Formation (Fig. 2d, e). These sandstones are located close to some faults, which connect to the unconformity that developed at the end of the Eocene period. In these thin sections, the remnants of ferroan calcite cements were irregular and developed dissolved pores (Fig. 2d, e). The low-oxygen isotope data (−15.02‰carbonate cements and the maximum depth (1920 m–1960 m with temperatures of 75–80 °C) suggest that the fluid that formed these carbonate cements had negative δ18O data (lower than −8‰carbonate dissolution) (Matthews and Katz 1977), which support massive meteoric water flux in these sandstones (Fayek et al. 2001; Harwood et al. 2013).

(4) Type IV: Extensive feldspar dissolution vs. little carbonate dissolution

Macquaker et al. (2014) reported the fabric observation of kaolinite precipitation (byproduct of the dissolution of aluminosilicate minerals) and no dissolution of the associated calcareous textures (Fig. 2f) in the Kimmeridge Clay Formation mudstones and regarded the phenomenon as surprising and significant (Macquaker et al. 2014). In both mudstones and sandstones, such phenomena have not yet received much attention, although they were mentioned in some publications (Armitage et al. 2010; Baker et al. 2000; Ceriani et al. 2002; Dos Anjos et al. 2000; Dutton and Land 1988; Fisher and Land 1987; Girard et al. 2002; Hendry et al. 1996; Mil-liken et al. 1994; Salem et al. 2000; Tobin et al. 2010). Using thin sections and scanning electron microscopy (SEM) from samples from the northern steep slope zone of the Dongying Sag, we identified the phenomena of typical extensive dissolution of feldspar grains with no/little dissolution of carbonate cements and detrital carbonate grains in the lower Es3 Formation and the Es4 Formation (Fig. 2g–k). The carbonate cements occurred as connected patches (Fig. 2a), single crystals (Fig. 2j, k) or grain-coating carbonate (Fig. 2g, h), and individual crystals commonly exhibited euhehedral crystals faces abutting open pore space (Fig. 2k). The euhehedral ankerite engulfed by the stage-II quartz overgrowths (Fig. 2j) suggests that the carbonate minerals were not leached when the stage-II feldspar dissolution and quartz cementation occurred in the acidic geochemical system. In addition, the detrital carbonate grains and grain-coating carbonate cements show no evidence of dissolution (Fig. 2g–i); moreover, carbonate overgrowths are often found accompanying the detrital carbonate grains. However, the feldspar grains engulfed by early grain-coating carbonate cements or close to detrital carbonate grains are dissolved extensively (Fig. 2g–i).

Overall, petrography textures suggest that carbonate mineral dissolution is not likely to occur all the time. Only in two cases, extensive carbonate dissolution in the sandstone reservoirs is likely to occur.

### 2.2 Porosity-depth data

The porosity evolution model proposed by Schmidt and McDonald (Fig. 1a) was initially accepted and embraced by many geologists (Bjørlykke and Jahren 2012; Boggs 2011; Burley 1986; Dutton and Willis 1998; Higgs et al. 2010; Khidir and Catuneanu 2010; Schmidt and McDonald 1979a; Shanmugam 1984; Wilkinson et al. 1997) to explain the fairly common occurrence of intergranular porosity in sandstone buried to significant depth. However, as a general rule, global porosity-depth data show a steady decrease in the sandstone P50, P10, and the maximum porosity trends as the depth increases (Fig. 3) (Ehrenberg et al. 2009; Ehrenberg and Nadeau 2005), which is inconsistent with the porosity evolution model proposed by Schmidt and McDonald in 1979 (Fig. 1a).

Although anomalously high porosities do exist in some deeply buried sandstones (Bloch et al. 2002; Warren and Pulham 2001), studies on the origin of the anomalously high porosities suggest that the dissolution of grains or preexisting cements are just one subordinate aspect of this porosity. These anomalously high porosities can be attributed to early emplacement of hydrocarbons (Bloch et al. 2002; Gluyas et al. 1993; Wilkinson and Haszeldine 2011), fluid overpressure, or grain coats and grain rims (Bahls and De Ros 2013; Bloch et al. 2002; Ehrenberg 1993); the mixture of porosity of rocks with different lithology from shallow to deep depths may also lead to the occurrence of anomalously high porosities in a porosity-depth profile (Bjørlykke 2014; Bjørlykke and Jahren 2012).
The Eocene sandstones in the northern steep slope zone in the Dongying Sag are an example exhibiting the impact of fluid overpressure, hydrocarbon emplacement, and mineral dissolution. Detailed geological settings are available in some papers (Cao et al. 2013; Guo et al. 2010, 2012; Yuan et al. 2013a, b). Subaqueous fans and lacustrine fans were deposited in the Eocene Es4–Es3 Formations in the northern steep slope zone together with contemporary organic-rich mudstones and shales. The development of anomalously high porosities in the reservoirs has been reported (Cao et al. 2014). In this paper, two types of porosity-depth profiles were plotted and presented, using the 7936 core porosity data collected from the Shengli Oilfield Company. The lithology and oil-bearing properties of these samples were analyzed with core-logging materials. The fluid pressure relevant to these samples was analyzed using the equivalent depth method (Gao et al. 2008) using acoustic logging data with the constraint of the measured formation fluid pressure. And a database of the reservoir properties was established using the information of the porosity, depth, lithology, oil-bearing properties, and fluid pressure data. Type-A porosity-depth profiles of the combined lithology (Fig. 4a1) show that anomalously high porosities do exist at the depth intervals of 2.8–3.7 km and 3.9–4.4 km, and the porosity-depth profiles of each individual lithology also show the existence of anomalously high porosities in some specific depth intervals (Fig. 4a2–a7). However, the Type-B porosity-depth profiles (Fig. 4b1–b7) show no existence of the anomalously high porosities when the impact of the fluid overpressure and hydrocarbon emplacement on the reservoir porosity was removed. This analysis suggests that even where anomalously high porosities exist in deeply buried reservoirs, significant dissolution of carbonate cements may not be the cause. This is consistent with the petrographic evidence of selective dissolution of feldspar in the presence of carbonate minerals and the precipitation of authigenic clays and quartz cements following the feldspar dissolution in these rocks (Yuan et al. 2013a, b).

