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HAL Id: hal-00932777
https://hal-brgm.archives-ouvertes.fr/hal-00932777
Submitted on 17 Jan 2014

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Well injectivity during CO$_2$ storage operations in deep saline aquifers –

Part 2: Numerical simulations of drying, salt deposit mechanisms and role of capillary forces

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Submitted to

International Journal of Greenhouse Gas Control
Abstract

The injection of CO₂ into geological reservoirs or deep saline aquifers is being studied to control global warming by limiting greenhouse gas emissions. CO₂ is captured from exhaust gases in power plants or industrial units and stored in underground geological reservoirs. Return on experience with CO₂ injection in the oil industry clearly shows that injectivity problems can be encountered due to several mechanisms including mineral dissolution/precipitation and physical alteration due to the complete desaturation of the near-wellbore zone. This study describes numerical modelling that is able to reproduce the experimental results of drying of brine-saturated sandstone cores by gas injection in the laboratory. The evolution of water and gas saturation profiles and the precipitation of salt inside the samples are followed with injection time. Numerical results agree well with experimental observations highlighting the key role played by capillary forces during the desiccation process (see the companion paper; Peysson et al., 2013). A tentative extrapolation of experimental results from laboratory scale to the near-well field scale is proposed. This approach is of major importance because it makes it possible to determine the optimal CO₂ injection flow rate according to both the intrinsic petrophysical properties of the porous medium and initial brine salinities.

Keywords: supercritical CO₂, drying-out effects, coupled modelling, relative permeability.
1. Introduction

Geological sequestration of CO₂ in deep saline aquifers offers a promising solution for reducing net emissions of greenhouse gases into the atmosphere. Nevertheless, this emerging technology based on massive CO₂ injection in saline reservoirs can cause a major disequilibrium of the physical and geochemical characteristics of the host reservoir. Recent numerical simulations indicate that the near-well zone is the area most sensitive to and impacted by the injection of supercritical CO₂. Geochemical reactions (e.g., CO₂ dissolution, pH variation of original brine, mineral dissolution/precipitation) are expected to occur mainly in this zone (Ennis-King and Paterson, 2007; Audigane et al., 2007; André et al., 2007, 2010; Huq et al., 2012). Moreover, physical phenomena (e.g., variations in temperature, pressure, gas saturation) will also generate major perturbations of the equilibrium of the host reservoir (Bielinski et al., 2008; Kopp et al., 2009; Okwen et al., 2011, Menhert and Okwen, 2012; Vilarrasa, 2012). Consequently, all of these processes might combine to enhance or alter its initial porosity and permeability and long-term well injectivity. Detailed studies need therefore to be carried out before effective gas injection in order to predict the sustainability of the system and optimize the injection phase.

Among these perturbations, the desiccation of the porous medium appears to be a major phenomenon with various positive or negative implications such as salt precipitation (Peysson et al., 2010, Ott et al., 2010a, b), modifications of the local geomechanical constraints and internal forces (Peter-Borie et al., 2011), and impacts of injected fluids on the interfacial tensions including capillary/osmotic phenomena (Leroy et al., 2010). Desiccation of porous media subjected to gas injection has been investigated at the laboratory scale on cores (Mahadevan, 2005; Mahadevan et al., 2007; Peysson, 2012) and on a chip (Kim et al., 2013). This process has also been observed at the field scale during the injection of dry gas into oil and gas reservoirs (Kleinitz and Tölcke, 1982; Kleinitz et al., 2001, 2003; Vandeweijer et al., 2011). Salt deposits cause a pressure draw-down in injection wells or decrease productivity in production wells. The modelling of CO₂ storage in deep saline aquifers has
emphasized the importance of drying mechanisms on injectivity (Pruess and Muller, 2009; Giorgis et al., 2007; Hurter et al., 2007; André et al., 2007; Zeidouni et al., 2009; Alkan et al., 2010) and has led to new research efforts on that topic.

Mechanistically, the massive and continuous injection of CO$_2$ in an initially saturated porous medium causes water displacement and evaporation. First, mobile water is pushed out by the injected supercritical CO$_2$ according to two-phase displacement (brine - CO$_2$). At the end of this phase, immobile residual water trapped in pores or distributed on grain surface as thin films comes in contact with the flowing dry CO$_2$. Consequently, a continuous and extensive evaporation process leads both to the formation of a reactive drying front moving into the medium, and the precipitation of salts and possibly secondary minerals in residual brines (Mahadevan et al., 2007). This desiccation and the following salt precipitation are also highly influenced by capillary forces, which draw the brine towards the injection points (Giorgis et al., 2007).

This study investigated the consequences of near-wellbore drying-out on well injectivity through the evolution of the petrophysical properties of the porous medium. The objective was to determine the key processes controlling the drying of the porous media during the injection of dry gas, and identify the parameters needed to predict the behaviour of the salt (dissolved and solid) inside the core. Our understanding of these processes is essential for the long-term management of the evolution of gas injection into the reservoir.

The work involved both laboratory experiments and numerical modelling. The drying potential of a gas phase was studied in the laboratory on sandstone plugs (6 cm in length and 4.9 cm in diameter) (Peysson et al., 2013). These experiments were then interpreted using a numerical modelling approach coupling hydraulic, thermal and thermodynamic processes able to simulate the evolution of liquid and gas saturation in time and space. A very fine discretization of the plugs enabled us to capture the continuous evolution of water, gas and salt profiles in the porous medium and estimate the porosity and permeability variations during the drying process.
The parameters established at the centimetre scale were then used without modification in order to extrapolate the results at the near-wellbore scale and predict the impact of CO₂ injection on the petrophysical properties of the host rock. A precise description of fluid (gas and water) flows inside the reservoir enabled us to identify the key processes, and, for given rock properties, the clogging risks due to salt precipitation related to the CO₂ injection flow rate.

