CO₂ dissolution and convection in oil at realistic reservoir conditions: A visualization study

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Abstract

CO₂ convective mixing has been extensively studied for CO₂ dissolution in saline water but very limited with the presence of oil. The objective of this work was to visually study the supercritical CO₂ (sCO₂) dissolution and convective mixing into oil at realistic reservoir temperature and pressure conditions with and without the presence of porous media.

A specially designed high-pressure 2D-cell was used to investigate the sCO₂ mixing into oil phases. Schlieren imaging method was used as the visualization method. The experiments were carried out at 100 bar and 50 °C using n-octane, n-decane, and crude oil as the main oils. Porous media with different permeability was prepared using glass beads.

Convective fingering was found to accelerate the mixing of CO₂ with n-octane and n-decane. It was not possible to visualize the CO₂ convective fingering in crude oil due to the low opacity of the oil phase. The CO₂ dissolution into oil phases was quite instantaneous and fast without the presence of porous media. The swelling of oil was measured as 55%, 50% and 11% for n-decane, n-octane and crude oil respectively without the presence of porous media. Boundary effects were affecting the CO₂ mixing due to the circular shape of the 2D-cell. Having a water layer below the oil layer tends to dampen the CO₂ transport from the oil phase to the water phase. CO₂ dissolution into oil saturated porous media was slower compared to that without the presence of porous media. The mixing of CO₂ was faster at higher permeability than at lower permeability. Visualization of CO₂ convective mixing/fingers inside oil-saturated porous media using a Hele-Shaw cell yet to be achieved experimentally.

1. Introduction

For the past few decades, global temperature has been rising beyond human control mainly due to Greenhouse gas (GHG) emissions. Effects of global warming have started affecting day to day life of all living beings and ecosystems on the planet (Gale and Kaya, 2003). Global organizations and different authorities have been trying to come up with directives and legalization to mitigate climate change by reducing greenhouse gas emissions (Mintzer, 1987; United Nations, 2015; European Commission, 2019). Geological storage of CO₂ of captured CO₂ through an environmentally safe path is one way to contribute to the fight against climate change (Gibbins and Chalmers, 2008).

It is very important to optimize and improve the understanding of the Carbon Capture and Storage (CCS) value chain to maximize CO₂ geological storage with minimum cost (Budinis et al., 2018). Injection of CO₂ to ongoing and abandoned oil fields is a well-identified solution for the commercial utilization of CO₂. During CO₂ injection into existing oil fields for Enhanced Oil Recovery (EOR), the added CO₂ will increase the oil recovery percentage. EOR for CO₂ utilization can also reduce a significant cost of the whole CCS value chain (Brock and Bryan, 1989; Blunt et al., 1993).

After CO₂ injection into the reservoir, CO₂ will be trapped by four mechanisms which can be identified as structural trapping, residual trapping, mineral trapping, and solubility trapping (Zhang and Song, 2014; Zhang et al., 2016; Sun et al., 2020). Solubility trapping is one of the significant mechanisms for long term CO₂ storage and EOR (Bachu, 2008; Zhang and Song, 2014). During CO₂ EOR, CO₂ dissolution in oil and/or water will result in, oil swelling, viscosity reduction, and in addition to wettability alteration by reduced pH (Sohrabi et al., 2009; Fjelde and Asen, 2010). Once the injected CO₂ is mixed with oil due to diffusion, the swelling will build up differential pressure at the pore level to move the oil drops or ganglia out of the pore space, accompanied by
Reduced viscosity. The mixing process can be further accelerated by the gravity difference between the original oil in place and the oil-containing CO$_2$ which may have higher density, thus leading to the convection-driving dissolution of CO$_2$ as shown in Fig. 1 (Mojtaba et al., 2014; Gasda and Elenius, 2018).

