Zeta Potential of a Natural Clayey Sandstone Saturated With Carbonated NaCl Solutions at Supercritical CO₂ Conditions

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Abstract The zeta potential is a measure of electric potential at the mineral-electrolyte interfaces. The zeta potential of natural sandstones depends on mineralogy, electrolyte pH, concentration, amount of dissolved CO₂, and temperature. We report for the first time the zeta potential measured on clayey sandstone comprising quartz, kaolinite, illite, albite and microcline saturated with NaCl solutions at supercritical CO₂ conditions. Our results demonstrate that zeta potentials in clayey sandstone samples at supercritical CO₂ conditions are significantly different from similar measurements conducted under ambient conditions and from those obtained with clean sandstones. Supercritical CO₂ zeta potential remains negative but is influenced by clays and feldspars due to their significant presence and exposure to large pores, which yields less negative zeta potential compared to quartz, under identical conditions. Our results have significant implications to natural subsurface systems such as CO₂ geo-sequestration sites, aquifers, geothermal sources and hydrocarbon reservoirs.

Plain Language Summary Sandstone is a type of porous rock commonly found in natural subsurface settings. Generally, the main mineral found in sandstone is quartz. However, other minerals such as clays and feldspars are also often present in sandstones. When the formation water flows through porous sandstone, the rock surface becomes electrically charged as water contacts minerals of the rock. The property that describes electrical properties of rock-water interface is termed the zeta potential, and it can be used to characterize subsurface flows and the wetting state of rocks. The zeta potential can be measured using the streaming potential method in the laboratory under conditions representative of subsurface systems. In this study we show that the zeta potential of clayey sandstone at supercritical CO₂ conditions is significantly different from the published data on clayey samples at ambient conditions and from clean sandstones comprising mostly quartz. Therefore, ignoring the effect of high content of dissolved CO₂ at high pressure and elevated temperature and the fact that other minerals in sandstone can modify the zeta potential might lead to misinterpretation of data and model estimates used a broad range of earth and environmental science applications.

1. Introduction

Zeta potential is a physicochemical property of interfaces between aqueous solutions (termed brines or electrolytes for simplicity) and minerals or other fluid phases. The mineral-brine zeta potential is interpreted from the measured streaming potential that arises due to pressure gradients in rocks saturated with electrolytes (Hunter, 1981; Jackson et al., 2012). Many experimental and modeling studies have reported the importance of the zeta potential for a broad range of applications for silica-brine systems, due to abundance of such formations. These applications include CO₂ geo-sequestration (CGS, e.g., Moore et al., 2004), hydrocarbon recovery (e.g., Jackson et al., 2016), geothermal resources (Jardani et al., 2008; Revil & Pezard, 1998), characterization of flow through fractured systems (Jougnot et al., 2020; Vinogradov et al., 2022), and management of groundwater aquifers (e.g., MacAllister et al., 2018).

The zeta potential of sandstone saturated with electrolytes at various conditions has been widely investigated experimentally covering high salinity (Walker & Glover, 2018), elevated temperature (Vinogradov & Jackson, 2015), multi-phase flows (Revil & Cerepi, 2004; Sprunt et al., 1994) and complex composition of brines...
(Thanh & Sprik, 2016). However, the only published data on zeta potentials of natural sandstones at CGS conditions (elevated temperature, high pore pressure, high CO₂ content) are limited to quartz dominated rock sample (Hidayat et al., 2022, hereafter referred to as H22), where measurements were carried out on a “clean” Fontaineb-leau (>99% wt quartz) sample at pore pressures up to 10 MPa and temperatures up to 40°C were reported. The authors found that the zeta potential remained negative, decreased with increasing content of dissolved CO₂, which resulted in a lower equilibrium pH. Many natural sandstone reservoirs comprise a broad range of various minerals, including clays, micas, feldspars and carbonates. These minerals are known to have a significant impact on the zeta potential (e.g., Hoxha et al., 2016; Peng et al., 2020) through the electrochemical interactions between individual minerals and ionic species at the mineral-water interface. These interactions result in establishment of micro-scale local zeta potentials for each mineral, which combined together define to the so-called effective or macro-scale zeta potential that is measured in cm-scale porous rocks with complex mineralogy (Collini & Jackson, 2022). The dependence of the macro-scale zeta potential on mineralogy of clayey sandstones under varying pore pressure, brine concentration and CO₂ content conditions, relevant to CGS remains largely unknown.

