SAG Foam Flooding in Carbonate Rocks

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Abstract

Foam is used in gas-injection EOR processes to reduce the mobility of gas, resulting in greater volumetric sweep. SAG (Surfactant Alternating Gas) is a preferred method of injection as it results in greater injectivity in the field, but designing a successful process requires knowledge of foaming performance at very high foam qualities (gas fractional flows).

Here the use of foam in low-permeability (~1 mD) Indiana Limestone cores for SAG foam applications is studied. Coreflood experiments were performed for a range of foam qualities at high pressure (100 bar), elevated temperature (55°C), high salinity (200,000 ppm) and in the presence of crude oil. The effectiveness of the foam was studied by differential pressure measurements along the core. Foam was still able to form under these stringent conditions, but it was a relatively weak foam (i.e. its ability to reduce gas mobility is modest). Still, further analysis of the experimental results show that the foam was able to maintain mobility control over the displaced phase, thus providing a stable displacement front, and that it can be used in a SAG foam process in these formations.

In addition, further coreflood experiments were carried out using heterogeneous, vuggy Edwards White cores with even lower permeability (~0.5 mD). These experiments were performed to determine whether foaming is possible in heterogeneous media and especially to investigate the effects of disconnected vugs on the foaming performance. CT scans were taken during the period of foam injection to determine saturation profiles within the core. Foam was able to form inside these cores, but inside the vugs foam segregation was observed with liquid pockets visible in the bottom of the vugs and gas in the remainder. This segregation was only a local effect though, confined to the vug itself, and foam was able to persist in the rest of the core.

Introduction

A common method of improving oil recovery from a reservoir is through the injection of gas in order to displace oil and maintain pressure. The main problem with gas injection is that it suffers from poor sweep efficiency. Reasons for this include viscous fingering (due to the mobility contrast between the injected gas and the displaced phase), channeling (gas prefers to flow through high-permeability layers, thus low-permeability layers remain unswept) and gravity override (the injected gas has a lower density than the oil in place, therefore flowing to the top of the reservoir, causing an unswept lower region in the reservoir). Foam can help alleviate these problems by trapping the gas in bubbles, thereby reducing its mobility. This results in a more stable displacement front and thus an increase in volumetric sweep (Schramm (1994); Rossen (1996)).

In this study we focus on the application of foam in SAG injection in carbonate formations. These rocks pose some specific challenges which need to be overcome in order for foam to be an effective method of enhancing oil recovery. One of the main problems with carbonate reservoirs is that they typically have low permeability compared to sandstone reservoirs. Ehrenberg and Nadeau (2005) provide average values of porosity and permeability for over 40,000 reservoirs worldwide. As shown in Figures 1a and b, they find that carbonates on average have a significantly lower porosity than sandstone reservoirs, and a lower average permeability as well. The effectiveness of foam in controlling gas mobility is greater in higher-permeability media, as was first identified by Bernard and Holm (1964). Other studies, including Falls et al. (1988); Khatib et al. (1988) and Moradi-Araghi et al. (1997) also find reduced effectiveness of foam in lower-permeability media. Still, Szlendak et al. (2012) found that in-situ foaming enabled mobility control in tight (~10 mD) formations.
Figure 1: Global average (a) porosity vs. depth and (b) porosity vs. permeability plots for sandstone and carbonate reservoirs. The lines respectively represent the P10, P50 and P90 values (from Ehrenberg and Nadeau (2005)).

Foam has been studied extensively in heterogeneous rocks. When it comes to large-scale heterogeneities, such as layered reservoirs, foam injection is often considered as an enhanced oil recovery method. This is because of the aforementioned advantage of foam that it reduces the mobility more in high- than in low-permeability zones.

Another complication in carbonate formations is existence of vugs: i.e., large voids (much larger than pores). Vugs could act as stagnant zones where foam segregates and breaks, as in foam half-life tests in the laboratory. On the other hand, Fernø et al. (2015) showed foam generation during flow into vugs in fracture networks. Interconnected vugs pose a special challenge for foam. Here we consider isolated vugs. Other complications in carbonates not examined here include adverse wettability and natural fractures. Boeije et al. (2015) proposed a screening protocol to identify surfactants capable of foaming in oil-wet porous media.

