Preinjection Characterisation and Evaluation of CO₂ Sequestration Potential in the Haizume Formation, Niigata Basin, Japan

Geochemical Modelling of Water-Minerals-CO₂ Interaction

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Résumé — Caractérisation avant injection et évaluation du potentiel de séquestration de CO₂ dans la formation de Haizume, bassin de Niigata, Japon. Modélisation géochimique des interactions eau-minéraux-CO₂ — L'Institut de recherche en technologie innovante pour la terre (RITE) a réalisé une expérience à petite échelle d'injection de CO₂ dans un champ, afin d'étudier la faisabilité d'une séquestration géologique de ce gaz à effet de serre au sud-ouest de la ville de Nagaoka (préfecture de Niigata, Japon). Avant cette expérience, les réactions géochimiques résultant de l'injection ont été évaluées à l'aide du modèle géochimique EQ3/6. La formation dans laquelle la séquestration est envisagée est la formation de Haizume, dans le groupe Uonoma, constituée de dépôts sédimentaires marins d'âge pléistocène, et surmontée par une couche d'argilites. La minéralogie de la formation de Haizume est constituée par le quartz, le plagioclase, le feldspath potassique, le pyroxène et les argiles (smectites et chlorites). La taille des grains varie de moyenne à grossière. Le grès montre une faible consolidation. L'eau de formation présente une teneur en sels totaux dissous d'environ 6100 mg/l et une concentration élevée en calcium (> 20 % de la concentration de Na⁺).

Le modèle géochimique a été utilisé pour ajuster la chimie initiale de l'eau de formation, puis pour modéliser les interactions entre le CO₂ injecté, l'eau de formation et les minéraux du réservoir. Les résultats indiquent une forte réactivité des minéraux en présence d'une quantité élevée de CO₂, ainsi qu'un forfait de conversion de ce CO₂ en minéraux. À l'état final, environ 23 mol de CO₂ ont été consommées par kg d'eau, dont plus de 90 % ont été précipités sous forme de minéraux carbonatés. Ces calculs de modélisation géochimique renferment une incertitude importante en ce qui concerne les temps de réaction, qui devront être évalués de façon plus précise dans des modélisations futures.
Abstract — Preinjection Characterisation and Evaluation of CO₂ Sequestration Potential in the Haizume Formation, Niigata Basin, Japan. Geochemical Modelling of Water-Minerals-CO₂ Interaction — The Research Institute of Innovative Technology for the Earth (RITE) is carrying out a small-scale CO₂ injection field experiment to investigate the feasibility of geological sequestration of CO₂ greenhouse gas in the south-west of Nagaoka City, Niigata Prefecture, Japan. Prior to the injection geochemical reactions caused by CO₂ injections were investigated using the geochemical modelling code (EQ3/6). The injection formation is the sedimentary marine Haizume Formation (Pleistocene) in the Uonuma Group, which is covered by a mudstone seal. The formation is mainly composed of quartz, plagioclase, feldspar, pyroxene, and clays (smectite, chlorite). The sandstone shows minor consolidation and grain size is medium to coarse sand. The total dissolved solid (TDS) of the formation water is approximately 6100 mg/l and the water contains a high Ca²⁺ (> 20% of Na⁺ concentration). The geochemical model was used for an initial adjustment of the formation water chemistry to the formation conditions and a modelling of the formation water-mineral-CO₂ reactions. The modelling results showed a high reactivity of the minerals in the CO₂ rich environment and high mineral conversion rate within the formation. At the final state, approximately 23 mol of CO₂ were taken into 1 kg of formation water and more than 90% of this was stored within carbonate minerals. In this simulation, some uncertainty is associated with the time scale and a more detailed investigation is planned and will address accurate evaluation.

INTRODUCTION

The Research Institute of Innovative Technology for Earth (RITE), Japan is conducting research to develop and establish new technology for long term geological sequestration of CO₂ in Japan. In this context, RITE is carrying out a small-scale CO₂ injection field experiment to investigate the feasibility of geological sequestration of CO₂ greenhouse gas in the south-west of Nagaoka City, Niigata Prefecture, Japan (NEDO, 2002). The injection Zone 2 is Pleistocene in age and in the Ic layer of the Haizume Formation in the Uonuma Group. In addition to the preinjection formation mineralogy and water chemistry investigation at Zone 2 and adjacent zones, an intermittent geophysical logging and time lapsed cross well seismic survey are being carried out to monitor CO₂ movement after injection. Laboratory experiments are also conducted. However, prediction of the long-term reaction between formation water dissolving CO₂ and formation rock is difficult by short term experiments and geochemical monitoring after CO₂ injection, such as water sampling is not planned. Therefore, a computer geochemical modelling was performed to evaluate the reactions after CO₂ injection and estimate the capacity of mineral storage of CO₂. Within this study the pre-injection mineral water equilibrium state was investigated using formation mineralogy and water chemistry, and subsequently possible CO₂-water-minerals reactions in the Haizume Formation were quantitatively simulated using the geochemical modelling code EQ3/6 version 7.2 (Wolery, 1992; Wolery and Daveler, 1992).