The convection-driven dissolution has been extensively studied experimentally for accelerated CO$_2$ dissolution in saline water under gravity for CO$_2$ storage inside 2D Hele-Shaw cells or 3D cylindrical cells (Song et al., 2003; Neufeld et al., 2010; Kneafsey and Pruess, 2011; Khosrokhabhar et al., 2014; Mojtaba et al., 2014; Vosper et al., 2014; Faisal et al., 2015; Taheri et al., 2018; Thomas et al., 2018; Mahnoodpour et al., 2019; Amarasinghe et al., 2020). Detail review of those studies can be found in Amarasinghe et al. (2020). Furthermore, numerically CO$_2$ convective dissolution have been studied widely with different scales and conditions (Farajzadeh et al., 2007; Hassanzadeh et al., 2009; Pau et al., 2010; Cheng et al., 2012; Elenius et al., 2014; Vosper et al., 2014; Emami-Meybodi et al., 2015; Gasda and Elenius, 2018). Elenius and Gasda (2021) have specifically investigated CO$_2$ convective mixing which is driven by non-monotonic density. In their numerical investigation they have investigated CO$_2$ convection behaviour in oil depend on its originated location, i.e. from above or below the oil zone. They have concluded that, CO$_2$ from above, provides a classical convective mixing with acceleration (due to viscosity decrease) while CO$_2$ from below flows upward (due to buoyancy), but is countered by downward convection due to the heavier mixture density. Furthermore, they have concluded that bottom convective system is notably more complex and efficient than from above.

![Fig. 1. Simplified sketch of convection-driving dissolution of CO$_2$ with oil inside the reservoir. CO$_2$ plume under the caprock is shown in yellow colour. Enlarge image shows how CO$_2$ from the plume mix with reservoir fluids.](image)

To make the CO$_2$ convective flow inside fluids (water or oil) analogous to 2D, Hele-Shaw cells are used. (Hartline and Lister, 1977). Hele-Shaw cells are constructed by two parallel flat plates with a small gap in between (generally less than 10 mm). The thickness of the cell should be enough to allow a clear visualization. The Rayleigh number (Ra) should not be less than or equal 4$\pi^2$ to get substantial natural convection (Lindeberg and Wessel-Berg, 1997; Faisal et al., 2015). Ra number is a dimensionless ratio between free convection to diffusion as shown in Eq. (1). Here $\Delta \rho$ is the density increase of oil and/or water due to CO$_2$ dissolution, $g$ is the acceleration of gravity, $k$ is the permeability of the porous media, $H$ is the height of porous media or the bulk fluid, $\mu$ is the dynamic viscosity of oil or water, $D$ is the molecular diffusion coefficient of CO$_2$ in oil or water and $\phi$ is the porosity of porous media. If a bulk fluid is being used, permeability ($k$) is calculated by Eq. (2) where $b$ is the cell thickness while porosity ($\phi$) is considered as 1.

$$Ra = \frac{\Delta \rho \times g \times k \times H}{\mu \times D \times \phi}$$  \hspace{1cm} (1)

$$k = \frac{\rho^2}{12}$$  \hspace{1cm} (2)

Many experimental studies have been carried out using surrogate fluids at atmospheric conditions to mimic actual CO$_2$ and water inside the experimental setup (Neufeld et al., 2010; Taheri et al., 2012; Tsai et al., 2013; Agartan et al., 2015; Wang et al., 2016; Teng et al., 2017). Few studies have conducted CO$_2$ convective dissolution in bulk water using actual CO$_2$ (Song et al., 2003; Kneafsey and Pruess, 2011; Khosrokhabhar et al., 2014; Mojtaba et al., 2014; Faisal et al., 2015; Thomas et al., 2015; Lv et al., 2016; Lu et al., 2017; Taheri et al., 2018; Teng et al., 2018). Vosper et al. (2014) and Mahnoodpour et al. (2019) have conducted CO$_2$ convective dissolution into water-saturated porous media using a 2D Hele-Shaw cell under gas conditions. Amarasinghe et al. (2020) have conducted their experiments inside a Hele-Shaw cell at supercritical conditions (100 bar, 50°C) where CO$_2$ was dissolved into water-saturated porous media. Wei et al. (2017) have carried out a visualization study on oil swelling due to CO$_2$ miscibility inside a high-pressure cylindrical cell under reservoir conditions.