Here we study foam in a low-permeability carbonate rock. Our main set of foam-flood experiments are performed at high pressure (100 bar back-pressure), elevated temperature (55°C) and high salinity (200,000 ppm). Initially we focus on tests without the presence of crude oil. Foam behaviour at high foam qualities (gas fractional flow) is of special interest, to show whether the foam is suitable for a SAG injection process (Shan and Rossen (2004); Rossen and Bruining (2007)). Rossen and Boeije (2015) show that it is crucial to know the behaviour at high foam quality in order to determine whether a SAG process can be successful. Afterwards we perform experiments with crude oil added to the co-injection fluids to determine its influence on foaming performance. Another experiment is aimed at determining the influence of rock heterogeneity and vugs on foaming performance. This experiment is carried out using a heterogeneous, vuggy carbonate core which is placed inside a CT scanner.

Experimental Setup and Procedure
The setup used for the main set of foam-flooding experiments consists of the following components. The porous medium that is used is a 40-cm-long, 4-cm-diameter consolidated Indiana limestone core in a PEEK core holder. The operating conditions in all of these experiments is the same (100 bar back-pressure, 55°C and 200,000 ppm salinity). The core holder is mounted vertically in a Tamson TV7000 thermo bath which is used to maintain the desired temperature of 55°C inside the porous medium. The setup allows for co-injection of gas, surfactant solution and oil. The gas phase in these experiments is methane (CH₄), which is injected using a Bronkhorst ELFLOW F230M mass flow controller. A Quizix QX-6000 low-flow-rate pump is used to inject the aqueous phase. In some of the experiments described here, crude oil (32° API) was also injected along with the other fluids using a Gilson 305 Piston Pump along with a transfer vessel. Dresser Mity-Mite Model 91 gas-loaded back-pressure regulators were installed to maintain the system at the desired operating pressure. To analyze the pressure inside the core, four differential pressure transducers (Endress+Hauser Deltabar S PMD75, range 0-3
bar) were installed to measure pressure drops within different sections of the core. In addition, another differential pressure transducer (Endress+Hauser Deltabar S PMD75, range 0-40 bar) was installed to measure the pressure drop over the entire core. Absolute pressure transmitters were installed up- and downstream of the core to monitor absolute pressure in the system; they can also be used to check whether the differential pressure measurements are correct. In all of the experiments described here, the flow direction is from the bottom of the core to the top. An overview of the setup used for this set of experiments is given in Figure 3.

Experiments were performed with two different surfactants at a salinity of 200,000 ppm. The surfactants were chosen based on their foaming performance as described in Boeije et al. (2015). We selected a non-ionic alkylpolyglycoside (hereinafter APG) and an anionic alcohol ethoxy sulphate (hereinafter AES), at a concentration of 0.5 wt./wt.% active content. The brine composition is given in Table 1.

Table 1: Brine composition for 200,000 ppm brine.

<table>
<thead>
<tr>
<th>Ion</th>
<th>Concentration [ppm]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cl</td>
<td>122,500</td>
</tr>
<tr>
<td>Na⁺</td>
<td>55,900</td>
</tr>
<tr>
<td>Ca²⁺</td>
<td>19,500</td>
</tr>
<tr>
<td>K⁺</td>
<td>2,100</td>
</tr>
</tbody>
</table>

In addition, another foam-flooding experiment was performed with a shorter, 17 cm long, carbonate core inside a CT scanner (Siemens, 3rd generation). The core was mounted vertically inside the opening of the CT scanner. This limits the size of the core that can be used. The rock used in this experiment was Edwards White, which is a heterogeneous, vuggy carbonate. These experiments were performed to determine the fluid saturations within the core, and in particular to test whether the high level of heterogeneity of the cores, in combination with their vuggy nature, has a significant impact on the foaming performance. We were not trying to mimic reservoir conditions in these CT experiments, but rather to probe foam behaviour in general. Therefore we performed the experiments at room temperature using zero-salinity solutions. For safety reasons we made use of nitrogen gas for the CT experiments rather than the methane that is used in the other experiments in this study. The back-pressure was kept at 100 bar (i.e. the same as the other experiments) to avoid significant effects of compressibility. The surfactant used for this experiment was the non-ionic alkylpolyglycoside (APG) that was also used in the longer-core, Indiana limestone foam floods. A schematic of this experimental setup is given in Figure 4.
Two differential pressure transducers were installed to measure the pressure drop over the entire core and the middle segment of the core, respectively. In addition, two absolute-pressure transducers were used to monitor the pressure at the outlet of the core and at the end of the middle segment. In this way, the pressure in each segment could be monitored separately.