3 Water–rock experiments

3.1 Samples and methods

Pure calcite crystals were crushed, and the calcite grains with a size of 2–4 mm were selected. In each experiment, one grain with a polished surface was employed to investigate the dissolution features after the experiments. The grains were ultrasonically cleaned with analytical-grade distilled water three times to remove submicron-to-micron-sized particles adhering to the grains. The calcite grains were dried at 60 °C for 12 h and examined with a Coxem-EM-30 plus scanning electron microscope (SEM) to check the total removal of the small particles. Calcite grain samples were prepared using a high-precision electronic balance (error < 0.005 g). High salinity waters with different salinity were prepared with 99.99% NaCl, 99.99% CaCl2, and deionized water (DW). Glacial acetic acid with a purity of more than 99.5% was used to prepare acidic water with different pH.

The detailed experiment conditions are listed in Table 1. The calcite dissolution experiments at different temperatures (20 °C, 90 °C) were conducted in Hastelloy Reactors. For experiments with participation of CO2, CO2 gas with a purity of more than 99.995% was injected into the reactor by pumping to reach the designed pCO2 of 50 bar. The experiments were conducted for 3, 8, and 15 days, respectively. After the experiments at 20 °C, the calcite grains were separated from the water quickly, while for the experiments at 90 °C, the reactor was firstly cooled to approximately 20 °C using cold water in less than 1 h, and then, the calcite grains were separated from the water. The water pH was tested after the separation of the water from minerals. The reacted
calcite minerals were cleaned in DW three times to remove possible salt precipitated on the mineral surfaces. And the reacted calcite minerals were weighed after being dried at 60 °C for 12 h.

**3.2 Experimental results and geological implication**

The weight loss and relevant volume changes of the calcite minerals are presented in Table 1. The experiments A1–A3 demonstrate that low pH water with acetic acid (pH = 3.93–3.98) can dissolve calcite at 20 °C. As the water salinity and the Ca²⁺ concentration in water increase, the dissolution capacity of the acidic water decreases dramatically. Even with a high water/rock volume ratio (45:1), the ratio between the mass loss after dissolution and the primary weight of the calcite mineral prior to the experiments suggests that only a small amount of calcite was dissolved, and this can only have resulted in a few secondary pores in the calcite grains (less than 1%); even the dissolved calcite was not re-precipitated (Fig. 5). The results of the experiments D1–D7 at 90 °C show a similar trend.

The results of the experiments B1–B3 show that deionized water and saline water with a partial pressure of CO₂ \( (P_{CO₂}) \) at 50 bar can dissolve calcite at 20 °C. A comparison of the results of the experiments B2 and B3 shows a decrease in the corrosion ability of the acidic water as the salinity increases. A comparison between the results of the experiments B1 and B3 shows that the calcite–CO₂ interactions reached dynamic equilibrium in 8 days (maybe in an even shorter time) after the dissolution of 0.212 g calcite and a longer (15 days) exposure of calcite to the CO₂-charged water did not result in more dissolution. This result indicates
Table 1  Data of calcite-dissolving experiments at low and high temperatures

| Expt No. | Before interaction | After interaction |  |  |  |  |  |  |
|----------|--------------------|-------------------|---|---|---|---|---|---|
|          | Composition of solution | Water volume, mL | pH | Water salinity, g/L | Ca\(^{2+}\) concentration, g/L | Calcite weight, g | p\(_{CO_2}\), bar | Weight loss of calcite, g | Volume change of calcite, % | Water/rock volume ratio | T, °C | Time, day |
| A1       | DW + HAC           | 500               | 3.97 | 0 | 0 | 30.060 | N/A | 6.29 | 29.718 | 0.283 | 0.943 | 45 | 20 | 8 |
| A2       | DW + HAC + NaCl + CaCl\(_2\) | 500 | 3.93 | 20 | 2 | 30.012 | N/A | 7.11 | 29.856 | 0.156 | 0.520 | 45 | 20 | 8 |
| A3       | DW + HAC + NaCl + CaCl\(_2\) | 500 | 3.98 | 80 | 5 | 30.006 | N/A | 6.82 | 29.952 | 0.054 | 0.180 | 45 | 20 | 8 |
| B1       | DW + NaCl + CaCl\(_2\) | 500 | 8.51 | 80 | 5 | 30.003 | 50 | 6.05 | 29.791 | 0.212 | 0.700 | 45 | 20 | 8 |
| B2       | DW                | 500 | 7.16 | 0 | 0 | 15.007 | 50 | 6.92 | 14.698 | 0.309 | 2.059 | 90 | 20 | 15 |
| B3       | DW + NaCl + CaCl\(_2\) | 500 | 8.72 | 80 | 5 | 15.000 | 50 | 6.33 | 14.785 | 0.215 | 1.433 | 90 | 20 | 15 |
| C1       | DW + HAC + NaCl + CaCl\(_2\) | 500 | 3.97 | 80 | 5 | 29.955 | 50 | 6.41 | 29.259 | 0.696 | 2.323 | 45 | 20 | 8 |
| D1       | DW + HAC           | 350 | 3.43 | 0 | 0 | 14.897 | N/A | 6.66 | 14.568 | 0.329 | 2.208 | 63 | 90 | 3 |
| D2       | DW + HAC + NaCl + CaCl\(_2\) | 350 | 3.44 | 20 | 2 | 14.512 | N/A | 6.56 | 14.317 | 0.141 | 0.972 | 65 | 90 | 3 |
| D3       | DW + HAC + NaCl + CaCl\(_2\) | 350 | 3.46 | 40 | 4 | 14.844 | N/A | 6.5 | 14.630 | 0.214 | 1.442 | 63 | 90 | 3 |
| D7       | DW + HAC + NaCl + CaCl\(_2\) | 350 | 3.46 | 80 | 5 | 15.065 | N/A | 7.02 | 15.009 | 0.056 | 0.372 | 63 | 90 | 3 |
| D5       | DW + HAC + NaCl + CaCl\(_2\) | 350 | 3.46 | 150 | 7.5 | 15.190 | N/A | 6.13 | 15.074 | 0.116 | 0.764 | 63 | 90 | 3 |
| D6       | DW + HAC + NaCl + CaCl\(_2\) | 350 | 3.47 | 200 | 10 | 15.129 | N/A | 6.4 | 15.090 | 0.039 | 0.258 | 63 | 90 | 3 |
| D7       | DW + HAC + NaCl + CaCl\(_2\) | 350 | 3.47 | 300 | 15 | 14.948 | N/A | 5.28 | 14.905 | 0.043 | 0.288 | 63 | 90 | 3 |

\(DW\) distilled water, \(HAC\) acetic acid, \(N/A\) not applicable
that in a relative closed geochemical system with a fixed $p_{CO_2}$, the available water volume dominates the dissolution volume of the calcite, even if CO$_2$ is available in sufficient quantities. Also, with a high water/rock volume ratio (45:1 or 90:1), only a small amount of calcite (less than 2%) was dissolved by the CO$_2$-rich water.

