A modelling study to evaluate the effect of impure CO₂ on reservoir performance in a sandstone saline aquifer

Mohammed Dahiru Aminu a,*, Vasilije Manovic b

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ABSTRACT
Carbon capture and storage (CCS) is expected to play a key role in meeting greenhouse gas emissions reduction targets. In the UK Southern North Sea, the Bunter Sandstone formation (BSF) has been identified as a potential reservoir which can store very large amounts of CO₂. The formation has fairly good porosity and permeability and is sealed with both effective caprock and base rock, making CO₂ storage feasible at industrial scale. However, when CO₂ is captured, it typically contains impurities, which may shift the boundaries of the CO₂ phase diagram, implying that higher costs will be needed for storage operations. In this study, we modelled the effect of CO₂ and impurities (NO₂, SO₂, H₂S) on the reservoir performance of the BSF. The injection of CO₂ at constant rate and pressure using a single horizontal well injection strategy was simulated for up to 30 years, as well as an additional 30 years of monitoring. The results suggest that impurities in the CO₂ stream affect injectivity differently, but the effects are usually encountered during early stages of injection into the BSF and may not necessarily affect cumulative injection over an extended period. It was also found that porosity of the storage site is the most important factor controlling the limits on injection. The simulations also suggest that CO₂ remains secured within the reservoir for 30 years after injection is completed, indicating that no post-injection leakage is anticipated.

1. Introduction

The global emissions of carbon dioxide into the atmosphere, caused mainly by the burning of fossil fuels such as those from industrial processes (e.g., cement, steel and lime production), power and transportation sectors, have caused global warming and climate change [1]. Carbon capture and storage (CCS) has been considered as one of the viable climate change mitigation technologies, and it is expected to help in reducing over 20% of the global greenhouse gas (GHG) emissions by 2050, while its exclusion can cause over 70% increase in the global cost of meeting emission reduction targets [2].

The main options for CO₂ storage in underground geological formations are saline aquifers, depleted oil and gas reservoirs, unmineable coal seams, basalt formations, hydrate storage of CO₂ within the subsurface environment, and CO₂-based enhanced geothermal systems [3]. Among these options, storage in saline aquifers is considered as one of the most feasible choices for technology deployment, since they provide the largest potential storage volume [4] and are unsuitable for other uses.

Depending on the source of the captured CO₂, some impurities (usually <5%vol. [5, 6]) are contained in the CO₂ stream. The type and amount of impurities in the CO₂ stream may have a significant effect on the physical qualities of the storage reservoir. The presence of impurities can alter the molar volumes and shift the boundaries in the CO₂ phase diagram to higher pressures, which implies higher operating pressures and cost may be required for CO₂ storage [3]. Therefore, the overall storage capacity can be adversely affected [7]. On the other hand, the possible short term effects of impurities on the reservoir performance may indicate the necessity of a change in injection strategy [7]. While it is crucial to quantify the long-term effects of impurities on storage sites in real life applications, it is equally important to note that laboratory experimental approaches are complex, costly and inadequately slow. Instead, numerical simulations can be used to accelerate the evaluation of impurity effects. Several numerical studies have been conducted to evaluate the long-term reservoir performance when impure CO₂ is injected into geological formations [7, 8, 9, 10], and such studies suggest that the presence of impurities may cause a considerable influence on injection and storage mainly due to chemical reactivity of the host rock with formation water resulting in changes in permeability and porosity. For example, the presence of SO₂ could lead to the formation of strong acids, which could then cause further dissolution and precipitation of

* Corresponding author.
E-mail address: mdaminu@mautech.edu.ng (M.D. Aminu).

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rock minerals. Additionally, the presence of impurities in the injected CO₂ and the subsequent geochemical reactions with saline water and minerals in the host rock can lead to the precipitation of authigenic or secondary mineral phases which could effectively lock up the stored CO₂ in immobile secondary phases for geologic timescales [11].