2. EXPERIMENTAL APPROACH AT THE LABORATORY SCALE

2.1 Rock samples: characterization and hydraulic properties

Vosges sandstones were selected for this study because it is a rock-type representative of the Triassic formations from Eastern part of Paris Basin (France), which were identified as potential reservoirs for CO₂ storage (Sterpenich et al., 2006). Vosges sandstones, also well-known for their homogeneity, were sampled in a quarry. It is composed of quartz (70 - 80 %), microcline (10 - 20 %) and about 10 % of a mixture of muscovite, albite, smectite, anorthite and hematite (Sterpenich et al., 2006). Cylindrical cores 6 cm in length and 2.5 cm in diameter are prepared from the sampled block.

The hydraulic properties of Vosges Sandstones were characterized: porosity (Φ) and permeability (K₀) were 21.8 % and 74 mD, respectively. A high pressure mercury injection was used to measure the pore throat distribution (Peysson, 2012). This injection also allowed determining the gas-water capillary pressure curve (Fig. 1).

For next numerical simulations, a standard Corey model (Corey, 1954) was used to represent relative permeability curves of both the liquid (kᵢ₀) and gaseous (kᵢ_gas) phases, and a Van Genuchten model (Van Genuchten, 1980) was used to fit the experimental capillary pressure (P_cap) curve (Fig. 1):

\[
k_{i,\text{d}} = \hat{S}^\prime \quad \text{and} \quad k_{i,\text{a}} = (1 - \hat{S})^\prime \quad \text{with} \quad \hat{S} = \frac{S_i - S_w}{1 - S_w - S_o}
\]  

(1)
where \( S_l \), \( S_{lr} \) and \( S_{gr} \) are the liquid saturation, the irreducible liquid saturation and the irreducible gas saturation, respectively. For the Vosges Sandstone sample, \( r = 5.5 \) and \( p = 2 \). The irreducible water saturation \( (S_{lw}) \) was 0.19 and the irreducible gas saturation was nil.

\[
P_{cap} = -P_0 \left[ \left( S_l \right)^{1/m} - 1 \right]^{m} \quad \text{with} \quad P_0 = 1.82 \times 10^4 \text{ Pa} \quad ; \quad m = 0.675 \quad ; \quad S_l = \frac{S_l - S_{lw}}{S_u - S_{lw}} \tag{2}
\]

All the parameters used in Eq. (1) and (2) were adjusted from different drainage capillary curves recorded on Vosges Sandstone cores (results not shown here). Specific efforts were made in particular to estimate \( P_0 \), the entry pressure, because it is a key parameter in the prediction of the gas breakthrough. The selected value of 182 mbar was in agreement with values proposed by Peysson (2012).

### 2.2 Brine and non-wetting fluid

Before experiment, Vosges sandstone cores were fully saturated with brine composed of 55% KCl and 45% KI, the latter intensifying the density contrast for X-ray measurements. According to experiments, water salinity was 35 and 150 g/L.

The first objective of this work was to determine the drying potential of the gas phase on sandstone plugs. The best solution was to use \( \text{CO}_2 \) in a supercritical state (i.e. at pressures greater than 7.4 MPa and temperatures higher than 31.1 °C). But, because of experimental constraints (stability of large volumes of \( \text{CO}_2_{sc} \), maintenance of the devices, problems of injection...), nitrogen replaced \( \text{CO}_2_{sc} \) as the non-wetting fluid and temperature and pressure were set to 80°C and 50 bars, respectively. In these conditions nitrogen is a gas with drying properties close to those of \( \text{CO}_2 \) (as explained in paragraph 3.1). The experimental results with \( \text{N}_2 \) on core drying were thus comparable with those obtained on experiments performed with \( \text{CO}_2 \).
2.3 Experimental setup

In the laboratory, nitrogen was injected into Vosges sandstone cores encapsulated in X-ray transparent cells and placed in an oven under controlled temperature (80 °C) (Fig. 2). The cell is composed of a membrane in order to confine the core at a pressure of 50 bar. Experiments are performed by applying, between the inlet and the outlet of the column, increasing pressure plateaus in one case, and a constant pressure difference in other. The local water saturation in the sample was measured during the injection of dry gas (nitrogen) with X-ray attenuation techniques. The experiments continued until the cores were completely dry. The volume of water expelled was measured in a burette at the column outlet, and a gas flow meter recorded the outlet gas flow rate (Peysson et al., 2010, 2013; Peysson, 2012). Gas permeability and porosity profile of cores were measured at the beginning and at the end of the experiment to evaluate the role of the drying.

2.4 Experimental results

For the first experiment, three pressure plateaus (120 mbar, 800 mbar and 1500 mbar) were imposed to the core initially saturated with a 35 g/L of brine (Peysson et al., 2013). During the first one, a two-phase flow displacement occurred with a rapid gas breakthrough (at t = 5 hours) and a low gas flow rate (Fig. 4). The saturation profile in the core showed a typical capillary fringe close to the outlet of the sample. During the second plateau, core continued to desaturate, with a significant contribution of drying. The saturation profiles are flatter because of capillary back flows. For the highest pressure, drying is very rapid, lasting only a few hours. During this phase, a drying front moved from the inlet to the outlet of the core. The dissolved salt, initially present in the brine and transported in the capillary flows, accumulated as precipitate near the injection surface, involving a decrease of about 70 % of the initial permeability.

For the second experiment, a constant difference pressure (300 mbar) was imposed to the core initially saturated with a 150 g/L of brine. A rapid gas breakthrough was observed (less
than 2 hours) with an increase of the gas flow rate up to 26 hours. Then a plugging of the
core occurred with no gas flow after 46 hours. The salt accumulation near the injection
surface generated by capillary back flows was sufficient to block the gas flow.