**Procedure**

Procedures for both the long-core and the CT experiments were similar. Foam-flooding experiments are carried out using co-injection of gas and surfactant solution (0.5 wt./wt.% active content). The pressure drop over the core is monitored during the experiment. The heterogeneity in the core in terms of permeability is also assessed by monitoring and comparing pressure drop in the different sections of the core. For the long-core experiments, the main goal is to perform steady-state foam floods at varying foam qualities. This will result in a plot of $f_g$ vs. $\nabla p$ similar to those from various studies (e.g. Ma et al. (2013) and Kapetas et al. (2016)). We are especially interested in foam behaviour at very high foam qualities, because this is crucial for predicting the mobility of the gas front in a SAG-injection process (cf. Rossen and Boeije (2015)). Thus we focus our experiments at foam qualities larger than 0.5, up to foam quality 0.99. We aim to construct fractional-flow curves based on experimental results. To determine whether the foam could be applied in a SAG process, one must find the point of tangency to the fractional-flow curve and the mobility at that point, which corresponds to the mobility of the foam front in a gas injection cycle of a SAG process (Rossen & Boeije, 2015). Because we focus on behaviour in the high-quality regime, we do not expect strongly shear-thinning behaviour and make use of the version of the STARS foam model (Computer Modeling Group, 2006) that incorporates only the water-saturation-dependent function $F_2$. Thus we use the following functional form of the foam model:

$$FM = \frac{1}{1 + f_{mmob} \cdot F_2}$$

where $f_{mmob}$ is a reference mobility reduction factor and the function $F_2$ is defined as:

$$F_2 = 0.5 + \frac{\arctan(epdry \cdot (S_w - f_mdry))}{\pi}$$

For the CT experiment, too, we run the experiment until a steady state is reached. In addition to the overall steady-state pressure response we also observe the distribution of water saturation inside the porous medium. Especially we focus on the behaviour inside and around the vugs, as these are fairly large in size relative to the dimensions of the core, and may have significant impact on the development of foam.

**Results and Discussion**

**Results From Long-Core Experiments**

Before starting foam injection, we measured the steady-state pressure gradient due to a single-phase water flood for different flow rates in order to measure the liquid permeability. The core is fairly homogeneous and has a permeability of 1.01 mD. In a separate test, the porosity of small (3.5 cm length, 3.1 cm diameter) core plugs of the same rock was measured using a Quantachrome Ultrapycnometer 1000 which measures the pore volume of the sample. In total the porosity of 11 core plugs was measured and the average porosity was found to be 0.13.
The main results from the experiments performed using the long Indiana Limestone cores are pressure-drop measurements aimed at determining foam’s ability to reduce total mobility. As mentioned above, the mobility of the foam at the front in a SAG-injection process is defined by a point of tangency to the fractional-flow curve. Results presented here are mainly steady-state pressure gradient over the mid-section of the core, excluding the entrance and exit regions. The foaming performance in the entry and exit regions of the core may not be representative for the overall foaming performance and are thus neglected. For the long-core experiments this means we use the cumulative pressure difference as measured by the four internal pressure transducers.

The results are pressure-gradient measurements at varying foam qualities, but at fixed total superficial velocity, focused primarily on the high-quality regime. We can fit the STARS parameter values using the method outlined in Rossen and Boeije (2015). For the first experiment, we check whether the entrance and exit regions of the core significantly affect the foaming performance by comparing the pressure gradient over the entire core to that in the middle section. Figure 5a shows the pressure-gradient data for the experiment using the APG surfactant and an initial straight-line estimate as a fit. Figure 5b shows the result from the same experiment for the internal section of the core (i.e. the total core except for the entrance and exit regions). The pressure gradient in the latter plot is approximately 25% higher than the result including the entry and exit regions. The section of the core across which the pressure difference was measured is 67% of the total length of the core (26.8 vs. 40 cm) and the pressure drop measured in that section varied from 80 to 85% of the total pressure drop.