Waldmann et al. [7] studied the physicochemical effects of discrete CO₂–SO₂ mixtures on injection and storage in order to assess the impact on overall storage capacity in the Ketzin injection site, Germany. In their study, they considered geological conditions in the Ketzin site, and they assessed the impact of SO₂ on the physicochemical behaviour of the CO₂ phase by geochemical modelling of fluid-rock interactions. Since variations in porosity and permeability of the host rock can negatively influence CO₂ storage capacity, the results obtained from chemical models on porosity and permeability were integrated into the reservoir simulations to assess likely impacts. Results from their study revealed that the presence of SO₂ caused a significant reduction in porosity compared to pure CO₂ during the period of injection. Their results also suggested that the overall impact of SO₂ (<1%vol.) was low, although the morphology changes due to chemical reactions reduced the pore spaces available for storage. Similarly, Wang et al. [9] studied the impact of SO₂ on CO₂ injectivity in the Basal Cambrian sandstone saline aquifer in Western Canada. In their study, 2.5%vol. SO₂ was used, and they reported, for the first time, that enhanced quartz dissolution due to the effect of SO₂ apparently caused significant precipitation of NaCl, which could reduce the permeability of reservoir rocks.

To the best of our knowledge, numerical models capable of evaluating long-term reservoir conditions during injection and storage of CO₂ and impurities have not been previously utilised [8], given that the current research is mainly focused on the thermophysical properties of impurities that are inevitably present in the CO₂ stream. However, it is worth noting that these properties can be confidently predicted from knowledge of the temperature, pressure and brine chemistry [12]. To further de-risk CO₂ storage, it is important that research is conducted in understanding the effect of impurities introduced through the CO₂ stream on the thermophysical properties of the CO₂-brine system of the reservoir [12]. Therefore, in this study, we evaluate the effect of CO₂ and impurities (NO₂, SO₂ and H₂S) during injection and storage in a sandstone saline aquifer. We evaluate their effects on key reservoir performance indicators, i.e., well bottom-hole pressure, CO₂ injection rate, field pressure and cumulative gas injection using a single horizontal well injection strategy for a 30-year period, and an additional 30-year period of monitoring. The Bunter Sandstone formation (BSF) in the southern sector of the UK North Sea was selected as a case study. A simplified model for reservoir simulation to evaluate the reservoir performance during injection and storage using time and specific observation point indices was developed.

2. Numerical model

2.1. Reservoir model and boundary conditions

To develop the model of the BSF for flow simulations, we used simplified box models where the geology and domain of research were characterised as horizontal layers and representative bulk values were assigned for the various reservoir properties using data from laboratory experimental analysis [13] and the open literature. The injection and storage of CO₂ into the BSF saline aquifer was simulated using the ECLIPSE E300 compositional simulator which can model fluid flow as a multiphase and multicomponent system. A similar reservoir simulator (ECLIPSE E100 black oil) was used in previous studies [14, 15, 16, 17, 18] on CO₂ storage, without considering the composition of injected fluids and the effects of impurities on reservoir performance. For a compositional simulation involving a multicomponent run, the key parameters that account for each impurity case are the experimentally determined permeability values, volume of each component in the injection stream and diffusion coefficients of components in water.

The simulation process involves several steps (Figure 1) which include inputting data to the CO2STORE module of the simulator that conform with reservoir properties of the targeted storage formation; the use of lithostatic pressure gradient for model validation with respect to the injection depth; and conducting sensitivity tests with respect to grid dimensions in order to select the grid that provides optimum injection within the targeted reservoir. The rock and fluid properties used for multiphase flow simulations are presented in Table 1. A grid of 10 × 10 × 3 cells (300 grid cells in total) was used, and the model dimension was discretised into 90 km × 90 km × 3 km. Notwithstanding that the grid is coarse, an adaptive mesh refinement for optimising the reservoir performance was not conducted in the simulation as it is known that for a single injector that seeks to store CO₂ over a storage region of large lateral extent such as the BSF, even a less refined mesh provides an accurate representation of the reservoir performance [19, 20, 21].