1 GEOGRAPHICAL AND GEOLOGICAL SETTING

The CO₂ injection site is located in Minami-Nagaoka gas field, south-west of Nagaoka City, Niigata Prefecture, at Honshu Island, Japan (Fig. 1). Since 2001, three observation wells (CO2-1 to CO2-3) were drilled in the Iwanoihara base by Teikoku Oil CO., Ltd. Core samples were used for different laboratory experiments and analysis. The CO₂ injection target Zone 2 comprises the Pleistocene Ic layer in the Haizume Formation (0–2540 m) of the marine Uonuma Group, which is covered with mudstone of 20 m thick (Sekiyu Gijutsu Kyokai, 1993). The stratigraphy of this area varies significantly by location. Further geological and stratigraphic details of the study area can be found in recent references of Kobayashi (1996) and Kobayashi and Takano (2002). Two potential injection reservoirs are situated between 1139 and 1156.5 m depth (NEDO, 2002).

2 PREINJECTION FORMATION CHARACTERISATION

2.1 Preinjection Formation Minerals in the Zone 2

The reservoir rock from the Zone 2 is a greenish-grey-colour sandstone and the grain size is in the range of medium to coarse sand with some rounded gravel. The sandstone shows minor consolidation.

The porosity and permeability of the rock samples are approximately 20% and 200mD respectively. Figure 2 shows a thin section image from a reservoir rock sample collected from Zone 2, at 1104 m depth from the well CO2-1. It is mainly composed of quartz, feldspars, volcanic rock fragments, and matrix as well as minor minerals and glasses fragments. The feldspar fraction is dominated by plagioclase. The composition of plagioclase varies and the average plagioclase composition is Ab₅₀/An₎₀. Accessory minerals include pyroxene, amphibole, epidote and chlorite. Table 1 summarises the composition of major minerals.
Figure 1
Location of the injection site.

Figure 2
Thin section image of the formation rock sample from 1104 m depth from the well CO2-1.
TABLE 1
Mineral composition of the reservoir rock collected from the well CO2-1 at the Zone 2 in the Ic layer in the Haizume Formation analysed by an observation of a thin section by optical microscope

<table>
<thead>
<tr>
<th>Minerals</th>
<th>vol (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Quartz</td>
<td>21.1</td>
</tr>
<tr>
<td>K-feldspar</td>
<td>4.3</td>
</tr>
<tr>
<td>Plagioclase</td>
<td>15.7</td>
</tr>
<tr>
<td>Mica</td>
<td>0.3</td>
</tr>
<tr>
<td>Chlorite</td>
<td>1.1</td>
</tr>
<tr>
<td>Pyroxene</td>
<td>4.1</td>
</tr>
<tr>
<td>Amphibole</td>
<td>0.2</td>
</tr>
<tr>
<td>Epidote</td>
<td>0.1</td>
</tr>
<tr>
<td>Rock fragment</td>
<td>25.9</td>
</tr>
<tr>
<td>Matrix and cement</td>
<td>17.8</td>
</tr>
<tr>
<td>Glasses</td>
<td>8.5</td>
</tr>
<tr>
<td>Other minerals</td>
<td>0.8</td>
</tr>
<tr>
<td>Total minerals</td>
<td>100.0</td>
</tr>
</tbody>
</table>

2.2 Preinjection Water Chemistry and Saturation State of Minerals in the Zone 2

Two types of water sample were collected in the Haizume Formation from the well CO2-1 using an air pump. The water composition was monitored during the sampling to control potential contamination. One type of water sample was collected from the adjacent Zones between 2 and 5 and the other from Zones between 3 to 5. Considering that these waters were the mixture of formation waters from different zones, the water composition of the Zone 2 was calculated from the difference between the two waters and flow rate. Total dissolved solid (TDS) is approximately 6100 mg/l.