This means that the foam in the core as a whole is significantly weaker compared to the mid-section. We only use the pressure result from the mid-section of the core for the remainder of the analysis presented here. The total superficial velocity is kept constant in these measurements at 0.05 ml/min, which for a 4-cm-diameter core is equivalent to $6.63 \cdot 10^{-7}$ m/s (= 0.188 ft./day).

![Figure 5a](image1.png) ![Figure 5b](image2.png)

Figure 5: Steady-state pressure gradient ($\nabla p$) for the experiment using the APG surfactant as a function of foam quality ($f_g$). All data points are in the high-quality regime. Straight-line fit is first estimate and is used to estimate limiting water saturation $S_{w^*}$. (a) steady-state pressure gradients based on the pressure difference over the entire core (b) results without entry and exit regions.

All of the experimental data appear to be in the high-quality regime, which is indicated by the fact that with increasing foam quality $f_g$, the pressure gradient decreases (Alvarez et al., 2001). A straight line provides a first-estimate fit through the data-points. This rough fit to the straight-line trend also implies that the water relative permeability remains fairly constant here. We can use this to determine the limiting water saturation $S_{w^*}$, which corresponds to the STARS model parameter $fmdry$. We can determine the water relative permeability in the high-quality regime by rewriting Darcy’s law as follows:

$$k_w(S_{w^*}) = \frac{u_w \mu_w (1 - f_g)}{k \nabla p}$$  \hspace{1cm} (2)

where $f_g$ and $\nabla p$ are obtained from the straight-line fit through the data (cf. (Boeije & Rossen, 2015a)). For this data-set the value of $k_w(S_{w^*})$ is $5.0 \cdot 10^{-2}$. To accurately determine the value of $fmdry$, suitable gas and water relative-permeability functions are needed. Mohamed and Nasr-El-Din (2012) provide relative-permeability...
curves for several carbonate rocks, including low-permeability Indiana Limestone. We fit a Brooks-Corey-type relative-permeability function to their data with the following result:

\[
k_{rw} = 0.28 \left( \frac{S_w - 0.32}{1 - 0.32 - 0.08} \right)^{2.44}
\]

\[
k_{rg} = 0.05 \left( \frac{1 - S_w - 0.08}{1 - 0.32 - 0.08} \right)^{1.02}
\]

The value of \( S_w^* \) (or \( f_{\text{mdry}} \)) can be determined by rewriting equation 3 into an expression for the saturation and using the aforementioned value of \( k_{rw}(S_w^*) \). The resulting value of \( f_{\text{mdry}} \) is 0.59.

Accurately determining the value of \( f_{\text{nmob}} \) requires knowing the foam quality at the point of transition between the low- and high-quality regime (\( f_g^* \) (cf. Boeije and Rossen (2015a)). In this experiment, all the data points are in the high-quality regime, so there is no transition point in the data. This means that the values of \( f_g^* \) and \( f_{\text{nmob}} \) cannot be determined. This is not a problem in this case since we are interested only in the mobility at the gas front in a SAG process. As discussed in Rossen and Boeije (2015) the mobility at the gas front is determined by the point of tangency to the fractional-flow curve, which for a foam flood is in the high-quality regime. To determine the point of tangency, first we need to determine the liquid saturations for all data points. We can then use the saturations to estimate the fractional-flow curve. The saturation can be determined in the same way we determined \( S_w^* \), using the Darcy’s law for the water phase along with the known relative-permeability function \( k_{rw} \).

The shock line is a straight line from the point of tangency to the initial condition (which in this case is at \( S_w = 1 \)). The resulting fractional-flow curve fit to the data, using the approach of Rossen and Boeije (2015), along with the shock line from the point of tangency, is shown in Figure 6a. A detailed view of the same fractional-flow curve around the point of tangency is shown Figure 6b, from which we can see that the water saturation at the point of tangency is 0.581.

