Impact of karstification in trapping mechanisms of CO₂ storage

Sharidah Mohd Amin1 · Raj Deo Tiwari1 · Ana Widyanita1 · Prasanna Chidambaram1 · Siti Syareena Mohd Ali1 · Abd Hakim Mazeli1 · Chee Phuat Tan1 · M. Khaidhir A. Hamid1

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Abstract
A carbonate-depleted carbonate gas field located offshore Sarawak has been identified as potential candidate for CO₂ sequestration site in conjunction with another high CO₂ gas field development for commercialization efforts. The field has undergone a feasibility study to evaluate potential geochemical reactions specifically on trapping mechanisms and storage capacity associated with CO₂ injection. A detail 3D reactive transport modelling study was conducted to quantify on the four different trapping mechanism: structural, residual, solubility, and mineral trapping during and post-injection. The model was developed by first converted the available field history matched black oil simulation model into compositional 3D model, in which CO₂ is treated as separate component in the reservoir through the production and injection processes. The study covered 22 years of gas production history forecast followed by 27 years of injection and 1000 years of post-CO₂ injection in the gas column reservoir. The results show that the field has potential to store and sequestrate CO₂ up to for 79% structural trapping, 19% residual trapping, 3% solubility trapping and no mineral trapping after 1000 years of post-injection period.

Keywords Carbonate gas field · CO₂ storage · Geochemical reactions · Trapping mechanisms · 3D reactive transport modelling · Karstification

Introduction
A storage in depleted gas reservoirs is one way to reduce the increasing greenhouse gases in the atmosphere. Any large scale of CO₂ injection involves a variety of coupled physical and chemical processes including multiple fluid flows, solute transport, and chemical reactions between the resident fluids and reservoir minerals. To date, most of previous studies has focused on the reactivity and trapping mechanisms during post-injection of several thousand years (Gallo et al. 2006; Lindeberg and Wessel-Berg 1997, 2002, 2003; Liu et al. 2012; Pruess and García 2002). However, none of the previous study has discussed about the impact of karstification in the carbonate field. Thus, this study focuses on detailed geochemical impact in terms of trapping mechanisms due to the presence of karstification in the carbonate field. The main objectives of this study are:

- To obtain on the kinetic rate parameters from laboratory-calibrated model for 3D coupled model.
- To study on the short- and long-term reactions due to CO₂ injection in carbonate reservoir.
- To obtain on the CO₂ storage trapping mechanism during and post-injection of CO₂ in carbonate reservoir.
- To study on plume migration of CO₂ in the presence of heterogeneity in the carbonate reservoir.

Background
A carbonate build-up platform in the Central Luconia Province is located 260 km NW of Bintulu. The field was discovered in 1970, when X-1 well was drilled and appraised by X-2 well in 1991. Structurally, the field is a high relief carbonate build-up and forms part of the Mega Platform complex comprised of other two carbonate fields with a common regional aquifer (Fig. 1).

The main carbonate reef build-up for field is concentrated in the south east due to the prevailing SE-NW paleowind direction at the time of deposition. The depositional environments in this location are the rim-reef cores,
fore-reef to back-reef, where generally the primary porosity properties are good. Platform progradation and sediment shedding occurred in the downwind position towards the NW, where the depositional environment is mainly lagoonal. Generally, the reservoir properties are believed to deteriorate in this area. However, due to diagenesis, much of the porosity being attributed to secondary porosity enhancement, making the lagoonal reservoir properties superior compared to in the reef facies.

The reservoir porosity in the gas zone ranges from 15 to 32%, while the core permeability derived from well ranges between 9 and 200 mD. Production test interpretations indicate higher permeability of 200–480 mD probably due to karsts or fractures. The field has a strong underlying aquifer support.

### Materials and methodology

#### Geochemical data

The system was modelled using the initial formation brine composition from analogue field as in Table 1 and the primary mineral assemblage from the study field as input data. (Table 2).

#### Kinetic batch modelling

Prior to simulating reactive transport for the carbonate reservoir, a batch geochemical model of water–rock interaction was performed to generate an aqueous-phase chemical composition approaching the composition of typical formation brine. This was done by equilibrating the initial formation brine composition as in Table 2 with the presence of the primary minerals at depth of 6079ft as listed in Table 1.