A comparison of the results of the experiments C1, A3, and B1 shows that the coexistence of acetic acid and CO$_2$ in saline water promotes more calcite dissolution than with only acetic acid or CO$_2$ in the saline water. However, no more than 2.5% of the calcite was dissolved in the C1 experiments. Overall, the experiments with a high water/rock volume ratio, low pH, and sufficient CO$_2$ resulted in the dissolution of only a small amount of calcite.

As low temperature, low pH, high $p_{CO_2}$, and high water/rock ratio cannot generate a large volume of secondary pores by the dissolution of carbonate minerals, it is not likely that extensive carbonate dissolution will occur in buried sandstone geochemical systems with high temperature and low water/rock ratio. Many studies on water–rock interaction experiments also support this idea when the data were analyzed quantitatively, although dissolution does take place at low/high temperatures (Weibel et al. 2014). In addition, the dissolved carbonate minerals were commonly reported to be re-precipitated in long-term numerical simulation experiments (Bertier et al. 2006; Liu et al. 2012). As the initial pH values (<4) of the waters used in the experiments were much lower than those of most formation waters and the water/rock ratios were much higher than those in subsurface rocks (Birkle et al. 2009; Birkle et al. 2002; Egeberg and Aagaard 1989; Frape et al. 1984; Surdam et al. 1985), we conclude that the calcite dissolution in deeply buried sandstones without a favored pathway (e.g., faults) is likely to be weaker than in the experiments.

4 Aggressive fluids and mass transfer in sediments

4.1 Pyrolysis experiments of kerogen

Hydrous and anhydrous pyrolysis experiments with pure kerogen or source rocks have been used to investigate the maturation of organic matter in source rocks with respect to the generation of organic acids and CO$_2$ (Barth et al. 1988; Barth et al. 1996; Barth and Bjørlykke 1993). Using worldwide source rocks and different types of kerogens with various total organic carbon (TOC) contents and different maturities, more than 110 pyrolysis experiments have been conducted in the last 40 years to analyze the yield of organic
acids and CO₂ during kerogen maturation (Table 2) (Barth et al. 1988; Barth et al. 1996; Barth and Bjørlykke 1993; Chen et al. 1994; Kawamura et al. 1986; Kawamura and Kaplan 1987; Meng et al. 2008; Zeng et al. 2007; Zhang et al. 2009). The results of the experiments demonstrate that the maximum yield of acetic acids and total organic acids (TOA) is 0.685 × 10⁻³ mol/g TOC and 1.34 × 10⁻³ mol/g TOC, respectively. The experiments with the acetic acids yield more than 0.5 × 10⁻³ mol/g TOC account for only approximately 5% of the total experiments, and the experiments with the TOA yield more than 0.6 × 10⁻³ mol/g TOC account for 10% of the total experiments. In the pyrolysis experiments, CO₂ has a yield equivalent to (0.30–10.9) × 10⁻³ mol/g TOC, which is higher than that of the organic acids. Commonly, high TOC and high maturation result in low yields of organic acids and CO₂ of one unit kerogen.

4.2 Acids in pore water

The dissolution of feldspar grains is a natural consequence of water–rock interactions under conditions of increasing burial depth and temperatures (Giles and De Boer 1990; Taylor et al. 2010). Although organic acids and CO₂ were commonly suggested as the cause of feldspar dissolution (Giles and De Boer 1989; Schmidt and McDonald 1979a; Surdam et al. 1989; Surdam and Boese 1984), Giles and De Boer (1990) suggested that no unusual or special source of acidic pore fluids is required for this dissolution process (Giles and De Boer 1990). To dissolve carbonate minerals characterized by retrograde solubility (Giles and De Boer 1989), however, there must be a supply of a large amount of acidic water that has the capacity to provide H⁺.

(1) Various organic acids from kerogens are present in most of the formation waters in petroliferous basins. Acetic acid with a relative content of approximately 80% dominates the organic acids in most cases (Surdam and Crossey 1987; Surdam et al. 1989; Surdam and Boese 1984). It was suggested by Surdam et al. (1984, 1987, 1989) and Meshri (1986) that organic acids were more aggressive than CO₂ and could be responsible for the dissolution of silicate and carbonate minerals (Meshri 1986; Surdam and Crossey 1987; Surdam et al. 1989; Surdam and Boese 1984). The leaching of calcite by acetic acid can be expressed as CaCO₃ + CH₃COOH —> Ca²⁺ + HCO₃⁻ + CH₃COO⁻. Using the data of the concentration of organic acid in oilfield waters, Surdam (1984, 1987) further suggested that large volumes of water-soluble organic acids are generated during the thermocatalytic degradation of kerogen in the range of 80–120 °C and the concentration of organic acids can even reach up to 10000 ppm (Surdam et al. 1989; Surdam and Crossey 1987; Surdam and Boese 1984).