3. MODELLING TOOLS AND NUMERICAL PARAMETERS

TOUGH2 (Pruess et al., 1999) was used for all of the numerical simulations done in this
study (core experiments and field scale). This code couples thermal and hydraulic processes
and is applicable to one-, two-, or three-dimensional, physically heterogeneous geologic
systems. It is a modular code that can be adapted to a broad range of applications by simply
using interchangeable calculation modules. Two modules were used for this study:

- The EOS7C module (Oldenburg et al., 2004) is a fluid property module developed
  specifically to deal with mixtures of non-condensable gases (like CO$_2$ or N$_2$) and methane. It
can be used to model isothermal or non-isothermal multiphase flow in water/CH$_4$/\(\text{CO}_2\) or N$_2$
  systems. This module uses a cubic equation of state and an accurate solubility formulation
  along with a multiphase Darcy's Law to model flow and transport of gas and aqueous phase
  mixtures over a wide range of temperatures and pressures (See Appendix A).

- The ECO2n module (Pruess, 2005) is specifically developed to model isothermal or non-
  isothermal multiphase flow in water/brine/CO$_2$ systems in deep geological reservoirs
  (aquifers or depleted gas fields). It is based on the equilibrium between the supercritical
  phase and the aqueous phase assumed for CO$_2$ dissolution. Hence,

  \[
  \text{CO}_2(\text{sc}) \Leftrightarrow \text{CO}_2(\text{aq}) \tag{3}
  \]

  where(sc) and (aq) denote supercritical and aqueous carbon dioxide, respectively. An
  extension of Henry’s Law is used to estimate the dissolution of CO$_2$:

  \[
  K \Gamma P = \gamma C \tag{4}
  \]
where $K$ is the equilibrium constant depending on the temperature $T$, $P$ is the partial pressure of CO$_2$, $\gamma$ is the aqueous CO$_2$ activity coefficient, $C$ is the aqueous concentration (mol/kg H$_2$O), and $\Gamma$ is the fugacity coefficient depending on pressure and temperature.

### 3.1 Thermodynamic properties of non-wetting fluids (CO$_2$ and N$_2$)

Although our objective is to predict the behaviour of carbon dioxide, laboratory experiments were done with nitrogen in order to avoid ancillary geochemistry mechanisms and focus on the physical effects of drying. Calculations were therefore done to estimate the thermodynamic behaviour of CO$_2$ and N$_2$ and determine how the nature of the fluid might impact the desaturation process. Simulis® Thermodynamics, a code developed by PROSIM S.A. (http://www.prosim.net), was used to define the fraction of water in the two phases (CO$_2$ and N$_2$) according to temperature at 80 °C (Fig. 3). The code integrates various equations of state such as the Peng-Robinson (PR) equation used here for CO$_2$ and N$_2$. The coefficients $K_{i,j}$ used for calculations are those of Soreide and Whitson (1992). The curves obtained with the PR equation were compared to the curve obtained with the Raoult Law (assimilation of N$_2$ and CO$_2$ to perfect gases):

$$y_{\text{w, CO}_2} = \frac{P_{\text{water, sat}}}{P}$$

where $P$ and $P_{\text{water, sat}}$ are the total pressure and the saturated vapour pressure of water, respectively, calculated with the IAPWS95 equation of state (Wagner and Pruss, 2002). The fraction of water in the supercritical CO$_2$ phase is higher than in the N$_2$ phase (about 30 % at 50 bar and 80 °C), whereas the N$_2$ curve is close to the Raoult curve, indicating that N$_2$ can be assimilated to a perfect gas in this range of temperature and pressure (Fig. 3). In a first approximation, the substitution of N$_2$ by CO$_2$ in the modelling phase has a limited impact on the results since the evaporation potentials of the two fluids are similar in these pressure and temperature ranges. Only a shorter desaturation time is expected using CO$_2$. Moreover, the type of gas in the model can be changed by calculating the proper thermodynamic coefficients.
3.2 Modelling core experiments

Two sets of calculations were done to study the experimental results. The first set consisted in using the EOS7C module of the TOUGH2 code to simulate the injection of nitrogen with increasing pressure-steps into a brine-saturated sandstone plug (Fig. 2 and Peysson et al., 2013) in order to determine the time required to desiccate the porous medium and estimate the gas flow rates at the outlet of the core as a function of pressure and temperature (see paragraph 2). A two-phase Darcy’s flow equation was solved using relative permeability and capillary pressure curves. Thermodynamic equilibrium between concomitant phases (water- N₂), similar to the Eq. (3) defined for CO₂, assumes that phase exchanges are responsible for the core drying. This thermodynamic equilibrium is calculated at each time step to determine the watervapour fraction in the gas phase and the fraction of dissolved gas in the brine.

The EOS7C module (version 1.0) does not take into account the salinity of the brine, i.e., it cannot determine either the solubility of the gas in the brine according to its salinity or the amount of salt precipitated inside the core (Oldenburg et al., 2004). Consequently, an additional set of calculations were done with the ECO2n module in order to determine the quantity and location of the salt deposited in the core. In this simulation, CO₂ was injected instead of N₂. This simulation was also used to fit the most influential parameters, such as the relationship between porosity and permeability.

In a second set of calculations, the same procedure was used to study the impact of a constant gas injection flow rate and determine the influence of the desaturation path on the deposit locations inside the core.

A 1D-column model 60mm long was used as a conceptual framework for determining the evolution of the water content induced by the injection of N₂ and CO₂ in both time and space. The column was represented by 60 grid blocks making up the mesh. Each grid cell was 1.0 mm thick. In the lab experiments, the core was cylindrical, whereas in the model, the core is a parallelepiped. The section of the parallelepiped is therefore recalculated to obtain the same
pore volume as the original cylindrical sample. The porous medium (rock matrix) is assumed to be inert with respect to N\textsubscript{2} and CO\textsubscript{2}, i.e. no chemical reactivity. In the next simulations of lab experiments, the initial brine saturating cores only contains sodium chloride.