However similar studies with the presence of oil (or residual oil) are still very limited. This represents a gap in defining and validating the adequate mathematical models and upscaling procedures for CO$_2$ storage and EOR, and the lack of input parameters for uncertainty estimation. The only study in the literature where CO$_2$ was convectively dissolved into the oil phase with visualization inside a Hele-Shaw cell, was conducted by Khosrokhabar et al. (2014). In their study, they have used a high-pressure cell which is made from two glass windows of 2.5 cm diameter and a thickness of 11.6 mm. They have visualized gravity-induced convective dissolution fingers of CO$_2$ into the n-decane phase by the Schlieren visualization method at 39°C and 64 bars where CO$_2$ behaves as a sub-critical phase. They have observed a highly unstable and turbulent behaviour when CO$_2$ comes in contact with the oil phase. But clear distinguishable fingers are hard to identify in their images probably due to the small diameter of the cell and/or usage of a black and white camera. So they have left further interpretation of the results and images as further work. Amarasinghe et al. (2021b) have investigated CO$_2$ breakthrough time through an oil-saturated unconsolidated porous media at 50°C/100 bar due to CO$_2$ convective mixing. They have used n-decane and crude oil with different permeable porous packs and have come up with a relationship between Ra number and dimensionless time and Ra number and CO$_2$ transport rate. Furthermore, they have experimentally found out that pressure decay in the CO$_2$-crude oil system is very low compared to the CO$_2$-n-decane system. Doranegard and Dehghanpour (2020) and (Yassin et al., 2018) have visually investigated nonequilibrium CO$_2$/oil interactions at CO$_2$/oil surfaces at reservoir conditions. They have investigated oil swelling as well as CO$_2$ convective fingers around the CO$_2$/oil interface.
Furthermore, they have observed that immediate oil expansion upon introduction with CO\textsubscript{2} and they have observed that sCO\textsubscript{2} provide larger oil expansion compared to liquid and gaseous CO\textsubscript{2}. Zhao et al. (2011) have investigated and visualized CO\textsubscript{2} flooding in porous media inside a high-pressure PVT cell at 85 bars and 40 °C using MRI technology. Meanwhile, Seyyedsar and Sohrabi (2017) have visually investigated the formation of a new oil phase during immiscible CO\textsubscript{2} injection into heavy oil-saturated porous media under reservoir conditions. Liu et al. (2016) have experimentally investigated CO\textsubscript{2} diffusion into bulk n-decane and n-decane saturated porous media using micro-CT visualization. They have found out that initial pressure has a significant effect on the diffusion coefficient. Furthermore, due to the complications of the diffusion path in n-decane saturated porous media, they have found out that CO\textsubscript{2} mass transfer and diffusion coefficient reduces. Czarnota et al. (2018) and (Janiga et al., 2020) have investigated CO\textsubscript{2} mass transfer into the oil phase at reservoir conditions. They have used acoustic pulse-echo techniques to measure the oil volume expansion due to the CO\textsubscript{2} solubility in the oil phase. Meanwhile, Rezk and Foroozesh (2018) have estimated mass transfer parameters numerically for CO\textsubscript{2}-crude oil systems. They have considered the oil swelling effect with a moving boundary and estimated that mass transfer co-efficient, diffusion co-efficient and oil swelling percentage are higher in light oil-CO\textsubscript{2} systems compared to heavy oil-CO\textsubscript{2} systems. Mahzari et al. (2018, 2019) have experimentally investigated CO\textsubscript{2} EOR efficiency though carbonated water injection using a micromodel system with the presence of live oil. They have visually confirmed CO\textsubscript{2} transfer into resident oil and eventual swelling of oil phases. Meanwhile, Amarasinghe et al. (2021a) have conducted CO\textsubscript{2} convective mixing visualization experiments in a micromodel apparatus using 3-phase fluid system (CO\textsubscript{2}-water-oil) at reservoir conditions. In there, they have shown how CO\textsubscript{2} mixing with both diffusion and convection driven mechanism with the presence of 3-phase and how CO\textsubscript{2} migrate vis specific pore structures.