The depth to the crest of the Bunter storage structure was 1171 m [22], and the thickness of the BSF at the storage point was 215 m [22]. Therefore, we used 1171 m as depth of the top face of the grid block, and we adopted 1181 m as injection depth. To validate the model against lithostatic pressure gradient, 0.165 bar/m was used. This was chosen as a typical value for lithostatic pressure gradient reported by Energy Technologies Institute (ETI) [22] for the Bunter storage development plan and is close to values used by Noy et al. [23] and Williams et al. [17]. Additionally, in order not to suffer mechanical failures at any point during injection operations, the simulation was initially run (with the average porosity and average permeability values of the BSF derived by ETI [22]) and injection pressure was limited to 90% of the lithostatic pressure (175 bar) at injection depth (1181 m). The pressure constraints from our simulations were found to be in good agreement with anticipated pressure, and we established that our model runs within safe injection pressure. Moreover, the injection operations are expected to be free from any sanding risks due to geomechanical issues. The analysis of existing wells within the BSF suggest that the reservoir is relatively strong and sand failure events around the near wellbore area [22], which could cause subsequent operational problems, are unlikely.

Saline aquifers are usually very large as they cover hundreds of km² and the BSF is not an exception (Figure 2). Therefore, modelling for CO₂ storage in saline aquifers is usually focused on the area affected by the CO₂ plume distribution [17], which is usually around the injection point and surrounding regions. However, in selecting an appropriate reservoir size for modelling, it is important to note that a fairly considerable aquifer size should be adopted since pressure footprint due to injection can extend much further in the aquifer [17, 23, 24] to cause unrealistic pressure build-up in the model [17]. Therefore, in this study, the optimum reservoir size was obtained by conducting simulations with various grid sizes and correlating the results with available pressure data [22]. The grid dimension was kept constant while the grid block size was varied in the DX and DY directions by a deduction of 20 m from the previous grid block size in each subsequent simulation. The variation of these grid blocks was then limited to reservoir sizes that were capable of sustaining CO₂ injection for the specified 30-year period. In order to select the optimum reservoir size, we used a method proposed by ETI [22] for the Bunter storage development plan, which suggests that optimum reservoir size could be selected from a grid in which the injection rate can be maximally sustained.

It is generally agreed that the boundaries of a reservoir can influence the pressure build-up and storage capacity. The most conventional estimates of the boundaries are assumed to be horizontal, and vertical boundaries of the model are closed [19, 24]. However, for CO₂ storage applications over injection timescales and subsequent migrations of <100 years, the boundary conditions are found to have a negligible impact on reservoir performance [25]. This is due to the large areal extent of the Bunter region [25]. Therefore, we believe that our results are valid for both sets of open and closed boundary conditions since they are based on reservoir performance evaluation over this timescale [25].
Figure 1. The workflow for numerical simulation.

- Interpreted geology, geophysics and petrophysics
- Reservoir characterisation
- Run ECLIPSE Module
- Model production and pressure validation tests
- Sensitivity runs (Identify uncertain parameters)
- Tuning runs (Modify uncertain parameters)
- Actual CO₂ injection and storage
  - Existing wells continue to inject
  - New wells may be implemented
Table 1. Rock and fluid properties used for the simulations.