The concentrations of organic acids are lower when the temperatures are below 80 °C or above 120 °C due to the bacterial destruction and thermal destruction of the short-chained organic acids, respectively (Surdam et al. 1989; Surdam and Crossey 1987). The concentration data of the organic acids in the formation waters from global petroleum sandstone reservoirs show that more than 90% of the pore waters contain organic acids at concentrations less than 3000 mg/L (Fig. 6a) (Cai et al. 1997; Fisher 1987; Kharaka 1986; MacGowan and Surdam 1988; Meng et al. 2006; Meng et al. 2011; Surdam et al. 1989; Surdam and Crossey 1987; Wang et al. 1995, 2007; Xiao et al. 2005). In the petroliferous basins in China, the concentrations of organic acids in the formation waters are usually less than 2500 mg/L (Fig. 6b). With high geothermal gradients (around 35 °C/1 km), the highest concentrations of organic acids developed at the depth of 1500–3500 m in the basins in East China; in contrast, the highest concentrations developed at the depth of 4500–6000 m in the basins in West China with low geothermal gradients (approximately 20 °C/1 km) (Fig. 6b) (Cai et al. 1997; Fisher 1987; Kharaka 1986; MacGowan and Surdam 1988; Meng et al. 2006; Meng et al. 2011; Surdam et al. 1989; Surdam and Crossey 1987; Wang et al. 1995, 2007; Xiao et al. 2005).

In rocks with a high mudstone/sandstone ratio (e.g., 10:1), about 60 mol of acetic acids can be produced in 1 m³ source rocks if an average TOC of 5% in the mudstone and an organic acid yield of 0.5 × 10⁻³ mol/g TOC (Table 2) are available in the source rocks. Because organic acids concentrate at temperatures of 80–120 °C, most organic acids are assumed to be released from the source rocks to the reservoirs in the depth interval of 1500–4000 m. From 1500–4000 m, the sandstone porosities generally decrease from 80% to 15% and the mudstone porosities decrease from 20% to 5% (Gluyas and Cade 1997; Pittman and Larese 1991; Ramm 1992). As organic acids are water soluble (Barth and Bjørlykke 1993), we assume that all the pore water expelled from the mudstones to the sandstone reservoirs have a high concentration of organic acids (10,000 ppm acetic acid). In this case, the organic acids expelled to the reservoirs can dissolve only 0.46% volume of calcite with a thorough consumption of the available acids. In another case, if diffusion or hydrocarbon migration can transport more organic acids to the sandstone reservoirs (Barth and Bjørlykke 1993; Thyne 2001), only 2% volume of calcite can be dissolved in the sandstone reservoirs. The organic acids are weak acids and the equilibrium constant of the calcite-leaching reaction by organic acids decreases from 8.5 × 10⁻⁴ at 25 °C to 7.9 × 10⁻⁵ at 100 °C (Giles and Marshall 1986). Under constraints of the equilibrium constant, the calcite...
Table 2  Pyrolysis experiment data of global source rocks and kerogen. (data from Kawamura et al. 1986; Kawamura and Kaplan 1987; Barth et al. 1988; Barth and Bjørlykke 1993; Chen et al. 1994; Barth et al. 1996; Zeng et al. 2007; Meng et al. 2008; Zhang et al. 2009)

| Sample location                  | Sample type     | Sample amount | Kerogen type | TOC, % | $R_o$, % | CO$_2$, 10$^{-3}$mol/ gTOC | Acetic acid, 10$^{-3}$mol/ gTOC | Total organic acids, 10$^{-3}$mol/ gTOC | Publications       |
|----------------------------------|-----------------|---------------|--------------|--------|---------|---------------------------|-------------------------------|------------------------|-------------------|
| Well C11 in Huanghua Depression  | Mudstone        | 13            | II-1         | 2.46   | 0.42    | –                          | –                             | 0.010–0.081            | Meng et al. (2008) |
| $E_{51}$ Formation in Dongying Sag | Mudstone        | 1             | –            | 1.04   | 0.33    | –                          | –                             | 0.175–0.608            | Zeng et al. (2007) |
| $E_{54}$ Formation in Dongying Sag | Mudstone        | 1             | –            | 2.11   | 0.40    | –                          | –                             | 0.084–0.220            |                   |
| $E_{54}$ Formation in Dongying Sag | Mudstone        | 1             | –            | 3.70   | 0.42    | –                          | –                             | 0.086–0.128            |                   |
| $E_{54}$ Formation in Dongying Sag | Mudstone        | 1             | –            | 1.73   | 0.24    | –                          | –                             | 0.160–0.445            |                   |
| Well Chun11 in Dongying Sag       | Mudstone        | 5             | I            | 3.50   | 0.32    | –                          | –                             | 0.134–0.330            | Zhang et al. (2009) |
| Well Cao 13-15 in Dongying Sag     | Mudstone        | 3             | II-1         | 2.29   | 0.32    | –                          | –                             | 0.024–0.177            |                   |
| Well Ying 10 in Dongying Sag       | Mudstone        | 3             | II-2         | 1.19   | 0.48    | –                          | –                             | 0.051–0.341            |                   |
| Well YMian4-5-165 in Dongying Sag  | Mudstone        | 1             | I            | 1.32   | 0.36    | –                          | –                             | 0.676                  |                   |
| Well Lunnan 54 in Tarim Basin      | Mudstone        | 4             | II-III       | 8.04   | 0.61    | –                          | 0.002–0.005                  | 0.010–0.04             | Chen et al. (1994) |
| Well Lunnan 54 in Tarim Basin      | Mudstone        | 12            | II-III       | 8.04   | 0.61    | –                          | 0.004–0.084                  | 0.011–0.110            |                   |
| Well Tan26 in Jianghan Basin       | Mudstone        | 1             | II-III       | –      | 0.41    | –                          | 0.612                         | 0.700                  |                   |
| Green River Shale                  | Kerogen         | 7             | I            | 2.30   | –       | –                          | 0.007–0.036                  | 0.010–0.048            | Kawamura et al. (1986) |
| Monterey Formation                | Kerogen         | 2             | II           | 10.0   | –       | –                          | 0.015–0.035                  | 0.023–0.0060           |                   |
| Monterey Formation                | Kerogen         | 1             | II           | –      | –       | –                          | 0.025                         | 0.036                  | Kawamura and Kaplan (1987) |
| Green River Shale                  | Kerogen         | 1             | I            | –      | –       | –                          | 0.04                          | 0.056                  |                   |
| Tanner Basin                       | Kerogen         | 1             | II           | –      | Immature | –                          | 0.149                         | 0.278                  |                   |
| Sierra Bog sediments               | Humic acid      | 1             | III          | –      | –       | –                          | 0.14                          | 0.249                  |                   |
Table 2 (continued)