In the absence of experimental data of the zeta potential at such conditions, we report for the first time zeta potentials measured for a natural sandstone comprising quartz, feldspars, and clay minerals, in contact with CO₂-rich NaCl solutions. The experimental pore pressure and temperature conditions were varied between 0.2 and 10 MPa and between 23°C and 40°C, respectively. We demonstrate that the zeta potential response to varying pore pressure is unique and different from that of clean sandstone samples.

2. Materials and Methods

A cylindrical San Saba sandstone sample was used in this study. The petrophysical properties of the sample are provided in Table 1.

Prior to conducting the streaming potential measurements, the sample was cleaned following the procedure reported by Alroudhan et al. (2016) and at compatible with clay minerals conditions (see Supporting Information S1, and refer to McPhee et al., 2015). The porosity of the sample was measured using the gas (N₂) expansion method in AP-608 Automated Permeameter and Porosimeter (Coretest System Inc, USA). The liquid permeability and the formation factor (using NaCl solutions of concentration ranging between 0.05 and 1.0 M, M = mol/L) were calculated following the procedure reported in Vinogradov et al. (2010).

The experiments were conducted using NaCl aqueous solutions of varying ionic strength and were divided into two main groups corresponding to the electrolyte conditions: dead and live solutions. The former relates to a synthetic aqueous solution fully equilibrated with atmospheric CO₂, whereas the latter solution prepared under ambient pressure and temperature is brought into contact with pure CO₂ in a mixing reactor establishing thermodynamic equilibrium at elevated temperatures and pressures (for further details, refer to H22). To prepare dead and live electrolytes, we followed the experimental protocol reported by H22 to make sure the dead and live solutions reached full chemical equilibrium with the rock sample before the experiment commenced. The pH and conductivity of the dead electrolytes were regularly measured using a FiveGo pH meter (Mettler Toledo, accuracy of 0.01 pH units) and a Jenway 4520 conductivity meter (Cole-Palmer, 0.5% accuracy), respectively. Furthermore, an in-line high pressure pH meter (Corr Instruments, LLC) was used to measure pH of live electrolytes. The equilibrium pH and conductivity of all tested NaCl solutions are provided in Table 2.

The streaming potential measurements were conducted in a high pressure-high temperature (HPHT) coreflooding apparatus (see Figure 1 in H22). The streaming potential coupling coefficient (C_{SP}) was determined from the ratio between the stabilized voltage, ΔV, and the stabilized pressure difference, ΔP, across the sample following paired-stabilized method (Vinogradov & Jackson, 2011). We confirmed that the effect of surface conductivity was negligible in all our experiments (refer to Supporting Information S1) and therefore, the zeta potential (ζ) was calculated using the classical Helmholtz-Smoluchowski equation (Journiaux & Pozzi, 1995) while disregarding the Overbeek’s correction for the surface conductivity (Glover, 2015):

\[
C_{SP} = \frac{\Delta V}{\Delta P} = \frac{\varepsilon \varepsilon_0}{\mu \sigma_w} \tag{1}
\]

where ε is the electrolyte permittivity, μ is the electrolyte dynamic viscosity, and σ_w is the conductivity of the electrolytes (Table 1). The permittivity (ε) and dynamic viscosity (μ) of dead electrolytes were evaluated using
Table 1
Petrophysical and Mineralogical Properties of the San Saba Sample Used in This Study

<table>
<thead>
<tr>
<th>Sample</th>
<th>San saba</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mineralogy</td>
<td></td>
</tr>
<tr>
<td>87.7% wt. – 95.2% wt. quartz</td>
<td></td>
</tr>
<tr>
<td>2.3% wt. – 6.2% wt. microcline</td>
<td></td>
</tr>
<tr>
<td>1.9% wt. – 4.0% wt. albite</td>
<td></td>
</tr>
<tr>
<td>1.1% wt. – 3.3% wt. kaolinite</td>
<td></td>
</tr>
<tr>
<td>0.9% wt. – 1.0% wt. illite</td>
<td></td>
</tr>
<tr>
<td>Porosity</td>
<td></td>
</tr>
<tr>
<td>21.0% ± 1.0%</td>
<td></td>
</tr>
<tr>
<td>Liquid permeability</td>
<td></td>
</tr>
<tr>
<td>35.0 ± 1.0 mD</td>
<td></td>
</tr>
<tr>
<td>Dimensions</td>
<td></td>
</tr>
<tr>
<td>Length = 7.61 cm</td>
<td></td>
</tr>
<tr>
<td>Diameter = 3.82 cm</td>
<td></td>
</tr>
<tr>
<td>Formation factor, $F$</td>
<td>159 ± 8</td>
</tr>
</tbody>
</table>

Note. Mineralogy of the sample is provided as a range of the main constituents based on published studies of Connelly et al. (2019) and Al-Shajalee et al. (2020) and independently confirmed from X-ray diffraction and Scanning Electron Microscopy Analyses carried out in this study.