The initial brine composition was equilibrated with the mineralogy of the study field mineralogy (Table 2), and the resulting brine composition (Table 3) was used as a starting point for subsequent modelling in this study. From XRD analysis from the selected well in Table 1, the mineralogy composition (in average wt%) used in the model are as follows:

- Calcite–71.0 wt % to 92.9 wt % (average 83.8 wt %)
- Dolomite–0.4 wt% to 2.7 wt % (average: 1.8 wt%)
- Siderite–0.8 wt% to 1.0 wt% (average 0.9 wt%)
- K-feldspar–1.1 wt%—4.9 wt % (average: 2.4 wt%)
- Quartz–< 1.5 wt% (average: 0.5 wt%)
- Clay minerals–< 12.5 wt % (average: 6.6 wt%)

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**Table 1** Initial formation brine taken from analogue carbonate field

| Component | Concentration (mg/L) |
|-----------|----------------------|
| Na⁺       | 8996.5               |
| Ca²⁺      | 233.3                |
| Mg²⁺      | 47.8                 |
| Fe³⁺      | 0                    |
| K⁺        | 816.4                |
| Cl⁻       | 13,812.8             |
| SO₄²⁻     | 1275.6               |
| HCO₃⁻     | 1363                 |
| Well name | Sample depth | Calculated whole rock composition (Weight %) | Clay fraction (clay typing, % wt) |
|-----------|--------------|---------------------------------------------|----------------------------------|
|           |              | Quartz Plagioclase K-Feldspar Calcite Dolomite Siderate Pyrite Total clay Total | Kaolinite Chlorite Illite Mixed layer I/S Smectite |
| 5695.3L   | 0.3          | 2.2                                         | 2.1                               | 86.3 0.5 0.8 1.7 6.2 100 | 14.7 5.1 59.6 15.9 5.1 |
| 5695.3R   | 0.3          | 1.9                                         | 1.8                               | 88.6 0.5 0.8 1.6 4.6 100 | 15.2 5.9 58.1 16.0 4.8 |
| 5699.8    | 0.7          | 0.6                                         | 1.4                               | 92.9 0.5 1.0 0.0 2.9 100 | 18.3 11.1 50.2 15.5 4.5 |
| 5726      | 0.3          | 2.3                                         | 2.3                               | 83.0 0.8 0.8 1.9 6.9 100 | 13.9 7.1 56.3 17.4 5.0 |
| 5749      | 0.9          | 4.4                                         | 4.9                               | 71.0 2.7 1.0 2.6 12.5 100 | 17.6 9.6 54.9 13.3 4.6 |
| 5772.1    | 0.5          | 3                                           | 2.8                               | 81.9 2.1 1.0 1.3 7.5 100 | 13.8 6.1 60.8 14.4 5.1 |
| 5773.8    | 0.3          | 1.7                                         | 1.1                               | 89.2 1.5 0.9 1.2 4.1 100 | 13.5 7.2 59.2 14.9 4.7 |
| 5773.9    | 0.3          | 2.2                                         | 1.8                               | 85.0 2.7 0.9 1.5 5.6 100 | 14.2 7.6 57.4 16.0 4.6 |
| 5774.5    | 0.5          | 2.0                                         | 1.9                               | 83.5 1.9 0.9 2.4 6.9 100 | 12.2 6.8 62.6 13.8 4.9 |
| 5787      | 0.4          | 2.6                                         | 2.7                               | 82.1 1.4 0.8 2.1 8.0 100 | 15.5 7.8 55.0 16.5 5.4 |
| 5811.67L  | 0.8          | 2.5                                         | 3.2                               | 83.1 0.4 1.0 2.0 7.1 100 | 13.1 4.4 64.1 13.3 5.2 |
| 5811.67R  | 1.5          | 2.8                                         | 4.4                               | 76.9 0.5 1.1 2.8 9.8 100 | 18.7 5.7 54.8 15.2 5.9 |
| 5839.4    | 0.5          | 1.9                                         | 1.8                               | 85.0 3.0 0.8 1.2 5.9 100 | 12.8 7.7 53.7 20.7 4.8 |
| 5839.6    | 0.5          | 2.5                                         | 4.3                               | 80.0 3.5 1.0 2.0 6.2 100 | 12.2 6.0 60.9 16.2 4.9 |
| 5840.6    | 0.5          | 2.1                                         | 1.6                               | 84.9 2.4 0.9 1.4 6.4 100 | 11.7 5.9 63.3 14.7 4.8 |
| 5867.8    | 0.5          | 2.2                                         | 1.7                               | 83.0 2.7 0.8 1.8 7.2 100 | 12.8 6.8 56.4 18.9 5.1 |
| 5870L     | 0.6          | 3.8                                         | 4.4                               | 74.5 2.8 1.0 2.3 10.6 100 | 14.3 6.2 55.2 19.1 5.1 |
| 5870R     | 0.3          | 2.5                                         | 3.1                               | 80.0 3.7 0.8 2.3 7.2 100 | 13.8 7.7 53.7 19.9 5.3 |
| 5888.4    | 0.4          | 2.2                                         | 1.8                               | 85.3 1.3 0.9 1.6 6.4 100 | 12.2 6.5 60.2 16.2 5.2 |
| 5889.7    | 0.4          | 1.9                                         | 1.4                               | 87.3 1.0 0.8 1.3 5.8 100 | 12.0 5.7 61.7 15.2 5.0 |
| 5890      | 0.4          | 2.2                                         | 1.9                               | 86.7 0.9 0.8 1.6 5.4 100 | 14.5 7.9 59.6 12.6 4.9 |
| 5903.7    | 0.4          | 1.9                                         | 1.7                               | 86.0 1.9 0.9 1.5 5.6 100 | 13.2 6.9 56.5 18.1 5.0 |
| 5908      | 0.5          | 2.3                                         | 1.9                               | 84.7 1.4 1.0 1.6 6.8 100 | 12.9 6.8 55.7 19.4 5.2 |
| 5909.8    | 0.5          | 2.0                                         | 2.2                               | 85.7 1.2 0.9 1.5 5.9 100 | 12.5 7.0 55.3 19.8 5.1 |
| 5948.6    | 0.4          | 2.3                                         | 3.1                               | 84.9 1.3 0.8 1.8 5.4 100 | 12.6 7.0 56.2 17.9 6.1 |
| 6024.56   | 0.6          | 4.0                                         | 5.8                               | 72.7 2.6 1.1 2.5 10.6 100 | 17.7 1.2 55.8 18.6 6.5 |
| 6039.68   | 0.5          | 2.4                                         | 2.2                               | 83.9 0.6 0.8 1.8 7.7 100 | 14.3 5.8 54.7 19.9 5.1 |
| 6079      | 0.4          | 2.2                                         | 2.9                               | 85.3 2.0 1.0 1.5 4.8 100 | 1.39 6.7 54.9 19.2 5.0 |
| 6079.4    | 0.4          | 1.7                                         | 1.6                               | 8.7 2.4 0.9 1.2 4.1 100 | 14.2 7.3 56.3 17.0 4.9 |
| 6085      | 0.5          | 2.2                                         | 1.4                               | 87.2 0.7 0.8 1.3 5.9 100 | 13.2 6.0 56.9 18.2 5.3 |
After the equilibration of the formation water and mineralogy, a total of eight aqueous components were used as initial formation water in the model (Table 3). Simulations were for three-dimensional reactive transport modelling and was simulate at iso-thermal conditions of 275°F and average reservoir pressure of 2538 psi. Since the reservoir is dominantly dominated by carbonates, the model was simplified using calcite (0.90 of volume fraction) and dolomite (0.10 of volume fraction) composition. The model systems chosen were the initial formation water composition, and the primary mineral assemblage consists of Calcite (Volume fraction: 90%) and Dolomite (Volume fraction: 10%) from the aquifer. Laboratory-calibrated model was used to obtain on kinetic rate and reactive surface areas parameters for the field.