| Sample location | Sample type         | Sample amount | Kerogen type | TOC, % | \( R_m \), % | \( \text{CO}_2 \), \( 10^{-3} \text{mol/gTOC} \) | Acetic acid, \( 10^{-3} \text{mol/gTOC} \) | Total organic acids, \( 10^{-3} \text{mol/gTOC} \) | Publications                  |
|-----------------|---------------------|---------------|--------------|--------|--------------|--------------------------------|---------------------------|---------------------------------|-----------------------------|
| Kimmeridge oil shale, Dorset, Upper Jurassic | Oil shale          | 5             | –            | 12.6   | Immature     | –                            | 0.057–0.215                 | 0.104–0.345                    | Barth et al. (1988)            |
| Jurassic, the Norwegian continental shelf | Coaly shale        | 3             | –            | 14.3   | Mature       | –                            | 0.009–0.013                 | 0.011–0.016                    |                              |
| Lower Jurassic, the Norwegian continental shelf | Coal               | 3             | –            | 39.6   | Immature     | –                            | 0.069–0.100                 | 0.081–0.123                    |                              |
| Upper Jurassic, the Norwegian continental shelf | Mudstone           | 3             | –            | 5.03   | Immature     | –                            | 0.151–0.284                 | 0.263–0.412                    |                              |
| Kimmeridge outcrop, Dorset, UK | Mudstone           | 3             | II           | 12.60  | Immature     | 2.86–8.73                    | 0.141–0.231                 | 0.252–0.346                    | Barth and Bjørlykke (1993)     |
| Kimmeridge, North Sea | Coal               | 3             | II           | 5.03   | Immature     | 7.95–10.93                   | 0.154–0.282                 | 0.262–0.414                    |                              |
| Kimmeridge outcrop, Dorset, UK | Mudstone           | 3             | II           | 51.30  | 0.29         | 1.65–4.70                    | 0.052–0.116                 | 0.110–0.244                    |                              |
| Heather, North Sea | Mudstone           | 3             | II           | 6.49   | 0.40         | 1.23–2.16                    | 0.142–0.273                 | 0.177–0.341                    |                              |
| The Norwegian continental shelf | Mudstone           | 3             | III          | 14.30  | Mature       | –                            | 0.009–0.013                 | 0.011–0.017                    |                              |
| The Norwegian continental shelf | Coaly shale        | 3             | Coal         | 39.60  | Immature     | 3.18–4.72                    | 0.069–0.100                 | 0.081–0.124                    |                              |
| The Norwegian continental shelf | Coal               | 3             | Coal         | 23.10  | 0.38         | 0.30–1.99                    | 0.124–0.276                 | 0.160–0.377                    |                              |
| Western Germany | Coal                | 3             | Coal         | 70     | 0.26         | 1.72–2.41                    | 0.518–0.552                 | 0.609–0.705                    |                              |
Table 2 (continued)

| Sample location          | Sample type                      | Sample amount | Kerogen type | TOC, % | $R_o$, % | $\text{CO}_2$, $10^{-3}\text{mol/gTOC}$ | Acetic acid, $10^{-3}\text{mol/gTOC}$ | Total organic acids, $10^{-3}\text{mol/gTOC}$ | Publications |
|--------------------------|----------------------------------|---------------|--------------|--------|---------|---------------------------------------|--------------------------------------|-------------------------------------------|--------------|
| Draupne, the Norwegian continental shelf | Dicarbonated mudstone | 6             | –            | 3.70–7.19 | –       | 1.00–2.36 | 0.048–0.455 | 0.081–0.659 | Barth et al. (1996) |
| Draupne, the Norwegian continental shelf | Mudstone | 3             | –            | 3.52–6.21 | –       | 1.58–8.31 | 0.082–0.315 | 0.181–0.497 |
| Heather, the Norwegian continental shelf | Dicarbonated mudstone | 4             | –            | 1.06–5.49 | –       | 0.77–6.61 | 0.038–0.399 | 0.077–0.601 |
| Heather, the Norwegian continental shelf | Mudstone | 2             | –            | 1.85–7.79 | –       | 3.45–3.78 | 0.086–0.125 | 0.148–0.218 |
| Brent, the Norwegian continental shelf | Dicarbonated mudstone | 2             | –            | 3.46–5.25 | –       | 0.38–1.53 | 0.085–0.103 | 0.144–0.151 |
| Dulin, the Norwegian continental shelf | Dicarbonated mudstone | 2             | –            | 1.32–3.13 | –       | 0.35–10.00 | 0.008–0.695 | 0.105–1.349 |

— Not measured

**Fig. 6** Concentrations of organic acids in the pore water of sandstone reservoirs in oil and gas basins. **a** Organic acids in global sedimentary basins; **b** organic acids in sedimentary basins in China. (data from Kharaka 1986; Fisher 1987; Surdam and Crossey 1987; MacGowan and Surdam 1988; Surdam et al. 1989; Wang et al. 1995; Cai et al. 1997; Xiao et al. 2005; Meng et al. 2006; Wang et al. 2007; Meng et al. 2011)
volume that can be dissolved in the reservoirs is reduced significantly. In addition, most source rocks contain considerable carbonate minerals and silicate minerals (Ehrenberg et al. 2012; Giles and Marshall 1986; Taylor et al. 2010); these minerals first consume some of the organic acids generated in the mudstones, which also decreases the volume of acids expelled to the reservoirs and the leaching ability of the organic acids in the reservoirs (Barth et al. 1996; Barth and Bjørlykke 1993).

(2) CO₂ is present in most oil-gas sandstone reservoirs, though most natural gas accumulations contain less than 10% CO₂ (Seewald 2003). It was suggested by Smith and Ehrenberg (1989), Ribstein et al. (1998), and Seewald (2003) that the CO₂ content in natural gases generally increases with increasing temperature and burial depth, and the \( p_{\text{CO}_2} \) increases systematically in the temperature range from 40 to 200 °C (Fig. 7) (Curtis 1978; Ribstein et al. 1998; Schmidt and McDonald 1979a; Seewald 2003; Smith and Ehrenberg 1989). The CO₂ in the reservoirs originates from the degradation of organic matter or from water–rock interactions (Curtis 1978; Ribstein et al. 1998; Schmidt and McDonald 1979a; Seewald 2003; Smith and Ehrenberg 1989).