In this paper, we have carried out a visual investigation of CO\textsubscript{2} convective dissolution into oils with and without the presence of porous media inside a high-pressure 2D Hele-Shaw cell. This is the first time these kinds of experiments have been carried out in supercritical conditions using oil. Furthermore, in this study we have investigated CO\textsubscript{2} convective fingering in detail compared to Khosrokhavar et al. (2014).

**Fig. 2.** Front view of the actual experiment setup.

**Fig. 3.** Cross-section of the high-pressure cell highlighting the test volume, filter locations and gas injection location.
Fig. 4. 3D sketch of the high-pressure cell including main components.

Fig. 5. Particle size distribution of the glass beads.

Fig. 6. Piping and instrumentation (P & ID) diagram of the experimental setup.
Furthermore, this is the CO$_2$ convective experiments have been carried out with the presence of porous media together with oil which none of the previous studies have been unable to address. A practical outcome of the experimental results will be the visual quantification of CO$_2$ convective fingers inside oil together with oil swelling. The experiments results will be important in identifying field-scale impacts and

Fig. 7. Schematic sketch of the Schlieren imaging setup.

Fig. 8. Case 1 – Visual snaps from CO$_2$/n-decane system without porous media. The red dotted line at $t=0$ s indicates the initial CO$_2$/n-decane interface.
assessement of reservoir conditions through numerical simulations under which oil and gas recovery can be improved while simultaneously increasing the long-term CO$_2$ storage potential.
Table 1
Set of experimental cases at 50 °C and 100 bars.

| Test # | Fluid(s) | Porous media | Visual results |
|--------|----------|--------------|---------------|
| Case 1 | n-decane | –            | Figs. 8 and 9 |
| Case 2 | n-octane | –            | Fig. 10       |
| Case 3 | n-decane and water | – | Fig. 13 |
| Case 4 | crude oil | –             | Fig. 11       |
| Case 5 | n-decane | 76 D         | Fig. 14       |
| Case 6 | n-decane | 500 mD       | Fig. 15       |
| Case 7 | crude oil | 76 D         | Fig. 16       |

Fig. 9. CO₂ convective fingers at t=14 s for n-decane system without porous media (case 1). Distinguishable fingers are marked in arrows and widths are given in Table 2. White circles indicate how CO₂ mix along the boundary due to boundary effects.

2. Method

2.1. High-pressure 2D cell

The high-pressure 2D-cell was developed using stainless duplex steel and borosilicate glass. The 2D cell was designed for maximum working conditions of 150 bars and 100 °C. The test cell consists of several units, which are assembled to construct the final cell. The cell volume was formed by two glass plates separated by an adjustable spacer. A heating cable was wrapped around the cell with an automatic temperature controller. The front view of the real cell with insulation is shown in Fig. 2. A 3D sketch of the cell is shown in Fig. 4 while a cross-section of the cell is shown in Fig. 3. A specially designed filter module of shaped glass filter (pore size of 10–16 μm) plate was placed at the inlet and outlet to avoid penetrating the glass beads away from the test volume (see Figure 3 and 4). The location of the gas injection was placed in the middle of the top filter module unit as shown in Fig. 3. The diameter of the test volume is 200 mm while the sight disk glass diameter was 100 mm. The thickness is 5.0 mm. Amarasinghe et al. (2020) has provided further details of the high-pressure 2D cell and its design specifications.

2.2. Materials

2.2.1. Chemicals

N-decane, n-octane, and a North Sea crude oil were used as oils. To improve the visualization of the oil phase (with n-decane and n-octane), the oil-soluble dye Sudan III (0.04 wt%) was added to the oils. Aqueous 0.04 wt% bromothymol blue (BTB) pH indicator solution (deionized water mixed with 0.01M NaOH and BTB) of pH around 8 was used as the water phase. The BTB pH indicator is blue above pH = 7.6 and yellow below pH = 6.5, and green between these pH-values (De Meyer et al., 2014). When CO₂ is dissolved in water, the pH will be reduced due to the release of H⁺ ions which will lead to the colour change in the water phase from blue to yellow (Thomas et al., 2015). Due to the high light intensity, the colour transition to yellow was visible as dark black colour.