3.3 Modelling field applications

Two sets of calculation were done to study the field scale. In the first set, a 1D-radial model was used as a conceptual framework for determining the transient evolution of the water content induced by the injection of supercritical CO\textsubscript{2} (considered to be a non-wetting fluid). The generic 1 m-thick reservoir was centred around a vertical injection well. The maximum explored radial distance was 100 km. Along the radius axis, the discretization was very fine close to the well (5 1-mm grid cells, then 5 2-mm grid cells, 77 5-mm grid cells, and 96010-mm grid cells up to 10 m from the injection well). There were 100 grid cells between 10 and 100 m, 100 grid cells between 100 m and 10 km, and 20 grid cells thereafter, up to 100 km. In each interval (between 10 m and 100 km), the width of radial elements followed a logarithmic scale. The objective of such refinement near the injection well was to capture more precisely both the details and the migration of the desiccation front in the near-well region.

In the second set of calculations, a 2D-radial model was used to complement the 1D-radial studies and to include gravity effects. The reservoir was 20 m thick and had a homogeneous structure. The discretization along the radius axis was the same as the one used for the 1D-radial model.

For both calculations, porosity was 21.8 %, K\textsubscript{h} = 74 mD and K\textsubscript{v} = 7.4 mD (K\textsubscript{v}/K\textsubscript{h} = 0.1). The reservoir was initially fully saturated with sodium chloride brine. Capillary and relative permeability curves are given in figure 1. The initial temperature and pressure of the targeted reservoir were 80 °C and 180 bar, respectively. No regional flow was considered and a hydrostatic status was initially assumed for the pressure in the reservoir and maintained constant at the lateral boundary.
For the 1D-radial model, different injection flow rates were used and their impact on the near wellbore integrity was estimated according to the intrinsic rock properties. This was done to estimate the mechanisms able to influence well injectivity. For the 2D-radial model, the injection lasted 10 years with an injection flow rate of 1 kg/s, distributed on the entire thickness of the reservoir. This is the equivalent of a specific flow rate of 0.050 kg/s/m (total amount of injected CO$_2$ = 315.36 kt).

4. Results and Discussions

4.1 Modelling core experiments

**Core drying and salt precipitation for desaturation with pressure-steps**

Simulations with TOUGH2's EOS7C module involved modeling the N$_2$ injection in brine-saturated sandstone cores. The salinity of sodium chloride brine was 35 g/L. Three pressure plateaus ($\Delta P = 0.12, 0.80$ and $1.50$ bars) were applied between the inlet and the outlet of the column to progressively dry the core. The mean water saturation (calculated from the water saturation in each 1-mm core slice) and the outlet gas flow rates were calculated throughout the injection period, for each pressure step. Results obtained with numerical simulations agree well with experimental data (Fig. 4). During each pressure step, the mean water saturation was well predicted and the value obtained at the end of the second step (for $t = 40$ hours) was coherent with the irreducible water saturation ($S_{irr} = 0.19$). This means that all the mobile water was removed from the core, leaving only the residual water inside the medium.

Despite the application of a low pressure difference, the approaches used were sensitive enough to reproduce the overall evolution of the system, even the outlet gas flow rate. The model was able to accurately determine not only the gas breakthrough time at the end of step 1, but also the gas flow rate, regardless of the water saturation inside the column (even when the column is dry).
At the end of the nitrogen percolation experiment, X-ray measurements revealed the precipitation of salt (halite) inside the porous medium. The new estimated permeability was about 22 mD compared to the initial permeability of 74 mD, i.e. a reduction of about 70% due to salt precipitation. Salt precipitation appears to be heterogeneous inside the core, with salt accumulating near the injection surface (Peysson, 2012; Peysson et al., 2013).

This precipitation process was modeled with TOUGH2’s ECO2n module (see Appendix B). However, to understand the evolution of the permeability inside the core, a relationship between porosity and permeability needs to be selected. Verma and Pruess’s “tube-in-series” model (1988) was used because it allows us to reach zero permeability even if porosity is not nil, as experimentally observed. It uses the fraction of pore space occupied by precipitated salt and two other parameters: \( \Pi \), the fractional length of pore bodies and \( \phi_r \), the fraction of initial porosity for which permeability is reduced to zero. The evolution of permeability \( K \), according to the initial permeability \( K_0 \) is given by:

\[
K_{\text{red}} = \frac{K}{K_0} = \theta^2 \frac{1 - \Pi + \Pi / \omega^2}{1 - \Pi + \Pi [\theta / (\theta + \omega - 1)]^2}
\]

where

\[
\theta = \frac{1 - S_s - \phi_r}{1 - \phi_r}, \quad \text{and} \quad \omega = 1 + \frac{1/\Pi}{1/\phi_r - 1}
\]

where \( S_s \) is the solid saturation, i.e. the pore space volume occupied by precipitated salt.

Usually, \( \Pi = \phi_r = 0.80 \) as given in Pruess et al. (1999) and by various other authors (Alkan et al., 2010).

Calculations done with these \( \Pi \) and \( \phi_r \) values underestimated the decrease in permeability. The values were therefore modified and optimized in order to fit a decrease in permeability coherent with experimental data. This was achieved with \( \Pi = 0.80 \) and \( \phi_r = 0.91 \), i.e. a drastic decrease in permeability for a low decrease in porosity (Fig. 5). These values are not out of range since they have already been used in a geothermal field in the Philippines where silica scaling causes high permeability decreases (Xu et al., 2004). These results are also in agreement with the values obtained by Peysson (2012).
Calculations done with these new parameter values gave a decrease in permeability of around 55-60% of the initial permeability, which is not too far from the 70% determined in the experiment. The mean permeability was about 31 mD (compared to an initial 74 mD), with some variations around this value probably due to the boundary conditions chosen for these simulations (Fig. 6). The total amount of precipitated salt was about 1.73 g (compared to the 3.73 g of salt initially dissolved in the original pore water). Figure 6 also shows a large decrease in permeability in the first millimetre, as observed in the experiments. This is due to the accumulation of salt close to the injection zone in agreement with X-ray adsorbance measurements (Fig. 7) (Peysson et al., 2013).