### Kinetic rates

Dissolution and precipitation of minerals in the system were kinetically controlled. The rates of mineral dissolution and precipitation rates were derived from (Lasaga et al. 1995):

\[
rate_m = A_m k(T)_m (a_{H^+})^n [1 - (Q_m/K_m)]
\]  

where the subscript m is the mineral index, and ratem is the dissolution/precipitation rate (m/s), Am is the reactive surface per kg water (unit), k(T) is the temperature dependent rate constant, aH+ is the proton activity, n is the order of the reaction (0 ≤ n ≤ 1), Km is the equilibrium constant for the mineral water reaction written for the dissolution of 1 mol of mineral and Qm is the reaction quotient. The last term in Eq. 1 takes into account that the rate of reaction depends on how far the system is from chemical equilibrium, that is, the degree of over- or under-saturation. The rate constants in Eq. 2 for all minerals were extrapolated to the field condition temperatures of 348.15 K from reported rate constants at 298.15 K using the Arrhenius relation:

\[
k(T) = k_{25} \exp \left[-E_a/R(1/T - 1/298.1)\right]
\]

where \(E_a\) is the activation energy (J/mol), \(k_{25}\) is the rate constant at 250 °C (mol/m²s), R is the universal gas constant (8.314 J/mol K) and T is the temperature (K). The rate constants at 298.15 K \((k_{25})\) and parameters for the calculation of reaction rates for all minerals were taken from the laboratory-calibrated model as discussed in the subsequent section.