2.2.2. Porous media

Hydrophilic glass beads of different diameters were used to prepare porous media of different permeabilities (500 mD and 76 D). Fig. 5 shows the particle size distribution of each porous media type. The porosity and the permeability of the bead packs was determined by the waterflooding of tubes with packed porous media.

2.3. Experimental procedure

The experimental set-up with the high-pressure 2D test cell is shown in the piping and instrumentation diagram (P & ID) (see Fig. 6). A back-pressure regulator was included in the set-up to avoid sudden developments of high pressures inside the cell. CO₂ was introduced from a piston cell. A digital manometer was connected directly to the 2D cell for accurate reading of the absolute pressure.

All the experiments were carried out at 50 °C and 100 bars. 2D Hele-Shaw cell was pre-heated using the heating cables up to 50 °C. All the fluids were pre-heated inside a heating cabinet (at 50 °C) before adding them into the Hele-Shaw cell. To observe CO₂ convective dissolution in oil, a known volume of oil (n-decane, n-octane, crude oil) was filled into the cell manually. In the CO₂ dissolution experiment with n-decane, a water layer at the bottom was first filled into the cell followed by n-decane.

For the experiments with porous media, the test volume is filled with a known volume of oil first. And then, dry glass beads were filled manually from the top of the cell gradually so that the beads will settle smoothly inside the oil phase (filling dry glass beads first and flood oil through to saturate the pack was not possible due to uneven packing and possible cracking inside the porous media). When the desired level of porous media was achieved, small external vibration was applied to the cell by using a plastic hammer to further settle the glass beads. If any oil was above the porous media, it was sucked out using a syringe. After placing the filter module on the top of the cell, the cell was left for approximately half an hour to get even packing of the homogenous porous media and stable temperature (confirmed from a thermal image by a FLUKE® Ti25 thermal imaging camera).

The CO₂ piston cell was first filled to 80 bars at room temperature and then heated to 50 °C. After the gas expansion, the piston cell was pressurized to 105 bars. The fine valve connected to the CO₂ piston cell (see Fig. 6), was then opened slowly to allow CO₂ to flow into the 2D cell to avoid the splash of CO₂ into the liquid phase and movement of glass beads.

The visualization observations were carried out using the Schlieren method using a Nikon D5200 camera with video recording. ImageJ open-source image analysis software (Rueden et al., 2017) was used to analyze the width of CO₂ convective fingers and oil swelling due to CO₂ mixing. To measure the oil swelling, the height increase of the free oil phase was measured using the software and was converted to the volume.

2.4. Schlieren method

2.4.1. Theory of schlieren method

The Schlieren method has been used by (Thomas et al. 2015, 2018)
to visualize CO$_2$ convective dissolution in water and salt solutions inside a Hele-Shaw cell at ambient conditions. Detecting the deflection of the light beam sent via the cell is the basic principle of the Schlieren method. The density gradients occur due to the fluid mixing (CO$_2$/Oil in this case) is the reason for the deflection of the light beam. One of the drawbacks of the Schlieren method is that it can be only applicable to bulk fluid flows setup. i.e. cannot be applied with the presence of porous media. See Settles (2001) and Bunton et al. (2016) for more general information on the Schlieren method.

To visualize flow inside a 2D Hele-Shaw experimental setup, the Schlieren method can be applied in several ways depending on how the concave mirrors or convex lenses are used.