The interpretation of these results enabled us to determine the $k/\Phi$ relationship. The objective was then to apply this approach to another kind of simulation and to determine whether the results of numerical simulations were in agreement with experimental observations.

**Salt precipitation for desaturation with a constant pressure difference**

A sandstone core saturated with brine enriched in KCl (150 g/L) was dried in the laboratory with a constant pressure difference. The nitrogen flow rate at the outlet of the core decreased to zero during the experiment, which shows that clogging occurred inside the core (Peysson, 2012; Peysson et al., 2013). This experiment was modeled with the ECO2n module, by injecting CO$_2$ with a constant pressure difference of 0.3 bar. The previously determined $k/\Phi$ relationship was used to simulate the loss of permeability generated by the deposition of salt inside the core. In the simulation, because of sodium chloride brine, the precipitation of halite is examined.

The precipitation profile obtained numerically shows zero permeability close to the injection point and relatively high values close to the column outlet (Fig. 6). About 1.81 g of salt precipitated – 1.11 g in the first half of the column and 0.70 g in the second half, indicating a massive precipitation close to the column inlet.
Figure 7 shows that salt accumulated in the first millimeters of the core, close to the injection zone. This explains the clogging observed in our experiments (permeability reduced to zero). The results were compared to those obtained in the previous experiment, i.e. the drying according to pressure steps. The shapes of the curves are the same, with a homogeneous deposition of salt throughout the columns, except in the first millimeters where there is clogging. This can be explained by fluid (non-wetting and water) displacement inside the core. Numerical calculations showed that:

- Before the gas reaches the column outlet, both fluids (water and non-wetting) move from the inlet to the outlet of the core. Gas is injected in the column and pushes “free” water out of the core.

- When the gas flow rate is measured at the outlet of the core (i.e. when the column is partially desaturated), the gas continues to move from the inlet to the outlet of the column, whereas water moves in the other direction.

Gas injection results in water back-flow inside the desaturated core due to capillary forces. This transports dissolved salt towards the injection point where evaporation is a predominant mechanism. Consequently, a large amount of salt precipitates, accumulates and has an impact on porosity and then on permeability. The influence of the injection flow rate on the amount of salt deposited in the core cannot be assessed with these two experiments because the brines used did not have the same initial salinity. This is, however, discussed below.

4.2 Modelling at field scale (1D-radial approach)

In order to understand multiphase flow during CO$_2$ injection in saline aquifers, we extrapolated to field scale, first with a 1D and then with a 2D radial model (see paragraph 3.3). For these simulations, we considered that the set of $k_r$-$P_c$ parameters used for predictive simulation at the core scale would enable modeling of CO$_2$ injection at the reservoir scale. Bennion and Bachu (2005) advised that relative permeability characteristics have to be
determined under reservoir conditions to accurately parameterize reservoir models. However, Egermann et al. (2006) demonstrated that both experiments and large scale applications can be simulated using a single set of parameters. The predictive model for CO₂ injection at the reservoir scale is sufficiently accurate if gas-water relative permeability curves and thermodynamic data (solubility of CO₂ in the liquid water phase and water vapour pressure in the CO₂ gas phase) are taken into account in the model. Since TOUGH2 code uses the thermodynamic and thermophysical properties (density, viscosity and enthalpy) of the non-wetting CO₂ phase and the kₚ-Pₚ curves for simulations, we used, as an approximation, a single set of parameters to extrapolate to the reservoir scale.

Numerical simulations were then done at 180 bar and 80 °C, although the Pₚ curve was drawn for 50 bars and 80 °C. The Pₚ curve could have been modified using the Leverett function to take into account this difference in pressure. However, since the gas-brine interfacial tension (IFT) between 50 and 180 bars does not change considerably (Chiquet et al., 2007; Chalbaud et al., 2009), the Pₚ curve shown in figure 1 was used in a first approximation.

The 1D numerical simulations were done to study the possible correlation between brine salinity (sodium chloride content), CO₂ injection flow rate and the spatial distribution of salt (halite) precipitation. Sensitivity calculations are done to test the behavior of the system with respect to these parameters. First, CO₂ was injected into a reservoir containing brine. Two salinities were tested, 10 and 100 g/L, while the same over pressure was applied between the well and the reservoir (ΔP = 2 bars, equivalent to a mean injection flow rate of about 0.015 kg/s/m).

As expected, based on the core scale experiments, supercritical CO₂ injection caused the drying of the porous medium around the well, but results differed depending on brine salinity. The impact on permeability was much greater for the more concentrated brine, with probable clogging near the injection point (Fig. 8). The impact on permeability was not limited to the near-well zone but extended inside the reservoir formation. These calculations show that the
amount of precipitated salt is directly related to the salinity of the brine initially present in the porous medium.

Simulations were also done to determine whether clogging might be prevented with a different injection flow rate. The same simulation as above was done with three different pressure differences between the well and the reservoir. For low salinities (10 g/L) and $\Delta P > 5$ bars (for a mean injection flow rate $> 0.045$ kg/s/m), salt precipitation affected the permeability of the porous medium only slightly (about 5% decrease in permeability) (Fig. 9a). When the injection pressure decreased (low $\Delta P$ and low injection flow rates of around 0.010 to 0.015 kg/s/m), the quantities of precipitated salts increased and the near wellbore porous medium became clogged when the injection flow rate was lower than 0.010 kg/s/m.