| Parameter                          | Value                  | Unit | Reference     |
|-----------------------------------|------------------------|------|---------------|
| Grid dimension                    | $10 \times 10 \times 3$ | -    | -             |
| Grid block size                   | $300 \times 300 \times 10$ | m    | -             |
| Porosity (lower and upper limits for all model runs) | 10, 22 | % | Vincent [26] |
| Permeability (variable depending on impurity present) | $254^a, 140^b, 225^c, 255^d$ | mD  | Aminu et al. [13] |
| Depth to top face of grid block   | 1171                   | m    | ETI [22]      |
| Reference depth                   | 1181                   | m    | -             |
| Model datum depth                 | 1181                   | m    | -             |
| Initial reservoir temperature     | 37                     | °C   | ETI [22]      |
| Average reservoir pressure        | 124                    | bar  | ETI [22]      |
| Bottom-hole pressure limit        | 175                    | bar  | -             |
| Content of impurity in the CO₂ stream | 5                     | %vol | [5, 6]        |
| Diffusion coefficients of components in water | $3.0 \times 10^{-4}$ (H₂O), $1.4 \times 10^{-4}$ (CO₂), $1.0 \times 10^{-4}$ (NaCl), $1.21$ (NO₂), $0.0001$ (SO₂), $0.0002$ (H₂S) | m²/day | Cussler [33] |
| Rock density                      | 2300                   | kg/m³ | ETI [22]     |
| Lithostatic pressure gradient     | 0.165                  | bar/m | ETI [22]     |
| Pore compressibility              | $4.5 \times 10^{-10}$  | Pa⁻¹  | Zhou et al. [34]; Noy et al. [23] |
| Fluid injection rate              | $6 \times 10^3$        | Sm³/CO₂/day | -          |
| Injection period                  | 30                     | Year  | -             |
| Monitoring period                 | 30                     | Year  | -             |

Note: a, b, c, and d, denote permeability values obtained after exposure to CO₂, CO₂–NO₂, CO₂–SO₂ and CO₂–H₂S, respectively.

2.2. Reservoir and fluid properties

The BSF was deposited during the late Permian and Triassic periods (Figure 3). It comprises pebbly sandstones and sandstones intercalated with small amounts of conglomerates, mudstones and siltstones. The reservoir formation is typically 200 m thick and has fair to good porosity and permeability, and occurs at depths of about 1000–3000 m [23]. The BSF is also overlain primarily by the Triassic mudstones of the Haisborough Group which act as caprock or seal and it is underlain by the air-derived non-condensable species, e.g. N₂, O₂, Ar, is recommended to benchmark for CO₂ capture, utilisation and storage (CCUS) systems [43]. In any transport and storage applications, the concentration of air-derived non-condensable species, e.g. N₂, O₂, Ar, is recommended to not exceed 4%vol [43] while other species may not exceed 5%vol [5, 6] due to effects on compression and storage systems [43]. Consequently, the impurity levels adopted for this study agree well with recommendations in the literature for guidelines on conceptual studies.

2.3. Governing equations

For the multiphase flow model used in this study, the governing equations for simulation of CO₂ injection for storage into a saline aquifer are like those used for oil, water, and CO₂ flows through porous reservoirs. Darcy’s law, which incorporates mass and energy conservation equations, as reviewed by Jiang [44], was used for the simulations.

\[ q = - \frac{k}{\mu} (\nabla p - \rho g) \]  

(1)

Using Eq. (1), it is possible to calculate velocity through the porous medium:

\[ v = \frac{q}{\rho} = - \frac{k}{\mu \rho} (\nabla p - \rho g) \]  

(2)

In terms of the positive z-direction as vertically up (opposite to gravity), multiphase extension of Darcy’s law can be used for an individual fluid phase α, thus:

\[ v_α = \frac{q_α}{\rho_α} = - \frac{kα}{\mu_α} (\nabla p_α - \rho_α g \nabla z) \]  

(3)
For CO₂ storage, the flow is modelled as a multiphase which requires considering CO₂, brine, rock, and multicomponents such as the CO₂ and water system. Thus, the number of components and phases can differ depending on application. In Eq. (4), the conservation of mass is expressed by the balance of four components that represent all possible mechanisms of mass transfer, which are: temporal rate of change of mass at fixed point, which can also be referred to as the local derivative or fixed term; convective mass transport; diffusive mass transport; and sink or source term for mass.

\[ \frac{\partial}{\partial t} \left( \sum_{\alpha} (\rho_{\alpha} X_{\alpha}^i) \right) + \sum_{\alpha} \nabla \left( \rho_{\alpha} X_{\alpha}^i q_\alpha \right) - \sum_{\alpha} \nabla \left( \varphi S_{\alpha} r_{\alpha} D_{\alpha} \nabla X_{\alpha}^i \right) = S_i \]  \quad (4)