Due to possible oxidation and degassing of CO2 during the sampling process, the water chemistry was adjusted to the formation conditions using EQ3/6. The formation temperature is 50°C, the oxidation level (Eh) was set to reduced condition using barite equilibrium. The partial pressure of CO2 ($p_{CO2}$) was adjusted to calcite saturation (Bazin et al., 1997). The pH of the water at in situ condition was calculated from the $p_{CO2}$ and total aqueous carbonate.

![Figure 3](image-url)

Major ion composition of the adjusted formation water from Zone 2, Haizume Formation.

TABLE 2
Formation water composition of samples from Zones 2-5, 3-5, calculated composition of Zone 2 (Zone 2*), and the adjusted chemistry to the formation condition

<table>
<thead>
<tr>
<th></th>
<th>Zones 3-5 (mg/l)</th>
<th>Zones 2-5 (mg/l)</th>
<th>Zone 2* (mg/l)</th>
<th>Adjusted to the formation condition (mg/l)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Na</td>
<td>2694.9</td>
<td>1936.3</td>
<td>1699.0</td>
<td>1699.7</td>
</tr>
<tr>
<td>K</td>
<td>815.5</td>
<td>385.4</td>
<td>251.0</td>
<td>250.4</td>
</tr>
<tr>
<td>Ca</td>
<td>487.1</td>
<td>420.5</td>
<td>400.0</td>
<td>399.2</td>
</tr>
<tr>
<td>Mg</td>
<td>10.2</td>
<td>16.0</td>
<td>18.0</td>
<td>17.8</td>
</tr>
<tr>
<td>Fe</td>
<td>4.2</td>
<td>0.1</td>
<td>0.0</td>
<td>0.0</td>
</tr>
<tr>
<td>Al</td>
<td>0.2</td>
<td>0.1</td>
<td>0.1</td>
<td>0.1</td>
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<tr>
<td>Ti</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
</tr>
<tr>
<td>Mn</td>
<td>0.3</td>
<td>0.5</td>
<td>0.6</td>
<td>0.6</td>
</tr>
<tr>
<td>Sr</td>
<td>1.8</td>
<td>1.8</td>
<td>1.8</td>
<td>1.2</td>
</tr>
<tr>
<td>Ba</td>
<td>1.2</td>
<td>1.4</td>
<td>1.4</td>
<td>0.6</td>
</tr>
<tr>
<td>Cl</td>
<td>5258.6</td>
<td>3852.1</td>
<td>3413.0</td>
<td>3418.7</td>
</tr>
<tr>
<td>SO4$^{2-}$</td>
<td>84.9</td>
<td>76.6</td>
<td>74.0</td>
<td>74.0</td>
</tr>
<tr>
<td>HCO3$^-$</td>
<td>801.0</td>
<td>372.0</td>
<td>241.0</td>
<td>494.0</td>
</tr>
<tr>
<td>SiO2</td>
<td>91.5</td>
<td>104.5</td>
<td>109.0</td>
<td>109.0</td>
</tr>
<tr>
<td>pH</td>
<td>8.01</td>
<td>7.90</td>
<td>7.8</td>
<td>6.3</td>
</tr>
<tr>
<td>Eh (V)</td>
<td>2.5</td>
<td>2.5</td>
<td>2.5</td>
<td>-0.2</td>
</tr>
<tr>
<td>Flow rate (kl/day)</td>
<td>5</td>
<td>21</td>
<td>16</td>
<td>-</td>
</tr>
</tbody>
</table>
3 GEOCHEMICAL MODELLING OF REACTIONS BETWEEN CO₂-FORMATION WATER-MINERALS IN THE ZONE 2, HAIZUME FORMATION

3.1 Modelling Conditions

The geochemical reactions between CO₂, mineral and formation water after the CO₂ injection in the Zone 2 were simulated using the EQ3/6 code. Since CO₂ dissolution was much faster compared mineral reactions, dissolution kinetics of CO₂ was not considered in this simulation. First, the total aqueous carbonate equilibrium with pH₂ = 3(g) was calculated. Then maintaining the pH₂, the interaction between formation minerals and the CO₂ saturated formation water was simulated kinetically until all the minerals reached equilibrium in the closed system at constant formation temperature 50°C and formation pressure 11 MPa, assuming to be the hydrostatic pressure at depth of 1100 m. The porosity was assumed 20%.