Table 2
The Equilibrium pH and Conductivity of Tested Electrolytes in This Study

<table>
<thead>
<tr>
<th>Ionic strength, M</th>
<th>$P$, MPa</th>
<th>$T$, °C</th>
<th>Solution</th>
<th>$\sigma_e$, S/m</th>
<th>pH values</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.05</td>
<td></td>
<td></td>
<td>Dead</td>
<td>0.550</td>
<td>6.80 ± 0.1</td>
</tr>
<tr>
<td>4.5</td>
<td>0.2</td>
<td>23</td>
<td>Dead</td>
<td>0.557</td>
<td>6.75 ± 0.1</td>
</tr>
<tr>
<td>7.5</td>
<td>4.5</td>
<td>40</td>
<td>Dead</td>
<td>0.553</td>
<td>6.75 ± 0.1</td>
</tr>
<tr>
<td>10.0</td>
<td>7.5</td>
<td>40</td>
<td>Dead</td>
<td>0.555</td>
<td>6.80 ± 0.1</td>
</tr>
<tr>
<td>0.2</td>
<td>10.0</td>
<td>40</td>
<td>Dead</td>
<td>0.601</td>
<td>6.50 ± 0.1</td>
</tr>
<tr>
<td>4.5</td>
<td>0.2</td>
<td>23</td>
<td>Live</td>
<td>0.505</td>
<td>3.72 ± 0.05</td>
</tr>
<tr>
<td>7.5</td>
<td>4.5</td>
<td>40</td>
<td>Live</td>
<td>0.496</td>
<td>3.43 ± 0.05</td>
</tr>
<tr>
<td>10.0</td>
<td>7.5</td>
<td>40</td>
<td>Live</td>
<td>0.493</td>
<td>3.26 ± 0.05</td>
</tr>
<tr>
<td>0.10</td>
<td></td>
<td></td>
<td>Dead</td>
<td>1.015</td>
<td>6.60 ± 0.1</td>
</tr>
<tr>
<td>0.20</td>
<td>0.2</td>
<td>23</td>
<td>Dead</td>
<td>1.982</td>
<td>6.60 ± 0.1</td>
</tr>
<tr>
<td>0.50</td>
<td>0.5</td>
<td></td>
<td></td>
<td>4.110</td>
<td>6.70 ± 0.1</td>
</tr>
<tr>
<td>1.00</td>
<td></td>
<td></td>
<td></td>
<td>8.090</td>
<td>6.70 ± 0.1</td>
</tr>
</tbody>
</table>

Note. $P$ is the mean pore pressure which was kept 2–3 MPa below the confining pressure. $T$ is the temperature during the experiment, which was controlled by the oven. The live electrolyte conductivity was evaluated using an empirical correlation reported by Börner et al. (2015).
4. Discussion

A summary plot of the zeta potential as a function of pH for both live and dead NaCl solutions is presented in Figure 3. The zeta potentials of San Saba were found to be more positive compared with the values of clean Fontainebleau sandstone for live and dead NaCl electrolytes. We observed a steeper slope of zeta potential versus pH in San Saba (blue dashed line, Figure 3) compared with Fontainebleau (black dashed line, Figure 3) when the temperature changed from 23°C to 40°C for the dead electrolytes. Note, that the amount of dissolved CO₂ remained constant during these dead brine experiments at both temperatures. At the same time, a shallower slope of the zeta potential versus pH was observed in San Saba (purple dash line, Figure 3) saturated with live NaCl.
Figure 2.
solutions compared with Fontainebleau (green dashed line, Figure 3) over the range of tested experimental pore pressures between 4.5 and 10.0 MPa. Since the zeta potentials of San Saba and those of clean Fontainebleau samples were obtained under identical conditions of pore pressure, temperature and CO$_2$ content, we attributed differences in respective zeta potentials to the presence of clays and feldspars in San Saba.

A recently published study (Alarouj et al., 2021) suggested that the effective macro-scale zeta potential was an average of the micro-scale zeta potentials from each individual mineral-brine interface. That is, if the mineralogy of the rock sample was dominated by a single mineral such as quartz, then the micro- and macro-scale zeta potentials would be identical. If, on the other hand, various minerals were distributed along the pore walls, as in the case of San Saba sandstone, then the local micro-scale zeta potentials would vary from one mineral to another, and the macro-scale zeta potential would be an average of the micro-scale values depending on the portion of pore walls lined by different minerals.