The gas phase diffusion coefficient for each component is also accounted for in a compositional run. These are used to define diffusive flows in terms of vapour mole fractions. The normal diffusion coefficients are defined by the following condition (Reid et al. [45]):

\[ J_i = -cD_i \frac{\partial y_i}{\partial \bar{d}} \]  \quad (5)

These diffusion coefficients are used in the compositional model run to obtain gas inter-block diffusive flows, which take the form:

\[ F_{\text{diff}}^g = T_0 D_g \left( \rho b^2 - \rho \right) \Delta y_i \]  \quad (6)

For CO₂ storage reservoir simulations, the capillary forces are significant both in residual and structural/stratigraphic trapping. In the seal or caprock, the threshold of the capillary force can be high enough to keep the non-wetting phase, which can be CO₂ fluid or gas phase, from entering small pore throats in the seal. The capillary force also keeps bubbles of the CO₂ phase in an immobile state in small pore spaces of the reservoir during CO₂ migration. Thus, capillary pressure, which is the pressure difference between the non-wetting phase and wetting phase in a porous medium, is given as:

\[ P_c = P_w - P_n \]  \quad (7)

The compositional model run uses a modified Peng-Robinson equation of state [46] which is able to correctly compute the density, viscosity and compressibility of CO₂ as a function of temperature and pressure, as well as the mutual solubility of CO₂ and brine.

For the density of brine and CO₂, the determination of amount of CO₂ that will be dissolved in brine is important for the estimation of storage capacity and in understanding the interactions between CO₂-brine-rock because of low pH or acidity of CO₂-saturated brine [47]. Since the dissolution of CO₂ in brine influences the solution density, the influence of dissolved CO₂ on brine is expressed thus:

\[ \rho = \rho_b + M_{CO_2} C - C \rho_b V \]  \quad (8)

2.4. Model study design

In this study, we designed a reservoir simulation case to evaluate the limits of stability of various chemical phases in the CO₂ stream with respect to operational uncertainties. This allowed us to account for the likely uncertainties that could arise in the deployment of CO₂ storage in the BSF with different impurities in the injection stream using surface flow rate as the well control mode. In practice, a CO₂ injection project could operate with multiple injection wells, e.g., the In Salah project [48, 49, 50], or single injection well, e.g., the Sleipner project [1]. However, the number of injection wells that can be deployed for any storage project may vary depending on several factors such as the reservoir heterogeneity [19], the amount of CO₂ which can be injected into a single well without causing adverse overpressure [51], the trade-off between cost of

Figure 2. Distribution of the Bunter Sandstone formation in the UK sector of the southern North Sea [23].
adding a well [19] and/or the amount of CO₂ that could be taken by any single well.

Previous studies on CO₂ storage in the BSF [17, 19, 23, 39] suggest that the motivation for using multiple injection wells for storage is largely driven by the amount of CO₂ that could be taken by each well per year. Typically, industry practice has suggested that volumes ranging from 1-2 MtCO₂/year [19, 52] could be stored in geologic formations beneath the North Sea, and this can be achieved with 20 injection wells [19], or even 12 injection wells [19, 23]. Moreover, ETI [22] deployed 4 injection wells to store approximately 7 MtCO₂/year over a period of 40 years. A constant injection rate as well as a constant injection pressure for the entire period of injection were assumed. In practice, the CO₂ injection rate could vary to maximise injectivity. However, it must be noted that the limits of stability in the CO₂ phase diagram regarding the operational uncertainties occur during the well pressure build-up prior to stability. Thus, the variation of injection rate after peak pressure is attained could only lead to a reoccurrence of the shifts in the boundaries of the CO₂ phase diagram as it was in the case of the preceding injection rate. We also assumed that 6 × 10⁶ Sm³CO₂/day was made available from an industrial site and transported for injection at the storage site. Thus, 6 × 10⁶ Sm³/CO₂/day was set as injection rate, but the volume of CO₂ injected was initially dependent on the well control modes, although the injection will normally be affected by the pressure build-up in the reservoir. Since the allowable bottom-hole pressure for injection was set to 90% lithostatic pressure at depth, the constant injection pressure used during the storage period ensured that the limit was not exceeded.