![Figure 6](image)

Figure 6: (a) Water fractional flow for the experiment using the APG surfactant: points are experimental data (diamonds) where the liquid saturation is calculated using relative-permeability functions from data of Mohamed and Nasr-El-Din (2012). The resulting curve fit is obtained using the following STARS model parameter values: \( f_{\text{nmob}} = 20 \), \( f_{\text{mdry}} = 0.59 \), \( \varepsilon_{\text{dry}} = 600 \). Dashed line indicates shock from point of tangency (triangle) back to initial state. (b) detailed view of the same fractional-flow curve around the point of tangency.

We then use the water saturation to determine the total mobility at the leading edge of the foam bank using equation 5 and compare it to the mobility of the displaced phase (in this case water):

\[
\lambda_{\text{w}} \left( S_w' \right) = \lambda_{\text{w}} + \lambda_{\text{g}} = \frac{k_{rw} \left( S_w' \right)}{\mu_w} + \frac{k_{rg} \left( S_w' \right) \cdot \text{FM} \left( S_w' \right)}{\mu_g}
\]
where $S_w^t$ is the water saturation at the point of tangency to the fractional-flow curve. For the experiment using the APG surfactant, the resulting value of $\lambda_{rt}(S_w^t)$ is equal to $1.42 \times 10^3 \ [\text{Pa} \cdot \text{s}]^{-1}$. The experiment was performed at 55°C, at which the water viscosity is around 0.50 mPa·s (Kestin et al. (1978)). Therefore the mobility for water, which is the displaced fluid, is $1/\mu_w = 2.0 \times 10^3 \ [\text{Pa} \cdot \text{s}]^{-1}$. This means that the mobility of the leading edge of the foam bank is slightly lower than that of the water: $\lambda_{rt, \text{foam}}/\lambda_{rt, \text{water}} = 0.71$. Thus, the results indicate that, even though the foam is rather weak (i.e. has low apparent viscosity), it is still able to maintain mobility control over the displaced water phase. This result is similar to the ‘weak foam’ proposed by Boeije and Rossen (2015b). Foam like this has the advantage that the injectivity is relatively high, so a foam-injection process can be carried out in a shorter time. If used in the field, the resulting foam front may show some instabilities which could lead to lower sweep efficiency, as was shown in simulation studies by Boeije and Rossen (2015b) and Farajzadeh et al. (2016).

The fit through the pressure-gradient data of the APG experiment using the model parameters in Figure 6 is shown in Figure 7. Note that the transitional foam quality ($f^*_g$) is rather low for this set of parameters (around 0.25). This is a result of the chosen value of $f_{mmob}$. If a different value of $f_{mmob}$ were chosen, the transitional foam quality would shift left or right, but the straight-line fit through the data points in the high-quality regime would remain the same.

Figure 7: Steady state $f_g$ vs. $\nabla p$ data for the experiment using the APG surfactant along with STARS model fit using the following parameter values: $f_{mmob} = 20$, $f_{mdry} = 0.59$, $ep_{mdry} = 600$

We performed the same set of experiments using the same foam qualities with the AES surfactant. Figure 8 shows the pressure-gradient data as a function of foam quality. Apart from the lowest foam quality investigated
\( f_g = 0.5 \), all of the data points are in the high-quality regime. The straight line fit shown in the figure gives a first estimate of the behaviour in this regime.

The resulting water saturations are calculated using the same water relative-permeability function as before (i.e. equation 3) and are shown as fractional-flow data in Figure 9a. A more detailed close-up of the same fractional-flow data is shown in Figure 9b. Water fractional flow is not a monotonically increasing function of the water saturation. This result is similar to that of Wassmuth et al. (2001), Kibodeaux and Rossen (1997) and Xu and Rossen (2004). Rossen and Bruining (2007) argue that this kind of fractional-flow curve could represent a combination of two different curves, one for strong and one for weak foam. It is also possible that the large scatter in \( S_w \) values in Figure 6.9 may be the result of experimental uncertainty. However, to bring the datum of \( S_w = 0.62 \) (at \( f_g = 0.97 \)) down to \( S_w = 0.57 \), the measured pressure drop would have to be around 0.32 bar rather than the 0.21 bar that was measured. This is an increase of approximately 50\%, which is beyond the expected uncertainty in our pressure measurements.

According to Rossen and Bruining (2007), in such a case the shock goes to the weak-foam part of the fractional flow curve, directly below the lowest-\( S_w \) datum of the strong-foam part of the curve, as shown in Figure 10. The result is a high-mobility foam bank behind the shock, not capable of maintaining mobility control at the front. Our fitting approach is not suitable for such data-sets and therefore we cannot fit the results from this experiment.