In the range of 80–120 °C, the release of CO₂ from kerogen in the source rocks is inevitably one important source and it was suggested by Schmidt (1979a, b), Surdam (1984), and Surdam and Crossey 1987) that this CO₂ source is one of the most important carbonic acids for carbonate dissolution (Schmidt and McDonald 1979a; Surdam and Crossey 1987; Surdam and Boese 1984). Because of the constraints of the mass balance calculation, Lundegard et al. (1984) suggested that even if all the CO₂ generated from kerogen was expelled from the source rocks to the sandstones, only 1%–2% of secondary porosity could be generated (Lundegard et al. 1984).

In contrast to Schmidt (1979a, b) and Surdam (1984), Smith and Ehrenberg (1989) proposed that the increased CO₂ abundance results in precipitation rather than dissolution of carbonate minerals at the depth interval with temperature ranging from 80°C to 120 °C, in which the organic acids have the highest concentrations and control the alkalinity of the carbonate–silicate–organic acid–carbonic acid–\( p_{\text{CO}_2} \) system.

Using numerical simulations with the constraints of thermodynamics, Huang et al. (2009) calculated the pH values of different carbonate–H₂O–CO₂ geochemical systems in the equilibrium state (Fig. 8a) and the dissolution/precipitation volumes of the calcite or dolomite minerals in these systems at temperatures of 28–235 °C, pressure of 1–70 MPa, depth of 1–7 km, and a specific molar content of CO₂ (Fig. 8b–d). The results show that the systems with a higher CO₂ content have lower pH values and this results in the dissolution of more carbonate minerals at depths shallower than 2000 m. At depths deeper than 2000 m, however, more carbonate minerals are precipitated in the systems with more CO₂, even if the systems have lower pH values of approximately 4.8 (Fig. 8) (Huang et al. 2009). Using laboratory water–rock interaction experiments, Song and Huang (1990) also demonstrated that calcite can be precipitated even when the pH is lower than 5 (Song and Huang 1990).

As carbonate minerals are characterized by retrograde solubility, cooling of hot fluids have been suggested to dissolve carbonate minerals during the uplift stage of the formation or during injection of deep hot water to shallow formations. Using numerical simulations with the constraints of thermodynamics, Yuan et al (2015a, b, c) modeled the calcite dissolution in two systems with temperature decreasing from 200 °C to 50 °C (Fig. 9). In the system with fixed \( p_{\text{CO}_2} \) (223 bar) during the cooling processes, 1 kg of water may dissolve 5.01 g calcite (Case-1), while in the system when \( p_{\text{CO}_2} \) decreases from 223 bar at 200 °C to 0.32 bar at 50 °C (according to the equation \( \log p_{\text{CO}_2} = -1.45 + 0.019 T \) (Smith and Ehrenberg 1989), 1 kg of water can dissolve only 0.027 g calcite (Case-2). In such cases, the pore water in sandstones with 20% porosity can only dissolve calcite (with specific gravity of 2.7 g/cm³) to increase porosity by 0.037% and 0.0002%, respectively, with the occurrence of retrograde dissolution.
**4.3 Buffer system and pH needed for burial carbonate dissolution**

Diagenetic reactions in intermediate to deep burial regimes are mutually rock buffered (Bjørlykke and Jahren 2012; Hutcheon and Abercrombie 1990; Macquaker et al. 2014; Smith and Ehrenberg 1989; Taylor et al. 2010; Turchyn and DePaolo 2011). The carbonate minerals were commonly

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**Fig. 8** a Plot of pH values versus depth for CaCO$_3$–H$_2$O–CO$_2$ equilibrium systems with different CO$_2$ contents; b–d volume increment of calcite and dolomite by per liter liquid (V) versus depth for CaCO$_3$–H$_2$O–CO$_2$, CaMg(CO$_3$)$_2$–H$_2$O–CO$_2$ systems, the CO$_2$ mol fractions are 0.1%, 1%, and 10%, respectively, after the systems reached equilibrium. (after Huang et al. 2009)

**Fig. 9** Numerical simulation results of the cooling of hot fluids from 200 to 50 °C in a calcite–CO$_2$–H$_2$O system with initial log ($p_{CO_2}$) values of 2.35. Case-1: Simulation was conducted with a fixed $p_{CO_2}$; Case-2: Simulation was conducted with a variable $p_{CO_2}$ (Yuan et al. 2015a, b, c)
suggested to react faster with acids than the aluminosilicate minerals (Bjørlykke and Jahren 2012). In the buried aluminosilicate–carbonate mineral–acid system, however, Smith and Ehrenberg (1989), Hutcheon and Abercrombie (1990), and Turchyn and DePaolo (2011) suggested that the aluminosilicate minerals–water interaction rather than the carbonate mineral–water reaction was the main acid-buffering mechanism (Bjørlykke and Jahren 2012; Hutcheon and Abercrombie 1990; Macquaker et al. 2014; Smith and Ehrenberg 1989; Taylor et al. 2010; Turchyn and DePaolo 2011). The buffer intensity of silicate minerals can be ten times that of calcite in an acidic system at high temperature (Hutcheon and Abercrombie 1990). The pH of most current oil–gas waters is higher than 5.5 due to the buffering effect of various aluminosilicate mineral–water interactions (Birkle et al. 2009; Birkle et al. 2002; Eggeberg and Aagaard 1989; Frappe et al. 1984; Surdam et al. 1985), and the extensive dissolution of carbonate minerals is unlikely in reservoirs with such a relative weaker acidity. This concept is a rather radical departure from the conventional system, but it is now being verified by the significant fabric observation of extensive feldspar dissolution and no/little carbonate dissolution in many buried sandstones (Armitage et al. 2010; Baker et al. 2000; Ceriani et al. 2002; Dos Anjos et al. 2000; Dutton and Land 1988; Fisher and Land 1987; Girard et al. 2002; Hendry et al. 1996; Milliken et al. 1994; Salem et al. 2000; Tobin et al. 2010) and some mudstones (Macquaker et al. 2014; Turchyn and DePaolo 2011). Yuan et al. (2015a, b, c) proposed the mechanism of selective dissolution of feldspars in the presence of carbonate minerals to generate secondary minerals in buried sandstones by organic-origin CO₂ (Yuan et al. 2015a, b, c). In addition, the dissolution of feldspars can, in turn, promote the precipitation of carbonate minerals (Tutolo et al. 2015). The C–O isotopic data of feldspars can, in turn, promote the precipitation of carbon-derived and inorganic-derived CO₂. The most carbon in these various types of CO₂, however, is subsequently sequestered by the precipitation of carbonate cements in both source rocks and reservoirs (Curtis 1978; Giles and Marshall 1986; Seewald 2003).