While our modelling did not consider the likely effect of impurity on any change in reservoir porosity, so that initial porosity values from experimental data can be assigned in each impurity case as we did for changes in reservoir permeability obtained from laboratory experimental analysis [13], it does not imply that impurities could not have affected the porosity of the BSF, and hence its reservoir performance. However, our assumption for adopting the upper and lower limits of the BSF porosity as reported by Vincent [26] in these simulations was informed by the analysis of mineral phases of rock samples after exposure to CO₂ and impurities, as reported in our previous study [13]. From the results, we can infer that there could be a significant reduction in porosity due to dissolution of clay minerals such as mica [53, 54] and an increase in porosity due to the dissolution of quartz [55], feldspar and albite [56].

3. Results and discussion

After injection, CO₂ plume is expected to migrate to the top of a reservoir due to buoyancy and it gradually spreads out laterally. The force of buoyancy causes the less dense gas to rise over denser saline water. The pattern of plume migration occurs both laterally and vertically and is expected to behave in this way due to the absence of any geological barriers to flow. The presence of impermeable seals atop and beneath the storage formation is expected to keep the plume spreading across the radial distance of the reservoir. Figures 4 and 5 are sketches of flow processes during CO₂ injection and storage in the reservoir.

From a geochemical perspective, CO₂ and impurities can influence the storage system in basically three ways: formation of carbonic acid or bicarbonates by dissolution of CO₂ in formation water (Equations (9) and (10)); production of weak or strong acids due to the effects of impurities; and dissolution and/or precipitation of calcite as cementing material [13, 58, 59].

\[
\text{CO}_2 + \text{H}_2\text{O} \rightarrow \text{H}_2\text{CO}_3 \tag{9}
\]

\[
\text{CO}_2 + \text{H}_2\text{O} \rightarrow \text{H}^+ + \text{HCO}_3^- \tag{10}
\]

Unlike the dissolution of CO₂ in formation water, the introduction of NO₂ in the injection stream can produce a weak acid HNO₂ or a strong acid HNO₃, as given below (Equations (11) and (12)):

\[
2\text{NO}_2 + \text{H}_2\text{O} \rightarrow \text{HNO}_2 + \text{HNO}_3 \tag{11}
\]

\[
3\text{NO}_2 + \text{H}_2\text{O} \rightarrow 2\text{HNO}_3 + \text{NO} \tag{12}
\]

Similarly, the introduction of SO₂ in the injection stream can produce weak acids H₂SO₃ and H₂S and/or a strong acid H₂SO₄ in the formation water, as given thus (Equations (13), (14) and (15)):

\[
\text{SO}_2 + \text{H}_2\text{O} \rightarrow \text{H}_2\text{SO}_3 \tag{13}
\]
SO₂ + H₂O ⇌ \( \frac{3}{4} \) H₂SO₄ + \( \frac{1}{4} \) H₂S \hspace{1cm} (14)

SO₂ + H₂O + \( \frac{1}{2} \) O₂ ⇌ H₂SO₄ \hspace{1cm} (15)

The introduction of H₂S in the injection stream can be explained thus (Equations (16) and (17)):

H₂S + H₂O → HS⁻ + H₂O⁺ \hspace{1cm} (16)

HS⁻ + H₂O → S⁻ + H₂O⁺ \hspace{1cm} (17)

In addition, the dissolution and/or precipitation of calcite involves a cascade of simultaneous reactions [60, 61, 62, 63, 64], as given thus (Equations (18), (19) and (20)):

CaCO₃ + H⁺ ⇌ Ca²⁺ + HCO₃⁻ \hspace{1cm} (18)

CaCO₃ + H₂CO₃ ⇌ Ca²⁺ + 2HCO₃⁻ \hspace{1cm} (19)

CaCO₃ + H₂O ⇌ Ca²⁺ + HCO₃⁻ + OH⁻ \hspace{1cm} (20)

In this study, the well bottom-hole pressure (WBHP) build-up in the reservoir was monitored at two periods which correspond to the first day of injection and the time it took to reach the peak pressure, for each simulation case, respectively. These observations enabled the determination of any potential effect of impurities on pressure perturbations and stability during injection. The WBHP build-up for storage sites with 10% and 22% initial porosity for pure and impure CO₂ storage is shown in Figure 6. For all storage sites, we observed a continuous increase in pressure until a maximum of 175 bar was reached, which then remained constant throughout the period of injection. However, for different simulation cases, the time to reach the maximum WBHP varied.