Precipitation of secondary minerals depends on nucleation, Ostwald ripening, crystal growth processes, and reactive surface (Plummer 1978; Steefel and Cappellen 1990). Due to coating, interaction with the minerals is generally expected to occur only at selective sites of the mineral surface and the actual reactive surface area could be between one and three orders of magnitude less than the surface roughness based surface area (Lasaga et al. 1995). To account for these effects, a surface roughness factor of 10 was incorporated and defined as the ratio of the true (BET) surface area to the equivalent geometric surface area.

### Reactive surface areas

The composition of the mineral assemblages was taken from samples M3 carbonate field. The reactive surface areas for calcite and dolomite were calculated using geometrical calculation method due to unavailability of BET measurement in the laboratory (Table 4). Reactive surface areas were calculated using geometrical calculation—based on assumption of idealized geometry of mineral grains (spherical grains with grain diameters of 3.3 × 10⁻⁵).

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### Table 3 Initial formation water for laboratory-calibrated model

| Parameters          | Values          |
|---------------------|-----------------|
| Temperature         | 275°F           |
| Pressure            | 2538            |
| pH                  | 8.2             |
| Ionic Strength      | 0.4             |
| Activity Coefficient Model | Extended Debye Huckel (Truesdell and Jones 1973) |

| Elements               | Concentration (mg/L) |
|------------------------|----------------------|
| Na⁺                    | 70                   |
| Ca²⁺                   | 29                   |
| Mg²⁺                   | 116                  |
| Fe²⁺                   | 9178                 |
| K⁺                     | 12,626               |
| Cl⁻                    | 532                  |
| SO₄²⁻                  | 1000                 |
| HCO₃⁻                  | 0                    |

### Table 4 The reactive surface areas for calcite and dolomite

| Minerals | Chemical formula | Reactive surface area (m²/g) | Initial mineral composition (volume fraction) |
|----------|------------------|-----------------------------|---------------------------------------------|
| Calcite  | CaCO₃            | 6.71 × 10⁻²                 | 0.90                                        |
| Dolomite | CaMg(CO₃)₂       | 6.35 × 10⁻²                 | 0.10                                        |
Three-dimensional (3D) reactive modelling

Reactive transport model was taken from static model of 300 thickness with three (3) CO₂ injectors well and ten (10) gas producers. The model consists of 117,024 grid blocks with active cells of 91,475. Since the model was generated from static model, the model is heterogeneous in terms of the rock properties (porosity and permeability). The static model is then converted into static model in CMG before the model was run with geochemistry inputs (initial formation water, minerals composition, kinetic rate, and reactive surface areas) for geochemical modelling study. The simulation was run for 27 years of injection period and post-injection of 1,000 years.

Results and discussion

The results from kinetic batch modelling (laboratory-calibrated model) and three-dimensional reactive transport modelling for the study carbonate field are presented in this section.

Kinetic batch laboratory-calibrated model

The study used calibrated experimental batch model to analyse and interpret CO₂-injection-induced carbonate dissolution phenomena from pore to score scale. The same approach on using laboratory-calibrated models to obtain on the kinetic rate and reactive surface areas parameters for full field/3D reactive transport modelling has been used by several authors in the literature to study on the geochemical reactions due to CO₂ injection in reservoirs.

Static ageing experiment was conducted in the laboratory to evaluate on the effects of CO₂ injection on the kinetic reaction and mineralogy changes after 45 days (Fig. 2). The sample used for laboratory-calibrated model is the ones from aquifer zone, which is at depth 6079 ft. The sample is submerged into synthetic brine with supercritical CO₂, SCO₂, injection under reservoir pressure and temperature (2900 psi and 275°F).