For higher brine salinities (100 and 160 g/L), the application of strong over pressure ($\Delta P = 15$ to 40 bars, equivalent to mean injection flow rates of about 0.120 to 0.500 kg/s/m) caused the precipitation of salts around the injector well, decreasing the permeability by about 40 and 60% for brine salinities of 100 and 160 g/L, respectively (Fig. 9a). When the over pressure decreased (injection flow rates lower than 0.030 to 0.050 kg/s/m depending on brine salinity), salt precipitation increased up to the clogging of the near wellbore domain.

The over pressure used to inject CO$_2$ needs, therefore, to be adapted to the salinity of the native aqueous solution and the petrophysical properties ($K$, $kr$, $P_c$ and $\phi$) of the host reservoir. This sensitivity study showed that when the CO$_2$ injection flow rate is too low, a porous medium could become clogged near the injection well even in the presence of a low salinity-brine. An analogy can be made between the gas flow rate and the Péclet number ($Pe$) reported by Peysson (2012) and Peysson et al. (2013). For each salinity/injection flow rate couple, there is a threshold gas flow rate ($Q_t$) and a limit Péclet number (Fig. 9b). For the Vosges sandstone, a brine salinity of 10 g/L entails a threshold gas flow rate of about 0.010 kg/s/m (equivalent to a Péclet number of 430), whereas for higher brine salinity, $Q_t$ increases up to 0.030 kg/s/m ($Pe = 1200$) and 0.050 kg/s/m ($Pe = 2000$) for salinities of 100 g/L and 160 g/L, respectively. Below these threshold values, the system clogs due to the capillary back
flow that moves brine (and dissolved salts) toward the injection point generating/stabilizing an evaporation aureole accumulating salt deposits. The water is evaporated by the continuous injection of gas, leaving solid salt behind reducing permeability to zero according to the $k/\phi$ relationship used for this sandstone. These numerical results are in agreement with laboratory experiments done with this Vosges sandstone (Peysson et al., 2013). These authors observed a clogging of the core with a brine salinity of 150 g/L and $Pe = 360$. However, when the injection flow rate was higher than the threshold value, two-phase displacement seems to be the main process – the water saturation close to the injection surface was sufficiently low to decrease the capillary back flow and limit the back flow of dissolved salts. Salt precipitation near the injection point was thus lowered and salts tended to be deposited farther from the sensitive and strategic near-wellbore zone. Salt precipitated more homogeneously throughout a large near-wellbore zone, decreasing the global permeability of the medium but not clogging the core. In all of these calculations, salt diffusion was disregarded. Indeed, given the high Péclet number values, this hypothesis is valid.

**4.3 Modelling at field scale (2D-radial approach)**

A 2D-radial system was considered in order to develop an integrated approach able to represent the coupling of different thermodynamic mechanisms (such as the evaporation of brine and the equilibrium between gas and liquid phases) and physical processes (such as the transport of a non-wetting supercritical fluid in a saline reservoir, gravity and capillary forces, etc.). We simulated the uniform and continuous injection of CO$_2$ along the entire thickness of the reservoir for 10 years. The reservoir was initially saturated with sodium chloride (NaCl) brine with a salinity of 160 g/L. The specific injection flow rate used (0.050 kg/s/m) was voluntarily close to the $Q_t$ value defined for the 1D-radial model. The objective was to determine whether the clogging predicted by the 1D-radial model for this injection flow rate and brine salinity also occurs on the 2D-radial model.
Because of gravity forces and the supercritical CO$_2$ density (lower than that of the brine), the reservoir dries out faster at the top. Figure 10a clearly shows that the porous medium is dry over about 12 m at the top of the reservoir, whereas only the first meter is dry at the bottom. Inside the drying zone, solid salt (i.e., halite) precipitates. Although salt is present in all of the desaturated zone, the spatial distribution of salt deposits varies according to the prevalent transport forces (advection, diffusion, capillarity, evaporation, etc.) inside the reservoir: the quantities are higher at the bottom of the reservoir and more limited at the top (Fig. 10b) for simulation conditions and the specific characteristics of the reservoir. The pattern of solid saturation (= solid volume/pore volume) indicates that 40% of the porosity is filled with salt at the bottom, whereas only 10% is filled at the top. According to the k/$\Phi$ law established in paragraph 4.1, the porosity and consequently the permeability are more impacted at the bottom of the reservoir (Fig. 10c). The decrease in permeability is represented by the empirical function k-red (= K/K$_0$). This function clearly shows that high permeability reductions are expected near the well (skin effect in the cells adjacent to the well) and in the lower part of the reservoir (in the first meter inside the reservoir).

To explain this distribution of salt inside the reservoir and the massive deposits at the bottom of the aquifer, it is necessary to plot the fluid flows inside the reservoir. Figure 11 shows the directions and trajectories of gas (a) and brine (b) flow inside the reservoir, which initially had a homogeneous structure and petrophysical properties that facilitated fluid circulation throughout the entire thickness (20 m). The lengths of the flow vectors drawn for each grid cell are proportional to the intensity of the flow. The arrows show direction. The vectors are centered on the grid cells.

CO$_2$ vectors are oriented from left (the well) to right, with some deviation towards the top of the reservoir (Fig. 11a). The vertical permeability, ten times lower than the horizontal permeability, limits upward flow, but this is clearly seen in this figure. This figure also shows that gas fluxes are higher in the upper part of the reservoir (between z = 10 and 20 m), which explains the shorter time required to desaturate this zone and its lateral extension.
Brine fluxes in the reservoir are more complex (Fig. 11b). Trends differ depending on the location in the aquifer. Close to the well and within a radius of about 30 m, brine flows toward the injector whereas the gas phase flows away from it. Far from the well, the water flows from left to right and from up to down. Beyond 30 m, the gas phase displaces the water by a piston effect from left to right and, because the gas phase is less dense, the brine is pushed downwards, the size of the arrows indicating the intensity. Close to the well (0-30 m), brine flows towards the injection well. Saline water flows towards the well due to capillary forces that compete with the evaporation process. The length of the arrows shows that the highest flows are at the bottom of the reservoir. In this reservoir zone, capillary forces are great enough to move water in grid cells very close to the well. These flows move the water, which is then evaporated by the continuous injection of desiccant fluid (CO₂), depositing large quantities of salt in this region. At the top of the reservoir, the capillary brine back flows are weaker and less salt is therefore deposited.