The dissolution kinetics were modelled using the rate law (Lasaga, 1984). In general, the dissolution rate (rₘ) of mineral m can be expressed:

\[ r_m = A_m k_m \left( 1 - \frac{Q_m}{K_m} \right)^n \]  \hspace{1cm} (1)

where \(A_m\) is the specific surface area, \(k_m\) is the kinetic rate constant, \(Q_m\) is ion activity product and \(K_m\) is the equilibrium constant for the mineral-water reaction. The parameters \(n\) and \(n\) are two positive numbers, but they are taken to unity when experimental data are not available.

The kinetic rate constant \(k\) at 50°C were calculated using the Arrhenius equation (Lasaga, 1984):

\[ k = k_{25} \exp \left( \frac{-E_a}{R} \left( \frac{1}{T} - \frac{1}{298.15} \right) \right) \]  \hspace{1cm} (2)

where \(E_a\) is the activation energy (J/mol), \(k_{25}\) is the rate constant at 25°C, \(R\) is the gas constant (8.31 J/molK) and \(T\) is absolute temperature (50°C = 323.5K).

The precipitation of minerals was set to be instantaneous when a mineral reached to the saturation.

3.2 Input Data

3.2.1 Formation Mineral Composition

Based on the formation petrography and mineralogy study, fourteen minerals including quartz, K-feldspar, plagioclase, mica, pyroxene, amphibole, clay and some accessory minerals were introduced as reactants into the modelling. Other minerals potentially involved in the reactions as products include kaolinite, rutile, dolomite, whiterite, dawsonite, and solution solid calcite. The solution solid calcite (hereafter ss-calcite) is a carbonate with six poles (calcite, magnesite, siderite, stronitiate, rhodochrosite, and smithsonite) and the chemical formula is (Ca, Mg, Fe, Zn, Sr, Mn)CO₃.

3.2.2 The Reactive Surface Area

The reactive surface area for each mineral is generally calculated from the specific surface area and the quantity of each mineral assuming that the reactive surface is proportional to specific surface. However, in practical sense, it is impossible to measure the surface area of each mineral in the whole rock. The conventional method is to allocate the surface area to each mineral depend on the volumic composition and to bring the contribution of surface area from each mineral to the total surface area, for example measured by the BET method (Brunauer et al., 1938). However, this method tends to overestimate the surface of concentrated minerals such as quartz and underestimate the small-size minerals. Assuming that the total surface area is mainly contributed by small-size grains such as clay minerals, a different approach of estimation of surface area was applied (Kozaka, 1995). For this rock sample, the BET surface area measured was only available for a ground sample with a value of 6.7 m²/g. The rock sample was ground below 0.25 mm, and this is larger than most of the observed grain size. The total surface area was adjusted to the sum of a calculated surface for the individual minerals to a constant specific area of 60 m²/g for chlorite and smectite. A constant grain size (0.04 mm) was used to calculate geometric surface area for the remaining minerals. The reactive surface area was assumed the same as the surface area of each mineral. The input parameters for modelling including initial moles, specific surface area of the reactant minerals, dissolution rate constants at 25°C, activation energy, calculated dissolution rate at 50°C, and references are summarised in Table 3.

4 RESULTS

4.1 Initial Dissolution of CO₂ into the Formation Water

As a starting point for the simulation, the total aqueous carbonate equilibrium with pH₂ = 3(g) was calculated. As
shown in Figure 4 (log year = –6), the contact with high pressure CO₂ decreased the formation water pH and increased the redox potential (Eh). The pH value decreased from 6.3 to 3.6 and Eh increased from –0.20 V to 0.02 V. The amount of total aqueous carbonate is 2.04 mol/kg H₂O (89.8 kg CO₂/m³ H₂O) without mineral reactions.