4.4 Mass transfer problem

In order to generate enhanced secondary porosity, the solutes (Ca²⁺, Mg²⁺, HCO₃⁻, CO₃²⁻) released by the dissolution of carbonate minerals need to be removed from the dissolution zone in the sandstone reservoirs (Bjørlykke and Jahren 2012; Ehrenberg et al. 2012; Giles 1987; Taylor et al. 2010). Advection, diffusion, and convection are the three possible mechanisms that control the mass transfer in the sedimentary basins. However, none of the advective, diffusive, or convective mass transfer supports significant transfer of the solutes released from carbonate dissolution in the buried sandstones without favorable flow conduits (e.g., faults and fractures) in the mesodiagenetic stage (Bjørlykke and Jahren 2012; Ehrenberg et al. 2012; Giles 1987; Taylor et al. 2010).

(1) Advection transfer

Mass transport of a component by the advective flow in subsurface porous rocks can be expressed by

\[ q_A = qC \]  (1)

where \( q_A \) is the advective flux of the species, \( q \) is the specific discharge, and \( C \) is the component’s concentration. The solubility of calcite is a function of the \( p_{CO_2} \) and temperature in the burial sediments, and the calcite solubility is less than 0.01 mol/L in systems at temperatures ranging from 80 °C (with 1 bar \( p_{CO_2} \)) to 160 °C (20 bar \( p_{CO_2} \)) (Giles and De Boer 1989). Assuming that a set of sediments has a mudstone/sandstone ratio of 10:1 and the mudstone porosity decreases from 20% to 5% as the burial depth increases from 2000 m to 4000 m (Pittman and Larese 1991), all the water in the mudstone units would be expelled to the sandstone units. The water from the mudstone, even with very low salinity, can dissolve and remove only approximately 0.05% volume of the calcite mineral in the sandstone units under the mechanism of advective flow.

(2) Diffusive transfer

Mass transport by diffusion \( (M_i) \) in porous rocks can be expressed by Fick’s law:

\[ M_i = -\phi D_0 \times \theta^2 \times \frac{dC}{dx} \]  (2)

where \( M_i \) is the diffusion flux, \( D_0 \) is the diffusion coefficient of solutes in water (cm²/s), \( C \) is the component’s concentration, and \( \theta^2 \) is the tortuosity factor of the sedimentary rock. Tortuosity is generally a ratio of pore connectivity length to sediment sample length; thus, its value is always greater than 1. In porous sedimentary rocks, the tortuosity of the flow path is determined by porosity, permeability, and pore structure. Tortuosity can be expressed by Archie’s equation (Archie 1942) as:

\[ \theta^2 = \phi^{1-\eta} \]  (3)

where \( \eta \) is an adjustable exponent (Boudreaux 1996). The empirical fit value of \( \eta \) reported by Boudreaux (1996) is 2.14 ± 0.03, with an average value of 2.14. Diffusion in a porous sediment system is much slower than in an equivalent volume of water because the convoluted path the solutes must follow to circumvent sediment particles (Boudreaux 1996).

The pore water composition in the middle-deep buried sandstones is generally close to saturation with respect to most minerals after long-term contact of the pore water and
minerals. The solute concentration gradients (Ca$^{2+}$, Mg$^{2+}$, HCO$_3^-$, CO$_3^{2-}$) are generally very low in the sandstone beds with relative homogeneous composition (Bjørlykke 2014; Bjørlykke and Jahren 2012), which prevents the large-scale diffusive transfer of these masses, even in a long geological time.

(3) Convective transfer

Thermal convection is a potential mechanism for mass transfer in buried sandstones with high porosity and permeability. Mathematical calculations of thermal convection, however, demonstrated that even thin interbedded layers of mudstones within permeable sandstone sequences will split potentially larger convection cells into smaller units of sandstone beds which may then be too small to exceed the critical Rayleigh number. After extensive compaction and cementation, convective circulation of pore water is not likely to occur in the sandstone beds with interbedded mudstones characterized by extremely low porosity and low permeability (Bjørlykke 1993; Bjørlykke et al. 1988), particularly after the formation of tight marginal cemented barriers formed by precipitation of eodiagenetic carbonate cements in the marginal sandstones close to the mud/sand surfaces (Saigal and Bjørlykke 1987).

Overall, large-scale carbonate dissolution is not supported by any of the mass transfer mechanisms in buried geochemical systems without favorable flow conduits.

5 Published examples of carbonate dissolution in sandstones

Many papers on mineral dissolution in buried sandstones have been published in the last 40 years. After a careful analysis of the included petrography evidence on the dissolution of carbonate minerals, we categorize these publications into three groups.

5.1 Papers lacking convincing petrographic evidence but including hypothesis

Schmidt and McDonald (1979a, b) first proposed the idea that significant secondary porosity (up to 20%) can be generated through burial dissolution of carbonate cements by CO$_2$ during the organic maturity stage (Fig. 1) (Schmidt and McDonald 1979a). In their paper, however, no convincing carbonate dissolution phenomena such as the microscope photography in Taylor (1996) or the SEM microphotography in Weedman et al. (1996) and Khidir and Catuneanu (2003) were presented (Khidir and Catuneanu 2003; Schmidt and McDonald 1979b; Taylor 1996; Weedman et al. 1996). Instead, most intergranular pores without carbonate cements were interpreted as secondary pores formed by the dissolution of carbonate cements (Taylor et al. 2010). Based on the CO$_2$/organic acids leaching hypothesis, and the negative relationship between porosity and the amount of carbonate cements in reservoirs, extensive burial dissolution of carbonate minerals has also been suggested by many other authors in the last few decades (Dutton and Willis 1998; Gibling et al. 2000; Harris and Bustin 2000; Higgs et al. 2010; Irwin and Hurst 1983; Khidir and Catuneanu 2010; Kordi et al. 2011; Mcbride 1988; Shannugam 1984; Wilkinson et al. 1997). Similar to Schmidt and McDonald (1979a, b), no convincing petrography evidence on carbonate dissolution was provided in these publications.