The changes of main components ((Ca²⁺, Mg²⁺, K⁺, Na⁺) of formation brine components were measured using Inductively Coupled Plasma (ICP) measurement. The increases in Ca²⁺ as shown in Table 5 indicate calcite dissolution due to CO₂ injection. The pre- and post-sample analysis on porosity changes are conducted using Digital Core Analysis (DCA), where the results shown very insignificant changes in porosity, which is 0.47%. In addition, XRD analysis shown very minor changes in calcite mineral and other minerals, which is less than 0.5%.

The calibrated models using static ageing experiments conditions are developed to get kinetic rate and reactive surface area input data for full field/3D reactive transport modelling. The input parameters model for calibrated model used are shown in Table 6. The reactive surface area is calculated using geometrical calculation, and it is fixed in the model. Rate constants were adjusted to fit the model with the experimental data to match with Ca²⁺ changes in concentration for pre- and post-experiments. However, the laboratory data for ICP measurement on Ca²⁺ concentration changes is insufficient as there is only two data points to be matched (Fig. 3). The resulting kinetic rates for calcite and dolomite obtained from the calibrated models are shown in Table 7.

Table 5 ICP analysis results for main components for pre- and post-static ageing experiment

| Components | Unit | Pre-concentration | Post-concentration |
|------------|------|-------------------|--------------------|
| Ca²⁺       | mg/L | 70                | 271.5              |
| Mg²⁺       | mg/L | 29                | 24.5               |
| K⁺         | mg/L | 116               | 97.45              |
| Na⁺        | mg/L | 9178              | 8976               |
| Cl⁻        | mg/L | 12,636            | –                  |
| HCO₃⁻      | mg/L | 532               | –                  |
| SO₄²⁻      | mg/L | 1000              | –                  |
| Fe₂⁺       | mg/L | 0                 | 0.012              |
3D reactive transport modelling

Short-term fate (27 years of injection to 100 years of post-injection)

Gas saturation After the injection, the vertical flow of CO₂ rich gas is influenced by buoyancy. This is due to the difference in the density and viscosity of CO₂-rich gas and brine. The smaller viscosity of CO₂-rich gas compared to brine causes hydrodynamic instabilities. Eventually, counter current flow occurs as the gas flows upward and the brine downward. The profile of CO₂ mole fraction in the model domain during 27 years of injection and 100 years of post-injection is plotted in Figs. 4 and 5. The CO₂ plume size is illustrated using gas mole fraction of CO₂. The significant gas front saturation phase of injected CO₂ propagates laterally to the West of the injection well giving the high concentration of CO₂ saturation layer in gas zone.

Changes in pH Following the injection of CO₂ into the system, the pH immediately decreased from 7.4 to 6.3 and thereafter remain the same until end of injection period. The initial fugacity, fCO₂ controls this available CO₂ in the system and consequently the evolution of the pH over the course of the reaction. The dissolution and precipitation of the minerals, discussed below

Table 6 Input parameters model for calibrated model

| Parameters                  | Values |
|-----------------------------|--------|
| Temperature                 | 135    |
| Pressure                    | 2538   |
| pH                          | 8.2    |
| Ionic Strength              | 0.4    |
| Activity Coefficient Model  | Extended Debye Huckel (Truesdell and Jones 1973) |

| Elements   | Concentration (mg/L) |
|------------|----------------------|
| Na⁺        | 9178                 |
| Ca²⁺       | 70                   |
| Mg²⁺       | 29                   |
| Fe²⁺       | 0                    |
| K⁺         | 116                  |
| Cl⁻        | 1263.6               |
| SO₄²⁻      | 1000                 |
| HCO₃⁻      | 532                  |

Fig. 3 Laboratory-calibrated model plot. The kinetic batch model was developed based on static ageing experiment. The model was fit using trying and error method from Ca²⁺ concentration data obtained before CO₂ injection (initial formation brine) and after 45 days of reactions of static ageing experiment

Table 7 Laboratory-calibrated model input data for 3D full field reactive transport modelling

| Mineral   | Volume fraction | Surface Area (m²/g) | Log₁₀k (mol/m².s) | Activation Energy, Ea (kJ/mol) | Reference Temperature (°C) |
|-----------|-----------------|---------------------|-------------------|-------------------------------|-----------------------------|
| Calcite   | 0.90            | 6.71 × 10⁻²         | −3.84             | 25.7                          | 25                          |
| Dolomite  | 0.10            | 6.35 × 10⁻²         | −5.83             | 59.9                          | 25                          |
Fig. 4 Gas saturation after 27 years of injection. The spatial distribution of CO₂ dissolved in aqueous phase at time of simulations given in terms of mole fractions.