This simulation shows that a CO₂ injection specific flow rate of 0.050 kg/s/m causes the clogging of the porous medium in the lower part of the reservoir where capillary brine flows are higher (in agreement with 1D-radial model results). As expected from laboratory experiments, the permeability is clogged by salt deposits because of the injection flow rate which is below the Qₜ value. These results were obtained for a homogeneous reservoir 20-m thick (Figure 12). Although clogging occurs in the lower part of the reservoir, the well injectivity is only moderately altered. However, reservoirs are usually highly heterogeneous and very often made up of thick layers separated by less permeable layers that limit vertical flow. Gas cannot move vertically. Consequently, each reservoir compartment can be considered to be a 1D case with major horizontal gas flows and capillary effects in each sub-domain. Clogging risks are increased in each compartment with a potentially strong impact on well injectivity.
This study clearly shows that a specific integrated approach is necessary in order to take into consideration both thermodynamic mechanisms and physical processes and adapt injection conditions to reservoir properties.

5. Conclusions

Drying processes were studied with laboratory experiments and numerical simulations in order to determine the dynamics of the decrease in water saturation in sandstone and the consequences of induced salt depositions on rock permeability. Experiments at the laboratory scale revealed the major role played by capillary forces and the salinity of the initial brine on the changes in permeability. These experiments also highlighted the relationship between salt deposits, gas injection flow rate and the capillary properties of the porous medium subjected to non-wetting fluid injection, and demonstrated the need to use a coupled and dynamic approach to fully represent the mechanisms involved.

The numerical code used in this study was able to accurately reproduce both the evolutions of the water content in sandstone cores and the salt precipitation with a given brine salinity and various CO$_2$ injection flow rates. The thermodynamic and physical processes were well simulated at small scale on a homogeneous porous material.

Based on our experimental results (rocks characteristics) and several hypotheses (such as the extension of kr/Pc curves obtained at laboratory scale to a large scale, the homogeneity of the reservoir), this study also investigated the behaviors of water, salt and a non-wetting fluid at the reservoir scale, and more precisely in the near-well zone. This integrated approach combining experimental and numerical simulations showed that the salt precipitation process and the amount of salt deposited are related to various parameters such as the salinity of the initial brine. The higher the brine concentration, the greater the salt deposit. However, the location of the deposits inside the reservoir is strongly linked to the supercritical non-wetting fluid injection flow rate and the capillary forces generating backflows. The numerical simulations showed that the capillary properties of the rock prevent a
sudden evaporation of the irreducible water by continuously feeding the injection zone with
“new” brine coming from reservoir zones far from the injection well. However, a sufficiently
high gas injection rate can overcome the capillary forces and limit the precipitation of salts
close to the injection well. Sensitivity analysis made it possible to determine a threshold gas
flow rate ($Q_t$) dependent on brine salinity. Below this threshold value, there is a high risk of
clogging, whereas higher values ($Q > Q_t$) can reduce the risk of clogging. However, if high
gas injection flow rates in high saline aquifers can prevent clogging, decreased permeability
is expected, even far from the injection well zone.

This study highlights that all these parameters ($kr/Pc$ curves, brine salinity, pressure and CO$_2$
injection flow rates) must be known in order to improve the management of the industrial
injection of CO$_2$ in saline aquifers. The real structure of the reservoir as its heterogeneity (not
considered in this study) has also to be taken into account as a key parameter because it
could modify the capillary fluxes and thus, the localisation of the salt deposits.

**Acknowledgements**

This work was carried out within the framework of the “ProchePuits” project, co-funded by the
French National Agency for Research (ANR). The authors thank Marc Parmentier (BRGM)
calculating the water content in the gas phases with the Simulis© Thermodynamics code.
They are grateful to all of the project partners (TOTAL, GDF Suez, Schlumberger, Itasca,
CNRS, University de Lorraine, University of Pau) for authorizing the publication of this work.
Appendix A: Governing Transport Equations and Resolution

In TOUGH2 Code, the mass conservation equations governing the flow of multiphase, multicomponent fluids in permeable media can be written in the following form:

\[ \frac{d}{dt} \int_{V_n} M^\kappa dV_n = \int_{\Gamma_n} F^\kappa \cdot d\Gamma_n + \int_{V_n} q^\kappa dV_n \]  \hspace{1cm} (A1)

The integration is over an arbitrary subdomain \( V_n \) of the flow system under study, which is bounded by the closed surface \( \Gamma_n \). The quantity \( M \) appearing in the accumulation term (left hand side) represents mass per volume, with \( k = 1, \ldots, NK \) labeling the components (water, NaCl, CO\(_2\)). \( F \) denotes mass flux (see below), and \( q \) denotes sinks and sources. \( n \) is a unit normal vector on surface element \( d\Gamma_n \), pointing inward into \( V_n \). Eq. (A1) expresses the fact that the rate of change of fluid mass in \( V_n \) is equal to the net inflow across the surface of \( V_n \), plus net gain from fluid sources.

The general form of the accumulation term is:

\[ M^\kappa = \phi \sum_\beta S_\beta \rho_\beta X_\beta^\kappa \]  \hspace{1cm} (A2)

In Eq. (A2), the total mass of component \( \kappa \) is obtained by summing over the fluid (= liquid, gas) and solid phases \( \beta \). \( \phi \) is porosity, \( S_\beta \) is the saturation of phase \( \beta \) (i.e., the fraction of pore volume occupied by phase \( \beta \)), \( \rho_\beta \) is the density of phase \( \beta \), and \( X_\beta^\kappa \) is the mass fraction of component \( \kappa \) present in phase \( \beta \).