The WBHP of pure CO₂ injection into the storage site with 10% initial porosity reached its maximum pressure after 45 days of injection. In comparison to injection with impurities, the injection of CO₂–NO₂ resulted in the maximum pressure being attained within a relatively shorter period. Likewise, after the injection of CO₂–SO₂ and CO₂–H₂S, the time it took to attain maximum pressure in each case was the same as it was in the case of CO₂–NO₂.

For pure CO₂ injection into the storage site with 22% initial porosity, the WBHP reached its peak after 90 days of injection. In comparison to injection with impurities, the introduction of CO₂–NO₂, CO₂–SO₂ and CO₂–H₂S in each case, showed that maximum pressure was attained relatively faster than in the pure CO₂ case. The WBHP rise suggested consistency with reservoir porosity. Thus, for storage sites with lower porosity, the maximum pressure was attained relatively faster than for sites with higher porosity.

The corresponding well gas injection rate (WGIR) at three observation points is shown in Figure 7. The observation points were considered against the WBHP. These points correspond to the first day of injection.
(observation point index 1), the peak WBHP (observation point index 2) and the last day of injection (observation point 3). For all storage sites, we observed an initial injection rate of $6 \times 10^6$ Sm$^3$CO$_2$/day at observation point 1 and this rate remained constant until the WBHP attained stability when injection rate began to decrease with time until the well was shut.

The injection rate of pure CO$_2$ at the storage site with 10% initial porosity was $1.87 \times 10^6$ Sm$^3$ at observation point 2 (45 days). Since we did not consider a new injection rate at any time after the peak WBHP was attained, for all storage sites, we do not expect to have a significant volume of CO$_2$ to be injected when the well shuts. Therefore, the introduction of impurity, in each case, caused a significantly higher injection rate to be attained. For all the impurity cases, the injection volumes at this observation point were similar.

On the other hand, when pure CO$_2$ was injected in the storage site with 22% initial porosity, an injection rate of $3.1 \times 10^6$ Sm$^3$ was encountered at observation point 2 (90 days). The injection of CO$_2$–NO$_2$ caused a considerably higher injection rate to be encountered at the same observation point (59 days). Furthermore, the introduction of CO$_2$–SO$_2$ and CO$_2$–H$_2$S impurities, in each case, respectively, resulted in a considerably higher injection rate in comparison with the pure CO$_2$ case. It should be noted that these large degrees of difference in the injection rates at observation point 2, for all storage sites, may not necessarily be a strong factor that could cause a considerably lower injection volume in the long term.

The results from WGRG seem to show a correlation with the initial reservoir porosity and permeability, although the effect of porosity appears to be more overwhelming. We observed that a more porous injection site led to higher injectivity. However, this scenario was specific to observation point 2, for all cases.

Figure 6 shows the field pressure (FPR) at three observation points for sites with 10% and 22% initial porosity. The FPR was observed with respect to WBHP on the first day of injection (observation point index 1), peak WBHP (observation point index 2), and at the end of injection period (observation point index 3). For all cases simulated, we observed an increasing FPR from an initial pressure of approximately 124 bar, and this increases gradually, without stabilising at observation point 2 until a peak pressure of 274 bar was attained. The peak FPR attainment
time varies for each simulation case, and such pressure is always reached before the end of injection when the well is shut. The variation in FPR for various storage sites was a result of differences in the change in hydrostatic pressure which is affected by the bulk properties of the reservoir.