![Figure 9: (a) Water fractional flow for the experiment using the AES surfactant: points are experimental data where the liquid saturation is calculated using relative-permeability functions from Mohamed and Nasr-El-Din (2012). The fractional flow does not increase monotonically with water saturation. (b) detailed view of the same experimental data, more clearly showing the non-monotonicity in the fractional-flow curve.](image)

![Figure 10: Construction of shock formed upon gas injection in SAG using the data from the experiment with AES surfactant and the method of Rossen and Bruining (2007).](image)

The Effect of Oil

To determine the effect of oil on the foaming performance of both surfactants we performed an experiment where we added crude oil (32°API) to the co-injection process. The foam quality in this experiment was kept constant at \( f_g = 0.5 \). Without oil we managed to get a fairly strong foam at this foam quality with both surfactants (cf. Figures 5 and 7) so if there is any effect of the oil we should be able to clearly see it from a lower pressure
gradient over the core. We maintained the same injection rates of gas and surfactant solution that we also used in
the previous experiment (i.e. 0.05 ml/min = 6.63 \cdot 10^{-7} \text{ m/s}) and then we added the crude oil to the injection
process. We do not include oil flow rate in the definition of foam quality. The oil flow rate used in these
experiment is one quarter of the gas flow rate and since we use a foam quality of 0.5 this is also equal to one
quarter of the surfactant solution flow rate \((Q_o = Q_g / 4 = Q_w / 4)\). We first generated a foam in the core without oil
and then started oil injection to see its effect on the pressure gradient. Figures 11 and 12 show the resulting
pressure-gradient profiles for the APG and the AES experiment respectively. Both experiments were carried out
at a salinity of 200,000 ppm. These plots comprise both the injection process without oil until a steady state is
reached and the subsequent process where oil is co-injected along with the other fluids. The pressure gradient
without oil is somewhat lower (3.0 vs. 3.9 and 4.8 MPa/m) than it was in the previous experiments even though
the experimental conditions were the same. This difference could be the result of a change in permeability of the
core. For each surfactant, the same core was used for the initial experiments without oil and this experiment with
oil. Prolonged periods of liquid injection may have led to dissolution thereby affecting the permeability. It is also
possible that some pollutants entered the system (e.g. impurities in the surfactant solution) which may have had a
modest detrimental effect on the foaming performance.

Figure 11: Pressure-gradient results from the experiment with the APG surfactant when oil is added to the injection
process. Steady state without foam is reached around \( t = 14,000 \text{ s} \), after which oil injection is started, which reaches the
core shortly thereafter.

Figure 12: Pressure-gradient results from the experiment with the AES surfactant when oil is added to the injection
process. Steady state without foam is reached around \( t = 5,000 \text{ s} \), after which oil injection is started, which reaches the
core shortly thereafter.

Further observations show that both surfactants perform similarly. The maximum pressure gradient without oil is
3.0 MPa/m for both surfactants. Then when oil is added there is a short rise in the pressure gradient, because the
oil has not yet reached the core, but the injection rate increases slightly. This is then followed by an abrupt
decline in pressure gradient, implying oil is destroying the foam almost completely. However, after this initial
decline, a gradual increase in pressure gradient occurs, meaning that some mobility reduction is again taking
place. With oil, the steady-state pressure gradient is roughly halved, compared to the injection process without
oil, for both surfactants (\( \nabla p = 1.6 \) and \( 1.5 \text{ vs. } 3.0 \text{ MPa/m} \)).
We examined the foam structure qualitatively at the outlet of the core. We use RADEL tubing for our fluid lines, which can be used at high pressure, but is still transparent, allowing "by eye" observations of the fluids inside. From our qualitative observations we find that the fluids at the outlet of the core were more like an emulsion than a foam. A representative example of the observations is shown in Figure 13, which shows the fluids inside the RADEL tubing. Overall we find that foaming in the presence of oil is challenging in these rocks, but that some mobility reduction can still be achieved.