Method 1: Two convex lenses method (Khosrokhabar et al., 2014; Thomas et al. 2015, 2018)
Method 2: Two concave mirrors method (Eder and Jordan, 2001; Settles, 2001)
Method 3: Single concave mirrors method (Eder and Jordan, 2001; Settles, 2001)

2.4.2. Application of schlieren method for this study

As our visualization window of the 2D Hele-Shaw cell is large (i.e. 10 cm), above mentioned (see subchapter 2.4.1) two Schlieren visualization methods were not possible to use because of the requirement of either two concave mirrors or two convex lenses with a larger diameter. Due to the lack of space in our lab, we (have opted for a Schlieren visualization method that uses only a single concave mirror. According to the requirement, a single concave mirror of diameter 20.32 cm and an effective focal length of 162.56 cm was used.

The mechanism of the Schlieren method using a single concave mirror is described as follows (see Fig. 7). A point light source is applied from the front side of the cell as the main light source. The light source was placed at 2 times the focal length of the concave mirror. Before the start of the experiment, the deflected light beam coming from the concave mirror was converged to the pre-arranged knife-edge where the camera was focused on the converging light source.

During the experiment, light rays are deflected due to the refractive index variation caused by the CO$_2$/oil mixture. The light rays coming through the experimental 2D cell glass are scrambled due to the CO$_2$ concentration variation. These oncoming light rays will not exactly be focused on the two times of the focal length of the mirror but on a focal plane to the mirror. So, by adjusting the knife-edge the perfect focal point can be found so that any light rays will not get lost which could lead to loss of the image intensity. Then to obtain the images the camera was used. In Fig. 7, a schematic sketch of the applied Schlieren method for this experiment is shown.

2.5. Set of experiments

Seven sets of CO$_2$ convective dissolution experiments were carried out as shown in Table 1.

3. Results and discussion

3.1. Visual observations of CO$_2$ dissolution into the oil without porous media

CO$_2$ was introduced to the oil phase of n-decane (Fig. 8). Due to the time delay in the opening of the fine valve slowly, CO$_2$ starts to have the first contact with the oil phase at $t = 8$ s. Then after, sudden development of CO$_2$ fingering was observed into the oil phase (see $t = 10$ s in Fig. 8). The fingers were observed drawn towards the bottom of the cell due to the gravity-driven mechanism until 17 s. In Fig. 9, an enlarged image of CO$_2$ convective fingers at $t = 14$ s is shown. The width of the fingers was counted and measured (see Table 2). Within a 5 cm span, 9 fingers were distinguished. The width of the fingers was observed as random. Meanwhile, the fingers were observed to merge with time, and the width of the fingers and locations were continuously changing.
The circular boundary affects the direction of CO$_2$ fingers downwards (see CO$_2$ flow along the boundary in Fig. 9 circled in white colour). At $t = 17$ s and $t = 19$ s, CO$_2$ reaches the bottom of the cell along the boundary before the middle fingers reach the bottom. At $t = 21$ s the CO$_2$ came along the left and right side of the boundaries merge with the centre fingers. At $t = 26$ s, the merged convective flow of CO$_2$ reaching upwards through the centre of the cell was observed.

From $t = 33$ s to $t = 88$ s, a disturbance to the oil phase was observed due to the phase change of CO$_2$, i.e. from the time of opening the fine valve slowly and up to $t = 33$ s CO$_2$ behaves as a gas phase. With time when CO$_2$ filled up the 2D cell space, CO$_2$ changes its phase-type from gas to liquid followed by the supercritical phase. From $t = 171$ s (after the disturbance) CO$_2$ convective fingers were circulating inside the oil phase. Meantime more CO$_2$ was dissolved into the oil phase. This can be observed by comparing the oil level at $t = 8$ s and $t = 3000$ s. With time the CO$_2$ convective fingers got less concentrated which indicated that the oil phase was getting fully saturated with CO$_2$. At $t = 1170$ s, less amount of CO$_2$ convective fingers can be observed. At $t = 3000$ s a fully saturated oleic phase with CO$_2$ can be observed. Due to the presence of the high-pressure CO$_2$, the dissolution process was instantaneous. N-decane swelling was calculated at the end of the experiment to be 55%.