![Graph showing CO₂ distribution](image1.png)

GWC: 5880 ft

End of 27 years of injection

Fig. 5 Gas saturation after 100 years of injection. The spatial distribution of CO₂ dissolved in aqueous phase at time of simulations given in terms of mole fractions.

![Graph showing CO₂ distribution](image2.png)

GWC: 5880 ft

100 years of post-injection

Fig. 6 pH changes near to one of CO₂ injector well before and during CO₂ injection period of 27 years. There is a drop in pH when CO₂ injection starts in 2026.

![Graph showing pH changes](image3.png)

pH - {51, 28, 5}

pH change from 1995 to 2052
(Fig. 6), reflect the consumption of the acidic CO₂ by 
CO₂-brine-mineral reactions. These geochemical reactions 
result in an increasing pH at later period. A higher fco₂ increases the amount of dissolved CO₂ in the brine 
and as a result increase the extent of mineral dissolution. 
Minerals such as calcite dissolve rapidly, leading to an 
increase in pH.

**Mineral dissolution/precipitation** Buffering occurs when dissolved CO₂ reacts, for instance, with reactive carbonates such as calcite and dolomite (Reaction 1 until 3). These reactions may provide sufficient buffering capacity (via HCO₃⁻ alkalinity) to resist drastic changes in pH. The presence of reactive carbonates such as calcite in a host reservoir will have a major impact on how the chemical reactions evolve during injection of CO₂. If CO₂ dissolved into brine is in equilibrium with carbonate minerals, the total amount of dissolved inorganic carbon (CO₂(aq), HCO₃⁻(aq), CO₃²⁻(aq)) in the host reservoir.

**Reaction 1**

\[ \text{H}_2\text{O} + \text{CO}_2 \leftrightarrow \text{H}_2\text{CO}_3 \leftrightarrow \text{H}^+ + \text{HCO}_3^- \]

**Reaction 2:**

\[ \text{Calcite} \quad \text{CaCO}_3 + \text{H}^+ \leftrightarrow \text{Ca}^{2+} + \text{HCO}_3^- \]

**Reaction 3:**

\[ \text{Dolomite} \quad \text{CaMg(CO}_3)_2 + \text{H}^+ \leftrightarrow \text{Ca}^{2+} + \text{Mg}^{2+} + \text{HCO}_3^- \]

After 27 years injection of CO₂ into the aquifer, the super-
critical fluid is in contact with the residual brine in the gas

column. As a result, it forms H₂CO₃, a weak acid that almost 
immediately dissociates into hydrogen and bicarbonate. The 
low pH during injection of 27 years with CO₂ causes the 
brine to become highly under saturated with respect to car-
bonates especially calcite present in the carbonate reservoir. 
The main carbonates that dissolve progressively are calcite. 
The amounts are multiplied in the injected plume or at edges of the plume, where gradients in pH is 
greatest. The corresponding changes in calcite volume frac-
tion are shown in Fig. 7. From the figure, zone above the 
Gas Water Contact (GWC) which the CO₂ injector located 
shows very small changes in terms of calcite dissolution. For 
zone below GWC, there is no dissolution of calcite occurs 
due to the absence of CO₂ in this zone. There is a thin layer 
of high dissolution zone above the GWC of the aquifer. This 
is the plume region, with a gradient to lesser dissolution 
down this boundary during short term of reactions (less 
than 100 years).

**Changes of porosity** Temporal changes in porosity due to 
mineral dissolution and precipitation can affect fluid flow. 
The dissolution of minerals such as calcite changes the for-
mation porosity. Then, changes in this property cause the 
changes in the fluid flow patterns. When this occurs, there 
is an additional pore space to accommodate brine and CO₂. 
The changes of porosity were calculated from changes in 
volume fractions of minerals caused by the minerals reac-
tions at each time step of running simulations according to 
Eq. 3 below:

\[ \Delta \theta_i = \left(1 - \frac{\sum_j n_{ij} V_{ij}}{V_{\text{total}}} \right) - \theta_i = 0 \quad (3) \]

![Quick Plot - Mineral(Calcite)](image)

**Fig. 7** Calcite volume changes near to one of CO₂ injector well before and during CO₂ injection period of 27 years. There is dissolution of calcite occurs when CO₂ injection starts in 2026

\[ \sum \text{ Springer} \]
where, \( \theta_{\text{in}} \) is the initial porosity, \( n_{i,t} \) is the moles of dissolved minerals \( i \) at time of simulations, \( V_{i,t} \) is the molar volume of dissolved minerals \( i \) at time of simulations and \( V_{\text{total}} \) is the total initial minerals volume of the system.