To evaluate the impact of different minerals on the effective zeta potential of San Saba, additional investigations of the sample were carried out. From X-ray diffraction analysis (XRD), we found the mass fractions of minerals of the bulk sample, in which clays (kaolinite and illite), feldspars (albite and microcline) and quartz were identified as the main components (Table 1). However, the XRD analysis did not provide the necessary insights on how the identified minerals were distributed in the pore space. To address this question, we used the scanning electron microscopy (SEM), from which the same minerals were identified as the main constituents (Figure 4).

SEM image presented in Figure 4c identified a minor inclusion of chrome spinel, which appeared to be an extremely uncommon locality and was only found in a single spot of the thin section. Kaolinite and illite were found to extend toward the middle of large pores. Thus, kaolinite and illite were exposed to electrolyte during the streaming potential experiments and were expected to influence both micro- and macro-scale zeta potentials.

Figure 2. Zeta potential (a, c and e) and electrolyte pH (b, d and f) as a function of salinity, temperature and pore pressure. Zeta potentials (a) and pH (b) for the dead solutions were independent of pore pressure since the experiments were conducted in a closed system while the solutions were prepared with atmospheric level of CO$_2$ corresponding to partial CO$_2$ pressure of 10$^{-14.4}$ atm (Li et al., 2016). Temperature dependence of the zeta potential (c) and pH (d) for dead solutions is compared with previously published data. Live brine zeta potential (e) and pH (f) are compared with previously published data. Green syblos correspond to pore (hence, partial CO$_2$) pressure of 4.5 MPa, red symbols stand for 7.5 MPa and yellow symbols denote experiments at 10 MPa. The Fontainebleau data presented in these figures were extracted from H22.

Figure 3. Summary of zeta potentials versus pH for dead and live electrolytes. The purple symbols correspond to live electrolytes; the blue and red symbols correspond to dead electrolytes at temperature of 23°C and 40°C, respectively. The previously published data for Fontainebleau sample saturated with dead and live NaCl solutions under the same experimental conditions are denoted by black symbols, and these values were extracted from H22.
Similar situation was observed with albite, which was found to extend toward major pores thus interacting with brines during the experiments (Figures 4a and 4c). Hence, albite was also expected to contribute to the overall macro-scale zeta potentials. On the other hand, the microcline grains were primarily locked between quartz grains (Figures 4a and 4b) or had a small area exposed to large pores (Figure 4c), thus limiting the mineral’s interaction with electrolytes during the experiments. Therefore, the contribution of microcline to overall macro-scale zeta potentials was expected to be insignificant despite its relatively significant concentration of c. 2%–6%.

To understand the impact of each mineral, we compared the previously reported zeta potentials of the minerals with that of quartz. Yukselen-Aksoy and Kaya (2011) reported zeta potentials of kaolinite and quartz for pH range of 3–11 in water and found kaolinite zeta potentials to be less negative than quartz. Hussain et al. (1996) conducted zeta potential measurements with kaolinite and illite in water for pH range of 2.5–11 and found that the magnitude of illite zeta potentials was smaller than that of kaolinite. Furthermore, Wainipee et al. (2013)
measured zeta potentials of illite and kaolinite with 0.001 and 0.7 M NaCl and pH range of 1–9, and found that the zeta potentials of illite were less negative compared to kaolinite. Therefore, we concluded that the impact of kaolinite and illite on the negative zeta potentials should lead to the following relative magnitudes $|\zeta_{\text{illite}}| < |\zeta_{\text{kaolinite}}| < |\zeta_{\text{quartz}}|$, all shifting the macro-scale zeta potential of San Saba to more positive values compared with clean Fontainebleau.

A study by Vidyadhar and Rao (2007) reported that microcline zeta potential was more negative than quartz in water for pH range of 1.5–11. On the other hand, the zeta potential of albite was measured to be less negative than quartz in water for pH range of 1.5–11 (Vidyadhar et al., 2002; Wang et al., 2018). Demir et al. (2001) investigated the effect of ionic strength of NaCl electrolytes on the zeta potentials of albite and microcline over a wide salinity range (0.0001–0.1M NaCl), and found that zeta potential in albite was less negative compared to microcline. Based on these findings, we concluded that the magnitude of the negative zeta potential of albite is the smallest, followed by quartz and microcline ($|\zeta_{\text{albite}}| < |\zeta_{\text{quartz}}| < |\zeta_{\text{microcline}}|$).