4.2 Mineral Reactions with CO₂ Saturated Formation Water

4.2.1 pH and Eh

Figure 4 shows the evolution of pH and Eh of the formation water up to 10⁶ years. The pH and Eh evolutions were controlled by mineral dissolution and precipitation in the whole system. Dissolution of most of the minerals in this formation consumes H⁺ and increases the pH of the solution and precipitation of these minerals releases H⁺. The evolution of Eh is almost symmetric to that of pH, except an increase in Eh after 8 days (= 10⁻¹.7 years) corresponding to pyrite saturation.
As soon as the minerals were in contact with CO₂ saturated water, the pH started to increase. The rate of change of pH of the system slowed after 8 days (≈ 10⁻¹.⁷ years). The rate of change of pH again increased after 20 (≈ 10⁻¹.³ years) and accelerated after 117 (≈ 10².⁰) years. The pH decreased again after approximately 3160 (≈ 10³.⁵) years, and the decrease slowed after 15 300 (≈ 10⁴.²) years. These changes corresponded to different events in the systems such as a start of the dissolution of unstabilised minerals (just after the contact with CO₂ saturated water) and solid solution calcite (ss-calcite), dawsonite and pyrite saturation (8 days = 10⁻¹.⁷ years), and enhanced dissolution of fast reactive minerals such as smectite, chlorite, K-feldspar, muscovite, olivine, amphibole after 20 (≈ 10⁻¹.³ years). However, muscovite reached equilibrium after 117 (≈ 10².⁰) years and started to consume H⁺ therefore stabilising pH. The pH increased again after 238 (≈ 10².⁴) years when plagioclase dissolution became more dominant. When the fine-gained clay minerals such as chlorite and smectite are exhausted, and both ss-calcite and muscovite precipitated after 3160 (≈ 10³.⁵) years and the pH will decrease until kaolinite starts to precipitate at 15 300 (≈ 10⁴.²) years.

4.2.2 Aqueous Species

The evolutions of total concentration of major elements in the Zone 2 water up to 10⁶ years are presented in Figure 5. There were some different Ca source minerals, however the main source of the Ca was plagioclase and Ca was precipitated as ss-calcite and dolomite. The concentration of Mg was mainly controlled by dissolution of orthopyroxene and precipitation of dolomite and ss-calcite. The sources of K were K-feldspar, biotite and muscovite. Muscovite dissolved at the beginning of the simulation, but reached equilibrium at 117 (≈ 10².⁰) years. Subsequent released K was incorporated into muscovite. Al was released by all the alumino-silicates, in particular plagioclase and K-feldspar. Al concentration is sensitive to pH of the solution and the solubility of Al is low between pH 4.5 to 9 (Mason, 1966). During K-feldspar and plagioclase dissolution, Al was incorporated into muscovite and dawsonite. When the K-feldspar was depleted, Al was incorporated into kaolinite. Si concentration was controlled by quartz, which was in equilibrium with the solution throughout the simulation. Na was almost constant: released Na from plagioclase was incorporated into dawsonite. Total C (carbonates) was constant, corresponding to a constant pCO₂. Some of Fe was incorporated into ss-calcite. Ti released from titanite was included into rutile. Ba concentration in the water is controlled by witherite and barite; Ba was first precipitated as witherite then as barite as the S concentration in the water increased due to pyrite dissolution.

4.2.3 Mineral Phase

Figure 6 presents the evolution of the mineral assemblage up to 200 000 (10⁵.³) years, when plagioclase, which is the second most abundant mineral, was exhausted. The time axis of this evolution graph is linear. Initial major components were quartz, K-feldspar plagioclase, biotite, orthopyroxene, pyrite, and titanite. However, K-feldspar and plagioclase were completely dissolved after 10⁴.⁶, 10⁵.³ years and biotite after 10⁵.⁰ years, respectively. Orthopyroxene and titanite persisted until 10⁷.³ years. Instead, muscovite, kaolinite, dolomite, dawsonite, ss-calcite, and rutile appeared. The final assemblage mainly comprised quartz, ss-calcite, dawsonite, dolomite, muscovite, kaolinite, pyrite and rutile. Major reactions in this system were dissolution of plagioclase,
K-feldspar, orthopyroxene and precipitation of ss-calcite, dawsonite, muscovite, kaolinite and quartz. Initially Al released was incorporated into muscovite and when K-feldspar depleted, excess Al from plagioclase precipitated as kaolinite.

### 4.2.4 Stored CO₂

Injected CO₂ was stored in the aqueous and mineral phases. Aqueous phase storage was mainly in form of CO₂(aq) and minor HCO₃⁻. The total CO₂ stored in both aqueous and mineral phases and each mineral evolution are presented in Figure 7.

Approximately 2.0 mol/kgH₂O (90 kgCO₂/m³H₂O) of CO₂ was stored in the aqueous phase. This amount was constant during the simulation. Mineral storage (carbonate precipitation) started after 8 days (≈10⁻¹.⁷ years) and increased up to 21.3 mol/kgH₂O (940 kgCO₂/m³H₂O). Main mineral phases were ss-calcite, dawsonite and dolomite. The Zone 2 could store CO₂ in both aqueous and mineral phase 18.8 mol/kgH₂O (827 kgCO₂/m³H₂O) first 200 000 years and 23.3 mol/kgH₂O (1025 kgCO₂/m³H₂O) at the final stage.