5.2 Papers with convincing petrographic evidence

Some studies with convincing petrographic evidence of the dissolution of carbonate cements in buried sandstones have been published. Review of these works suggests that most of the extensive dissolutions were related to deep hot fluids (Taylor 1996; Taylor et al. 2010) and cold meteoric fresh water (Bouch et al. 2006; Cavazza et al. 2009; Khidir and Catuneanu 2003; Poursoltani and Gibling 2011; Yuan et al. 2017; Zaid 2012), while organic CO$_2$ leaching resulting in very limited dissolution (Lu et al. 2011; Weedman et al. 1996).

(1) Dissolution related to deep hot fluids. Taylor (1996, 2010) presented a striking exception in the deeply buried (4.9–5.2 km) Miocene sandstones in the Picaroon field, offshore Texas. The anomalously high porosities (20%–29%) in the sandstone reservoirs are largely a result of the porosity enhancement by the dissolution of carbonate minerals (Taylor 1996; Taylor et al. 2010). A detailed petrographic study has established evidence for the partial dissolution of pore-filling calcite cements and detrital carbonate grains in the most porous sands (Fig. 2f), and 6%–15% of the calcite cements were estimated to have been removed from the reservoirs. However, the aggressive fluids that leached the cements were not acids originating from kerogen maturation but deep hot fluids with high salinity and high concentrations of Sr, Ba, Fe, Pb and Zn. The Corsair fault systems played a very important role in introducing such hot fluids into the Miocene sandstones. As Taylor et al. (2010) suggested, the geological conditions in the Picaroon field that provide access to deep fluid sources are somewhat extraordinary (Taylor 1996; Taylor et al. 2010).

(2) Dissolution related to meteoric water. Convincing petrographic evidence of carbonate dissolution in sandstones with meteoric freshwater incursion was provided by some studies. The isotopic composition of the car-
carbonate cements or the burial history of the rocks suggests that the dissolutions were induced by meteoric freshwater during the early eodiagenetic stage or the uplift telodiagenetic stage (Bouch et al. 2006; Cavazza et al. 2009; Khidir and Catuneanu 2003; Poursoltani and Gibling 2011; Yuan et al. 2017; Zaid 2012). For example, Khidir and Catuneanu (2003) presented convincing SEM photomicrographs to show the dissolution of carbonate cements in the Scollard sandstones outcrop. The sandstones were not buried at depths where the temperature reached 120 °C, and the δ18O composition of the included calcite cements and relevant calculated water δ18O composition suggested a meteoric origin of the diagenetic fluids. Poursoltani and Gibling (2011) provided an example of the dissolution of carbonate cements in sandstones with developed fault systems that were formed during an uplift period; the calcite cements were suggested to be leached by meteoric water. Cavazza et al. (2009) provided excellent outcrop photographs, microscope photos, and SEM images to show carbonate cement dissolution in the Quaternary marine terraces outcrop sandstones; the dissolution was the leaching result of meteoric flow during the falling period of sea level.

(3) Organic CO2 leaching with weak dissolution. Some authors have presented convincing petrographic evidence of carbonate dissolution that was induced by organic CO2 originating from the thermal evolution of kerogen. The authors, however, also stated that only a small amount of carbonate minerals were dissolved under the constraints of fluid chemistry modeling or mass balance calculation (Lu et al. 2011; Weedman et al. 1996).

Overall, review of the published papers suggests that the extensive dissolution of carbonate minerals in the sandstones was generally attributed to the high flux of deep hot fluids provided via fractures or the meteoric freshwater available during the eodiagenetic and telodiagenetic stages.

5.3 Papers with dissolution of feldspars but no carbonate

Some studies have reported the phenomena of extensive feldspar dissolution in buried sandstones, but the carbonate cements and detrital carbonate grains in the sandstones showed no signs of extensive dissolution (Armitage et al. 2010; Baker et al. 2000; Ceriani et al. 2002; Dos Anjos et al. 2000; Dutton and Land 1988; Fisher and Land 1987; Girard et al. 2002; Hendry et al. 1996; Milliken et al. 1994; Salem et al. 2000; Tobin et al. 2010). For example, Yuan et al. (2015a, b, c) presented convincing photomicrographs to show the selective dissolution of feldspar minerals in the presence of carbonate minerals including detrital carbonate grains and early precipitated calcite cements. These carbonate cements display euhedral crystal faces where they border open primary pores, and the detrital carbonate grains show no corroded fabrics that occur in feldspar grains. Aside from the geological examples of buried sandstones, numerical simulation results also demonstrated that the feldspar dissolution induced by carbonic acid in the subsurface sandstones would be accompanied by carbonate precipitation in systems in the long term, and these relevant carbonate cements serve as an analogue of late-stage carbonate cements in the subsurface sandstones (Barclay and Worden 2000; Wilson et al. 2000).

6 Conclusions

(1) Four types of selective dissolution assemblages of feldspar and carbonate minerals can be identified in sandstones. A critical eye must be cast on the identification of mineral dissolution and intergranular secondary pores in sandstones so that possible subjective conclusions can be avoided.

(2) Petrographic data, porosity data, water–rock experiments, geochemical calculations of aggressive fluids, and mass transfer do not support significant mesodiagenetic carbonate dissolution in buried sandstones. A review of relevant publications suggests that the extensive dissolution of carbonate minerals was generally attributed to a high flux of deep hot fluids provided via fractures or the meteoric freshwater available during the eodiagenetic and telodiagenetic stages.

Acknowledgements This study was funded by the Natural Science Foundation of China Project (Nos. 41602138, 41872140, 41911530189), the National Science and Technology Special Grant (No. 2016ZX05006-007; No. 2016ZX05006-003), the Fundamental Research Funds for the Central Universities (18CX07007A), and the State Key Laboratory of Organic Geochemistry, GIGCAS (No. SKLOG-201709).

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