Numerous experimental studies investigated dissolution of kaolinite and albite, and the dissolution rates were reported to increase with increasing temperature and decreasing pH (Cama et al., 2002; Carroll & Walther, 1990; Harley & Gilkes, 2000; Palandri & Kharaka, 2004). Thus, we hypothesize that kaolinite, albite and illite dissolved during our experiments releasing multi-valent cations such as Ca$^{2+}$, Mg$^{2+}$, Al$^{3+}$ (Cama et al., 2002; Li et al., 2018; Yuan & Pruett, 1998), which caused their adsorption and/or exchange at the quartz surface of San Saba making the zeta potential more positive, as was suggested by published studies of the zeta potential on clayey sandstones (Alarouj et al., 2021; Li et al., 2018).

All in all, based on the bulk composition (Table 1) and pore-scale distribution of main minerals of San Saba (Figure 4) we concluded that the main impact on the macro-scale zeta potential came from kaolinite and albite, both of which had higher presence in the bulk (c. 1%–3% wt. and 2%–4% wt., respectively), and larger available surface area for interactions between ionic species of brines and minerals (i.e., substantial exposure of mineral surfaces to pore fluid in larger pores). Conversely, a smaller amount of illite (c. 1% wt.) and limited exposure to flowing brines of microcline made their contribution to the macro-scale zeta potential negligible. The experimental results of this study confirmed the hypothesized individual contribution of minerals and demonstrated that both, kaolinite and albite of San Saba made the effective macro-scale zeta potentials in this study more positive compared with Fontainebleau (H22). Moreover, these minerals were more reactive with NaCl solutions thus making the pH dependence of the zeta potential significantly different compared with quartz data across the tested range of pH and temperature (Figure 3).

5. Conclusions

We report for the first time zeta potentials measured on intact clayey San Saba sandstone sample saturated with NaCl solutions of various concentrations and under conditions of temperature, pH, dissolved CO$_2$ content and pore pressure consistent with CGS. The zeta potential of San Saba was negative for all tested solutions, but when in contact with live electrolytes it decreased in magnitude with increasing pore pressure due to increased amount of dissolved CO$_2$ and the corresponding decrease in pH. Conversely, the zeta potential of San Saba and dead electrolytes was found to be independent of the pore pressure condition but decreased with increasing temperatures reflecting respective changes in pH. Furthermore, the zeta potential with both, dead and live solutions was less negative when compared with previous experimental and modeling studies of clean sandstones at similar conditions. An anomalous salinity dependence of the zeta potential was observed with dead NaCl of ionic strength 0.05–1.0 M, which is uncommon but qualitatively similar to the previously reported data on a different clayey sandstone (Li et al., 2018).

We found that mineralogy of San Saba played a key role in defining the micro- and macro-scale zeta potentials. Specifically, a larger content of kaolinite and albite combined with their substantial presence in large pores made these minerals to chemically interact with flowing brines, which led to a smaller in magnitude macro-scale zeta potential compared with that of pure quartz. Furthermore, increased dissolution rate of kaolinite, illite and albite at elevated temperature and low pH (around 3) likely resulted in appearance of multi-valent cations in the electrolyte, which further led to specific adsorption and/or ion exchange at the mineral surface, eventually leading to smaller in magnitude zeta potentials compared with quartz-NaCl systems. On the other hand, microcline, mostly locked between quartz grains, and small concentration of illite in San Saba did not have noticeable impact on
the macro-scale zeta potential. Our results demonstrate that mineralogy of sandstones has a noticeable impact on electrochemical interactions at the mineral-brine interface and must be investigated to predict and explain the salinity, composition, pH and temperature dependence of the effective macro-scale zeta potentials.

The results of this study are novel and essential for a broad range of applications including underground gas storage (CO$_2$/H$_2$), monitoring of subsurface flows, geothermal sources and hydrocarbon recovery. However, additional measurements of zeta potentials are required and planned to investigate the effect of electrolyte composition, concentration, pore pressure, temperature and CO$_2$ content over extended ranges and in other clayey sandstone samples.

**Data Availability Statement**

Detailed information on rock sample preparation and petrophysical characterization is available for download in the Supporting Information S1. Summarized experimental data of San Saba sandstone in this study is available for download at [https://doi.org/10.5281/zenodo.6477225](https://doi.org/10.5281/zenodo.6477225).

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