### DISCUSSION

Dissolved CO₂ was stored in both aqueous and mineral phases in the formation. Mineral storage of CO₂ was by precipitation of dolomite, solid solution calcite, and dawsonite. The maximum stored CO₂ was 23.3 mol/kgH₂O, where mineral storage 21.3 mol/kgH₂O, which corresponded to 90% of total storage. The mineral reaction enhanced the uptake of CO₂ by a factor of 10. The mineral storage capacity of the formation was high, which reflected the high concentration of divalent-ion-source minerals of the formation such as plagioclase and orthopyroxene. The Haizume Formation is therefore an ideal target for mineral storage of CO₂.

Na released from plagioclase was assumed to be precipitated as dawsonite. Although this mineral has been observed at limited area (Smith and Milton, 1966; Goldbery and Loughman, 1970; Loughman and Goldbery, 1972), its dissolution/precipitation kinetics are not well investigated. Until the complete dissolution of plagioclase by 200 000 years, the major reactions of the system were dissolution of plagioclase, K-feldspar and orthopyroxene and the formation of ss-calcite, dawsonite, muscovite, kaolinite and quartz. K-feldspar dissolution and muscovite precipitation was supported by Al supply from plagioclase dissolution. Precipitation of muscovite contributed to lower the pH of the system. Plagioclase dissolution released Ca and Na and these elements precipitated as ss-calcite or dawsonite also removing CO₂ from the aqueous phase. As K-feldspar concentration decreased, the dissolution of K-feldspar decreased accordingly and excess Al released from plagioclase started to precipitate as kaolinite. Muscovite precipitation slowed and continued until biotite was depleted. In this time scale, the mineral storage capacity was 16.8 mol/kg H₂O.

In this simulation, as the dissolution kinetics of CO₂ was considered faster than mineral reactions, the reactions were modelled between minerals and CO₂ saturated formation water. In the real situation, the dissolution rate of the CO₂ may be of high importance for a short-time scale.

The surface area was assumed the same as the surface area 6.7 m²/g of the ground sample and a constant surface area of 60 m²/g for clays (chlorite and smectite) and the constant grain size of 0.04 mm was used to calculate geometric surface area for larger grains. However, this estimation is probably too high and more realistic estimation of the surface area of each mineral is therefore necessary. Since an input surface area influences the time scale of the simulation (Xu et al., 2001; Zwingmann, unpublished data) the time scale of the simulation also contains some uncertainty. A detailed sensitivity analysis will aid to evaluate the influence of surface area and other geochemical parameters. In addition, the flow rate of the formation water in the Zone 2 is unknown. Fluid flow and transport could be an important factor if present an application of reaction-transport coupled model would be required under this circumstance. Due to the estimated high amount of mineral precipitation, it would be important to predict where and when these authigenic minerals precipitate in the pore structure.

### CONCLUSION

From the preinjection analyses of the Haizume Formation, Zone 2 in the Uonuma Group was identified as a potential CO₂ injection formation target. The sandstone from the Zone 2
contains significant amount of reactive and divalent-ion-source minerals such as plagioclase, orthopyroxene, and biotite. There is also K-feldspar, which is not divalent-ion-source, but also reactive in the acid environment. The formation water from the Zone 2 is low TDS (6100 mg/l) Na+-Ca²⁺-Cl⁻ water. Ca²⁺ concentration is more than 20% of Na⁺ concentration.

The potential major mineral reactions simulated between formation minerals and CO₂ saturated water were dissolution of plagioclase, orthopyroxene and K-feldspar and precipitation of ss-calcite, dawsonite, dolomite, muscovite, kaolinite and quartz.

Storage of CO₂ in minerals started after few days and phases appeared as ss-calcite, dolomite and dawsonite. Reflecting these highly reactive minerals, the Zone 2 has a potential of CO₂ storage 18.8 mol/kgH₂O by 200 000 years and maximum 23.3 mol/kgH₂O at the final stage (10⁷.3 years). The CO₂ storage was greatly enhanced by the formation mineral despite the fact that the time scale of the simulation contains some uncertainty. Introduction of the dissolution kinetic of CO₂ and more accurate estimation of specific surface will aid to increase the accuracy of the model. If notable fluid flow is present in the injection target zone the modelling should be extended to a coupled flow model.

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