Advective mass flux is a sum over phases,

\[ F^\kappa = \sum_\beta X_\beta^\kappa F_\beta \]  \hspace{1cm} (A3)

and individual phase fluxes are given by a multiphase version of Darcy’s law:

\[ F_\beta = \rho_\beta \cdot u_\beta = -k_r \frac{k_r \rho_\beta}{\mu_\beta} \cdot (\nabla P_\beta - \rho_\beta \cdot g) \]  \hspace{1cm} (A4)

Here \( \mu_\beta \) is the Darcy velocity (volume flux) in phase \( \beta \), \( k_r \) is absolute permeability, \( k_{r\beta} \) is relative permeability to phase \( \beta \), \( \mu_\beta \) is viscosity, and \( P_\beta \) (the fluid pressure in phase \( \beta \)) is the sum of the pressure \( P \) of a reference phase (usually taken to be the gas phase), and the capillary pressure \( P_{r\beta} \) \( (\leq 0) \). \( g \) is the vector of gravitational acceleration. TOUGH2 simulator also considers diffusive fluxes in all phases, and includes coupling between diffusion and phase partitioning that can be very important for volatile solutes in multiphase conditions (Pruess, 2002). Diffusive flux of component \( \kappa \) in phase \( \beta \) is given by

\[ f_\beta^\kappa = -\phi \tau_0 \tau_\beta \rho_\beta d_{\beta\kappa} \nabla X_\beta^\kappa \]  \hspace{1cm} (A5)

where \( \tau_0 \tau_\beta \) is the tortuosity which includes a porous medium dependent factor \( \tau_0 \) and a coefficient that depends on phase saturation \( S_\beta \), \( t_\beta = t_\beta(S_\beta) \), and \( d_{\beta\kappa} \) is the diffusion coefficient.
of component $\kappa$ in bulk fluid phase $\beta$. Special TOUGH2 versions that include a conventional Fickian model for hydrodynamic dispersion have also been developed, but this option is not activated in the present study (see § 4.2 for more explanations).

By applying Gauss’ divergence theorem, equation (A1) can be converted into the following Partial Differential Equation (PDE):

$$\frac{dM^\kappa}{dt} = -\text{div} \ F^\kappa + q^\kappa \quad (A6)$$

which is the form commonly used as the starting point for deriving finite difference or finite element discretization approaches. However, in TOUGH2 an “integral finite difference” method (Narasimhan and Witherspoon, 1976) is used, in which space discretization is directly applied to the integrals in Eq. (A1).

The calculation of thermophysical properties for water/NaCl/CO$_2$ mixtures appearing in the equations given above is described in the ECO2n user’s guide (Pruess, 2005).

**Appendix B : Details about precipitation/dissolution of sodium chloride**

The TOUGH2 code includes several Equation of State (EOS) modules developed specifically for treating different mixtures in order to solve problems in the fields of hydrogeology, geothermal and petroleum engineering, nuclear waste disposal, and environmental pollution (Pruess, 1999). The ECO2n EOS module (Pruess, 2005) was developed to simulate flow problems in which the transport of a variable salinity brine and a Non Condensible Gas (CO$_2$) occurs. The multiphase system is assumed to be composed of three mass components: water, sodium chloride, and carbon dioxide. Whereas water and CO$_2$ components may be present only in the liquid and gas phases, the salt component may be dissolved in the liquid phase or precipitated to form a solid salt phase.

The treatment of precipitation/dissolution of sodium chloride has been introduced using the method employed to treat similar phenomena occurring for water-silica mixtures (Verma and Pruess, 1988). Dissolved salt concentration is characterized in ECO2n module by means of a specific primary variable $x_{\text{NaCl}}$ (fraction of total H$_2$O+NaCl mass). This variable is restricted to the range $0 \leq x_{\text{NaCl}} \leq x_{\text{eq}}$, where $x_{\text{eq}}$ is the solubility of salt according to temperature (Potter et al. 1977, Chou 1987):

$$x_{\text{eq}} = 0.26218 + 7.2 \times 10^{-5}T + 1.06 \times 10^{-6}T^2 \quad (B1)$$

with $T$ in °C. When salt concentration ($x_{\text{NaCl}}$) exceeds salt solubility ($x_{\text{eq}}$), this corresponds to conditions in which solid salt will be present in addition to dissolved salt in the aqueous phase. The mass of precipitated salt is then calculated and used to evaluate the reduction of rock porosity, as well as the related formation permeability, according to relationships given in the text (see § 4.1).
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Figure 1
Figure 2
Figure 3

Molar water fraction ($y_{\text{H}_2\text{O}}$) vs. Pressure (bar)

- CO2
- N2
- Raoult Law
Figure 4

Graph showing the relationship between water saturation and outlet gas flow rate under different pressure drops (ΔP) of 0.12 bars, 0.80 bars, and 1.50 bars. The graph includes data points for mean measured water saturation and calculated water saturation, as well as measured and calculated outlet gas flow rates.
Figure 5
Figure 6
Figure 7
Figure 8
Figure 9

(a) 

Mean injection flow rate (kg/s/m)

0.0 0.2 0.4 0.6 0.8 1.0

0.0 0.1 0.2 0.3

$k_{red} = K/K_0 (x = 0)$

10 g/L
100 g/L
160 g/L

(b) 

Peclet number

0 50 100 150

Gas flow rate (kg/s)

0.00 0.01 0.02 0.03 0.04 0.05 0.06

0 500 1000 1500 2000

Brine salinity (g/L)

Low capillary fluxes = decrease of clogging risks

High capillary fluxes = salt precipitation and clogging risks
Figure 10
Figure 11
Figure 12