The same behaviour as for n-decane was observed for n-octane (see Fig. 10). In Fig. 10 images from disturbance that occurred due to the phase change of CO$_2$ were omitted which happens between $t = 12$ s and $t = 60$ s. It was observed that CO$_2$ dissolution in n-octane was faster than for n-decane. The whole convective dissolution process was finished in about half time compared to n-decane for the same amount of oil volume. N-octane swelling was calculated at the end of the experiment to be 50%.

CO$_2$ convective dissolution into crude oil experiment was also conducted (see Fig. 11). Due to the thick dark colour intensity of crude oil, light cannot penetrate the crude oil. So, CO$_2$ convective dissolution fingers and convection flows cannot be identified using the Schlieren method. The swelling of crude oil was calculated as 11% at the end of the experiment.

3.2. Visual observations of CO$_2$ dissolution into the oil with a water layer without porous media

Another experiment was carried out with n-decane on top of the
water phase. Experimental images at different stages are given in Fig. 13. First, at the introduction of CO$_2$, CO$_2$ fingering was observed in n-decane (as explained earlier). At $t = 15$ s, CO$_2$ came along the boundaries level off on the top of the n-decane/water fluid’s plane. The merged CO$_2$ together with incoming CO$_2$ fingers along the middle of the oil phase started convective fingers upwards. At $t = 42$ s, CO$_2$ convective fingers into the water phase were observed (notice the small black fingers right below the water phase at $t = 42$ s). Several fingers were developed downwards up to $t = 59$ s while several fingers were merging when they reached the bottom. Boundary effects also were observed when CO$_2$ was dissolved into the water phase. Similar observations were made by Amarasinghe et al. (2020) when they visualized CO$_2$ dissolution into the water phase at the same pressure and temperature conditions (50 °C and 100 bar). From $t = 49$ s to $t = 135$ s, the water phase became saturated with CO$_2$ which changed the colour of the water phase from blue to yellow. Meanwhile, CO$_2$ convective fingers kept flowing in n-decane, and swelling of the oil phase continued. At $t = 1200$ s almost all the n-decane phase was observed saturated. Since 50 ml of n-decane was used compared to the previous case with 100 ml, the whole CO$_2$ convective dissolution process was finished faster, approximately in half the time with 100 ml n-decane. (see Fig. 12)}
3.3. Visual observations of CO$_2$ dissolution into oil with porous media

CO$_2$ convective dissolution into porous media was observed for 3 different cases. In the first, CO$_2$ was introduced to 100% n-decane saturated 76 D porous media (55 ml of n-decane was used). In the beginning, the levels of the porous media and n-decane were the same (see t = 0 min in Fig. 14). As mentioned in the introduction, the Schlieren method can only be applied for bulk fluids. Since light does not penetrate through porous media, CO$_2$ convective fingers inside porous media cannot be obtained using the Schlieren method. During this study, we have attempted to add a CO$_2$ soluble dye (C.I. Disperse Red 60) in the CO$_2$ phase in order to track CO$_2$ transport in oil-saturated porous media. But due to the lower solubility of the dye in CO$_2$ at 100 bar (our working pressure), the attempt was unsuccessful (Gupta and Shim, 2007).

When CO$_2$ was dissolved into the n-decane phase with time, the n-decane phase began to swell and produce a free n-decane phase above the porous media. CO$_2$ convective flows were observed inside the swelled free n-decane (see at t = 5 min in Fig. 14). With further time at t = 10 min and t = 20 min, n-decane kept swelling as well as CO$_2$ convective flow kept circulating inside the oil phase. At t = 170 min, CO$_2$ swelling has stopped along with further CO$_2$ convective flows in the oil phase which indicated all the oil inside porous media, as well as swelled oil, has been fully saturated with CO$_2$. A total oil swelling of 46% was calculated from the image analysis. Fig. 17 shows the oil swelling percentage as a function of time for the experiment case 5.

| Finger # | The width of the finger (mm) |
|----------|-----------------------------|
| 1        | 9.25                        |
| 2        | 3.80                        |
| 3        | 3.20                        |
| 4        | 5.30                        |
| 5        | 4.20                        |
| 6        | 4.85                        |
| 7        | 4.56                        |
| 8        | 8.26                        |
| 9        | 7.32                        |

Table 2: CO$_2$ convective finger’s width at t = 14s for the n-decane system without porous media (case 1) as per Fig. 9.