The changes in porosity affect permeability. Small increases in porosity will increase the permeability. This creates a migration pathway especially at the top of carbonate (cap rock) for CO\(_2\) into the surface. Thus, an assessment of the change in porosity is very crucial to understand the migration pathway of CO\(_2\) plume in the reservoir.

All the above reactions impact the porosity by changing of mineral volume due to dissolution and precipitation of minerals. In reservoir above the GWC zone, the delta porosity is very insignificant, which is with 0.03%. Zone below the GWC shows no changes in porosity due to the absence of CO\(_2\) in this zone during CO\(_2\) injection period of 27 years until end of 100 years of post (Figs. 8 and 9).

**Long-term fate study (100–1000 years post-injection injection)** Upon CO\(_2\) injection into the reservoir, the supercritical CO\(_2\) migrates upward quickly due to the density differences. As a result, most of the supercritical CO\(_2\) accumulated in this zone (Fig. 10). Over the years, the CO\(_2\) dissolved, and the dissolved plume migrates downward due to the larger density of brine with CO\(_2\) compared with the brine without CO\(_2\). It was reported in Ennis King and Paterson, 2003 that the larger density of brine with CO\(_2\) is approximately 10 kg/m\(^3\) higher compared with the brine without CO\(_2\). The brine containing dissolved CO\(_2\) in carried downward and is subsequently replaced by brine with less CO\(_2\). This process is known as convective.
It is acknowledged that the role of convection mixing is crucial during the long-term CO₂ dissolution (Ennis-King and Paterson 2005; Linderberg 2002). This phenomena can be seen in Fig. 11, where at 300 years, the dissolve CO₂ started to have the push-down effect. The denser CO₂-saturated brine moves downward due to higher
Fig. 13 Gas saturation after 1000 years of post-injection. CO₂ plume migration migrates laterally to the west of the reservoir due to the presence of impermeable layer at the east zone of the reservoir.

Fig. 14 Impact of karstification in the carbonate reservoir on the convection/fingering effect at top of the reservoir.
permeability zone due to karstification in the host reservoir (Figs. 12, 13, 14). The properties of the model were populated from the static model as in Fig. 14. At the end of 1000 years post-injection, the dissolved CO₂ is dominant in the aquifer zone. The same observation being made by Agartan et al. (2015). At the end of 1000 years post-injection, the dissolved CO₂ is dominant in the aquifer zone. However, due to the low permeability zone at the East part of the reservoir, there is no downward movement of heavily denser CO₂ (Fig. 14). Several other authors that observed the same findings which reported that vertical heterogeneity impact on the solubility trapping of CO₂ in the reservoir are tabulated in Table 8.

**Changes in pH, minerals, and porosity** The geochemical evolution for long-term reactions is shown in Figs. 16, 17 plotting pH, changes in mineral abundance and porosity against time. The reactivity of long-term reactions is best interpreted using the evolution of pH. The evolution of pH is related to the amount of CO₂ dissolved in the brine. The evolution of pH and minerals changes for long-term reactions is shown in Figs. 15 and 16. It was no significant changes in the pH and mineral volume. In addition, there is very insignificant changes in porosity for 1000 years of post-injection, which is 0.0021% in porosity increase (Fig. 17).

**Total amount of sequestered CO₂** The total amount of CO₂ stored in aqueous, gas, and solid phases over time during the injection period of 21 years in the reservoir is shown in Table 9. From this table, approximately 79% of injected supercritical CO₂ remains in gas phase over 27 years of injection. However, over long timescale the amount of supercritical (own phase), CO₂ decreases and CO₂ in the dissolved brine increases. The total amount of CO₂ sequestered is calculated by the changes in moles of carbon inventory multiplied by the molecular weight.

**Conclusions** This study set out to explore the effects of injected CO₂ on geochemical reactions and fluid dynamics over geological timeframe of 1,000 years in the carbonate depleted gas reservoir. Kinetic batch modelling/laboratory-calibrated model was developed to obtain on kinetic rates and reactive surface areas that were used as an input data for 3D reactive transport modelling.