The FPR for pure CO₂ injection at the storage site with 10% initial porosity was 263 bar at observation point 2 (45 days) and 273 bar at observation point 3. When CO₂–NO₂ was injected, there was a reasonable decrease in FPR at observation point 2 (31 days) and nearly the same pressure was maintained as in the pure CO₂ case at observation point 3. Additionally, the injection of CO₂–SO₂ caused a considerable decrease in FPR at observation point 2 (59 days), while similar FPR was encountered in observation point 3, relative to the pure CO₂ injection case. Lastly, the injection of CO₂–H₂S also showed a significant decrease in FPR at observation point 2 (59 days), relative to the pure CO₂ case.

The well gas injection total (WGIT) for sites with 10% and 22% initial porosity is presented in Figure 9. This is also the same as the cumulative injection. The observation points were considered against the WBHP on the first day of injection (observation point index 1), the peak WBHP (observation point 2) and at the end of injection period (observation point 3). For all cases simulated, we observed an increasing CO₂ injection until the well was shut. Besides the effect of impure CO₂, which was manifest at observation point 2, the initial porosity of the storage site significantly controlled the cumulative injection.

Figure 7. Well gas injection rate (WGIR) for pure and impure CO₂ storage for sites with 10% initial porosity ((a) CO₂–NO₂, (b) CO₂–SO₂, (c) CO₂–H₂S) and 22% initial porosity ((d) CO₂–NO₂, (e) CO₂–SO₂, (f) CO₂–H₂S).
both water and air, changes in viscosity, and interfacial tension, could all affect the reservoir performance to a certain degree. Though, the main cause of differences in the pressures and volumes (besides the properties of the storage formation) between pure and impure CO2 injection is controlled by the volume of impurity within the CO2 stream. Additionally, we observed that for all impurity cases simulated, the difference between target and actual injection rates is minimal and this can be due to the spatial variance in reservoir properties such as porosity and permeability and the implicit use of constant values for these properties in the model. Thus, the influence of non-linearity and heterogeneity as obtained in realistic settings is not considered in the model. This minimal difference between target and actual injection rates for the simplified models has been observed in a previous study on the BSF [25]. It is important to note that for all injection cases simulated, we monitored the cumulative CO2 injected both at the end of injection and at the end of monitoring periods, without any noticeable discrepancy. Nonetheless, the use of simplified models will remain helpful in the evaluation of opportunities for CO2 storage deployment in the UK offshore reservoir systems as a viable step toward decarbonisation of the global economy [25]. Due to its large areal extent, the BSF conforms with the use of simplified models to predict both regional and local reservoir behaviour in response to different injection scenarios.

4. Conclusions

The presence of impurities in the CO2 stream can significantly affect the physical quality of the storage reservoir and it may also alter the molar volumes and shift the operational boundaries in the CO2 phase diagram, affecting the limits of stability during injection operations. The main aim of this study was to evaluate the effect of CO2 and impurities on reservoir performance during injection and storage in a sandstone saline aquifer. These included the evaluation of key performance indicators such as bottom-hole pressure, well gas injection rate, field pressure and cumulative injection for each impurity case. The results suggested that the effects of impurities influenced the reservoir performance but mostly at the early injection stages, but not the targeted injection rate. It was also

Figure 8. Field pressure (FPR) for pure and impure CO2 storage for sites with 10% initial porosity ((a) CO2–NO2, (b) CO2–SO2, (c) CO2–H2S), and 22% initial porosity ((d) CO2–NO2, (e) CO2–SO2, (f) CO2–H2S).
found that the porosity of the BSF significantly affected CO2 injection. Additionally, it was observed that the effect of impurities does not affect the security of stored CO2 as neither the injected volume nor field pressure were found to decrease 30 years after injection had ceased. We expect that the conclusions from this study can be extended to other saline aquifers with similar depositional- and post-depositional history.

Declarations

Author contribution statement

Mohammed Dahiru Aminu: Conceived and designed the experiments; Performed the experiments; Analyzed and interpreted the data; Wrote the paper.
Vasilije Manovic: Analyzed and interpreted the data.

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Competing interest statement

The authors declare no conflict of interest.

Additional information

No additional information is available for this paper.
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