Fig. 13. Case 3 - CO$_2$/n-decane/Water system without porous media. The red dotted line at t=4 s indicates the initial CO$_2$/n-decane interface.
A similar experiment was carried out for n-decane (55 ml) saturated porous media of 500 mD (see Fig. 15). Similar observations were observed as in 76 D porous media. In order to fully saturate oil with CO₂ and stop the swelling of oil, it took 2000 min in 500 mD porous media which is approximately 117 times slower than in 76 D porous media. A total oil swelling of 32% was calculated from the image analysis. The difference of oil swelling between 76 D and 500 mD was understood as due to the small differences in the porosity.

CO₂ convective dissolution was studied also for 76 D porous media saturated with crude oil (55 ml) (see Fig. 16). Since it was understood that CO₂ convective flow cannot be seen even in the swelled oil phase, general images of the experiment were taken instead of using the Schlieren method. Only a small fraction of oil swelling was observed even after 2000 min of the experiment.

4. Suggested future work

Visualization of CO₂ convective mixing/fingers inside oil-saturated porous media using a Hele-Shaw cell yet to be achieved experimentally. With the use of high-tech such as X-ray or MRI technology, it may be possible to visualize CO₂ convective fingering inside porous media. Moreover, usage of reservoir rock types is recommended to investigate both CO₂ convective mixing and fluid-rock interactions together. Furthermore, the experimental results will be used to improve the numerical simulation which are developed using the Open Porous Media (OPM) framework (Lauser, 2014; Rasmussen et al., 2020).

5. Conclusions

- The available high-pressure 2D-cell and Schlieren imaging method have made it possible to investigate CO₂-dissolution and convective mixing into oil phases at pressures and temperatures realistic for reservoirs with CO₂-EOR and CO₂-storage.
- Convective fingering was found to contribute to the mixing of CO₂ into n-octane and n-decane without porous media. CO₂ convective fingering was not possible to visualize with crude oil due to the low opacity of the oil phase. The CO₂ dissolution into oil phases was quite fast without porous media.
At the end of the CO\textsubscript{2} mixing process, the swelling of the oil phase was 55% for n-decane and 50% for n-octane. Much less oil swelling was observed for crude oil which was calculated as 11%.

Boundary effects were affecting the CO\textsubscript{2} dissolution process due to the circular shape of the 2D-cell.

The CO\textsubscript{2} dissolution into oil-saturated porous media was slower compared to without porous media. The mixing of CO\textsubscript{2} was faster at higher permeability than at lower permeability.

Fig. 14. Case 5 - Experimental snapshots of CO\textsubscript{2}/n-decane system with 76 D porous media. The red dotted line at t=0 s indicates the level of porous media.

Fig. 15. Case 6 - CO\textsubscript{2}/n-decane system with 500 mD porous media. The red dotted line indicates the level of porous media.

- An oil phase above the water phase tends to damp the CO\textsubscript{2} dissolution transition process from the oil phase to the water phase.
- The CO\textsubscript{2} dissolution into oil-saturated porous media was slower compared to without porous media. The mixing of CO\textsubscript{2} was faster at higher permeability than at lower permeability.
• Oil swelling of 46% and 32% was observed with n-decane saturate porous media with 76 D and 500 mD respectively. CO$_2$ mixing process was 117 times slower in 500 mD compared to 76 D to fully saturate oil with CO$_2$ and stop the swelling of oil.
• The experiment results will further make it possible to better define and validate the simulation model for upscaling procedures for CO$_2$ storage and EOR.

Author statement

Widuramina Amarasinghe: Conceptualization, Methodology, Validation, Investigation, Writing, Visualization, Formal analysis. Ingebrit Fjelde: Conceptualization, Methodology, Validation, Investigation, Writing, Supervision, Funding acquisition. Ying Guo: Conceptualization, Methodology, Writing, Supervision, Project administration, Funding acquisition

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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