The main outcomes of the study are:

### Table 8 List of literature and key findings on effect of vertical heterogeneity/permeability on trapping mechanism

| Scope/Aspect | References | Key Findings |
|--------------|------------|--------------|
| Analytical simulations | Ennis-King and Paterson (2005). Role of convective mixing in the long-term storage of carbon dioxide in deep saline formations; SPE J. 10, 349–356 |
| | Ennis-King and Paterson (2007). Coupling of geochemical reactions and convective mixing in the long-term geological storage of carbon dioxide; Int. J. Greenhouse. Gas Control 1, 86–93 |
| | In the long term, the dominant mechanism for dissolution of CO₂ in brine is convective mixing rather than diffusion due to density instability. The time needed for the injected gas to dissolve completely is typically on the order of hundreds of years to tens of thousands of years, depending on the vertical permeability |
| Laboratory experiments | Agartan et al. (2015). Experimental study on effects of geological heterogeneity in enhancing dissolution trapping of supercritical CO₂, Water Resour. Res., 51, 1635–1648 |
| | In the presence of randomly distributed permeability zones, convective mixing occurs through the high-permeability connected pathways. Dissolution in formation water at a pH of 5–5.4 is the sole mechanism for CO₂ in silicate reactions. Carbonate reservoirs are more effective for CO₂ sequestration due to their higher densities and solubility |
| Natural analogues | Gilfian et al. (2009). Solubility trapping in formation water as a dominant CO₂ sink in enhancing dissolution trapping of supercritical CO₂, Water Resour. Res., 45, 14–36. |
| | Dissolution in formation water at a pH of 5–5.4 is the sole major sink for CO₂ in silicate reactions. Carbonate reservoirs are more effective for CO₂ sequestration due to their higher densities and solubility |
| Numerical simulations | Farajzadeh et al. (2007). Numerical simulation of density-driven natural convection in enhancing dissolution trapping of supercritical CO₂, Int. J. Heat Mass Transfer. 50, 6691–6700 |
| | At moderate heterogeneity, the flow is dominated by the permeability field structure, i.e., channel formation and CO₂ plumes progress along the high-permeability streaks |
| | Farajzadeh et al. (2011). The effect of heterogeneity on the character of density-driven natural convection of CO₂ overlying brine layer, Adv. Water Resour., 34, 327–339 |
Reactive transport modelling indicates that the geochemical processes in the reservoir is dominated by 79% structural trapping, follow by 19% residual and 3% solubility trapping.

The CO₂ inventory for post-injection until 1000 years is tabulated in Table 9, which the solubility trapping is observed to increase after 1000 years of reactions.

The results indicate that extent of trapping mechanisms and CO₂ plume migration are highly dependent on vertical reservoir heterogeneity which is the presence of karstification in the reservoir.

Limitation and recommendation for future studies are:

- The study predicts no mineral trapping as 1000 years of simulation is too short for the model to observe any
precipitation of minerals. It is anticipating that mineral trapping would take place once the model system reaches super-saturate state, and these processes are very slow which this could only potentially occur over 10,000 years of reactions. It was anticipating that the trapping of CO₂ in minerals would be very minimum and will not jeopardize on any integrity issue of the storage.

For future work, the same approach for up till 10,000 years of simulation period could be useful to assess the potential mineral trapping in carbonate depleted gas fields and its effect on integrity issues.

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Table 9 CO₂ inventory for end of 27 years of injections and post-injection of 1,000 years

| Trapping Mechanism          | End of injection (Tscf) | %       | 1000 years of post-injection (Tscf) | %       |
|-----------------------------|-------------------------|---------|------------------------------------|---------|
| Supercritical               | 0.667                   | 79      | 0.591                              | 70      |
| Residual                    | 0.158                   | 19      | 0.151                              | 18      |
| Dissolved in Water          | 0.022                   | 3       | 0.106                              | 12      |
| Mineral                     | 0.00                    | 0.00    | 0.00                               | 0.00    |
| Total                       | 0.85                    | 100     | 0.85                               | 100     |

Fig. 17 Porosity changes during 1000 years of post-injection. The simulations were first run for 27 years to account for the injection period followed by 1000 years to account for retention period. After the CO₂ injection, the dissolved CO₂ reacts with primary minerals, initially with calcite. The model was run with calcite mineralogy only as the study was simplified and calcite was chosen due to its fast kinetic rates compared to other minerals. The changes of porosity observed in the model are very small after post-injection of 1000 years.

Declarations

Conflict of Interest Not applicable.

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