![Figure 13: Fluids at the outlet of the core; appearing more like an emulsion than a foam.](image)

**CT Experiments**

CT scans depend on attenuation coefficients, which are different for every material. The original attenuation coefficients are commonly transformed into Hounsfield units (also known as CT numbers) using a linear transformation. Hence one can use these to distinguish the liquid (aqueous surfactant solution; no oil), gas and the rock itself. To calculate rock porosity and fluid saturations inside the core we make use of the method outlined in detail in Rangel-German et al. (1999). Here we only mention the parts of their method relevant to our study (i.e. limited to two-phase flow), but we use the same nomenclature as in their study. First we calculate the rock’s porosity by comparing attenuation coefficients of a dry core with one that is fully saturated with liquid:

\[
\phi = \frac{CT_{cw} - CT_{cd}}{CT_{cw} - CT_{a}}
\]  

where \(CT_{cw}\) is the CT number for the rock at a matrix location that is fully saturated with liquid. \(CT_{cd}\) is the CT number for a dry core at a matrix location. Both \(CT_{cw}\) and \(CT_{cd}\) may vary for every pixel in the CT image of the core. \(CT_{a}\) is the CT number for the liquid (water). Since we do not use any salinity in the liquid solution, \(CT_{a}\) is close to 0 for this study. \(CT_{n}\) is the CT number for air, which is about -1000. For two-phase, gas-liquid systems, the liquid saturation inside the core can be calculated as follows:

\[
S_w = \frac{CT_{sw} - CT_{cd}}{CT_{sw} - CT_{a}}
\]  

where \(CT_{sw}\) is the CT number for the core saturated with both liquid and gas. This is the CT number we obtain from the scans that are taken during the foam flooding experiments. Figure 14a shows a vertical scan through a core (17 cm in length and 4 cm in diameter) at vacuum. This scan clearly shows multiple distinct vugs that are several mm in size. These vugs appear as black spots on the scan, because of their lower attenuation coefficient. The resulting porosity map for the core is shown in Figure 14b and Figure 14c. The vugs have porosity equal to 1. Therefore, these dominate the porosity map and it is hard to distinguish details in the regions with lower porosity. Therefore, we plot the porosity figure again with the color axis clipped so every porosity value higher than 0.3 is shown as full scale.
We measured the both liquid permeability and porosity of this rock in similar fashion to the previous set of experiments; on average the permeability is around 0.5 mD and the porosity is 0.2. The foam floods in this experiment are all carried out using a very high foam quality $f_g = 0.99$ and total superficial velocity of $5.31 \cdot 10^{-7}$ m/s. This quality was chosen because the low permeability (0.5 mD) means that large pressure gradients are expected. Also, the absence of salts in the surfactant solution and carrying out the experiment at ambient temperature means that a fairly strong foam can be generated, leading to even higher pressure gradients. Since we aim only to visualize the influence of heterogeneities, this limitation of the foam quality is acceptable. Pressure-gradient results from this experiment are shown in Figure 15. We applied this relatively low superficial velocity in order to maintain a pressure drop over the core within the range of the transducer and also to avoid damaging the core. As can be seen from the graph, the maximum observed pressure gradient is roughly 18 MPa/m, which for a 17 cm core equates to a 30 bar pressure drop. This also means that there is some ambiguity in the terms Pore Volume (PV) and foam quality, because these are defined with respect to the 100 bar back pressure.

The figure also shows that, at least in terms of pressure gradient, a steady state is achieved in this core. There are still some observable pressure fluctuations after 2 PV of fluids has been injected, but these fluctuations are not significantly larger than those found in previous experiments using the more homogeneous Indiana Limestone cores. This implies that the rock heterogeneities, such as the vugs, do not cause large fluctuations in the pressure response on a macro-scale. The resulting steady-state pressure gradient is approximately 13.8 MPa/m. Under
these conditions, this is equivalent to a mobility reduction factor ($MRF$) of around 12.8 compared to the mobility of water. So even at very high foam quality it is still possible to generate fairly strong foam in this rock. The observed mobility reduction factor is also significantly higher than that in the previous set of experiments using the Indiana Limestone, cores even though those were performed in higher-permeability rock. This implies that the salinity and the elevated temperature used in that experiment have a significant impact on foaming performance. As mentioned, the experiment was limited to very high foam quality. An attempt was done to carry out measurements at lower foam quality ($f_g = 0.8$), but the generated foam was too strong resulting in excessive built-up of pressure. Even when we lowered the flow rate further to the minimum accessible value of the equipment, still the resulting pressure gradients exceeded the transducer range and thus this experiment was terminated.

**CT Scans During Experiment**

In total, five CT scans were made of the core during the foam-flooding experiment using $f_g = 0.99$. All of the scans were taken with the core at steady state in terms of pressure gradient. The first three CT scans were taken, as indicated in Figure 14, after 2, 2.25 and 2.5 PV of fluid injection respectively. After these three scans, we tried using a different foam quality, the aforementioned experiment with $f_g = 0.8$ which was terminated due to excessive pressure build-up. Afterwards the high-foam-quality of $f_g = 0.99$ was resumed and two more scans were taken once steady state had been reached once more. The latter two scans were taken to determine the reproducibility of the results.

By combining the CT results during the experiment with scans of a dry core and a core fully saturated with surfactant solution we can convert the resulting scans into liquid-saturation maps using Eq. 7. We analyze the results in three different ways. First we look at the average liquid saturation at every vertical position inside the core. That is, we take the arithmetic average of every horizontal line of saturation voxels and use the result as a measure of the variation in saturation throughout the core. Next, we look at the complete, non-averaged, data to see how the heterogeneities affect the saturations in the foam on the core scale. Finally, we focus on the saturations in the vugs and see how the fluids are distributed there and how this distribution changes with time.

The resulting average saturation profiles as a function of the height of the core from this experiment are shown in Figures 16a and b. The former shows just the saturation profile from the first scan (at 2 PV injected). This shows a relatively constant saturation of $S_w = 0.37$ throughout the core, although there are some minor fluctuations from this number. The only place where the liquid saturation changes drastically is towards the outlet of the core. This could be caused by the capillary end effect, which causes liquid hold-up towards the core’s outlet. The fluctuations in the saturation that are observed also do not coincide with the axial location of the vugs: thus at a core scale the vugs do not seem to have a great effect on the average saturation. Figure 16b shows the combined average saturation profile from all the CT scans. This shows there is only minor variation in the saturation in time, indicating that a steady state is reached and is not affected greatly by the heterogeneities in the rock.

![Figure 16: Average saturation taken at every vertical position throughout the core after the system reached steady state at: (a) t = 2 PV, (b) multiple points in time after state (a).](image-url)
Based on the average saturation profiles, we might conclude that the rock heterogeneities have no effect on the saturation inside the core, since the resulting average all appear to be constant throughout the core and also steady in time. However, a look at the complete non-averaged CT scans, as shown in Figures 17a through e, shows that on the core scale large variations in saturation are found. These figures show the five complete, non-averaged, saturation profiles. These are the profiles used as input data in the average profiles discussed above. Broad features are consistent among the scans, but there is also substantial fluctuation in saturation on the fine scale from image to image. This suggests either scatter in the CT images or fluctuation in saturation on the fine scale. The core needs to be analyzed in greater detail to fully explain these large fluctuations in saturation.

Finally we focus on the flow behaviour inside the vugs. As shown in the scan of the dry core (Figure 14a), there are three large vugs, several mm in size along the right-hand side of Figures 14a and b. Figures 18a to c show the saturation profiles in the vugs indicated on the leftmost figure. The second figure from the left shows a detailed view of the given vug. The second figure from the right shows the saturation profile in the vug from the first scan that is taken and the rightmost figure shows the result of the last scan. The resulting saturation profile remains constant through time for each of the vugs indicating that a steady state is reached. We find that two regions can be distinguished within each vug: one filled with liquid and the other filled with gas. The latter is located slightly above the liquid region. There is a clear interface separating the two regions which remains stable through time (that is, it is in the same location in the last scan as in the first). This means that foam segregates inside the vugs, but the effect of the vugs appears to be limited to the vugs itself. Directly outside the vugs we can already see changes in saturation (green colors in the saturation profiles) which implies foam is present there.

Conclusions
The long-core experiments show that foam EOR is possible in low-permeability carbonate rocks. However, foam strength is considerably reduced compared to previous experiments we conducted in high-permeability unconsolidated bead-packs. This finding agrees with previous studies (e.g. Khatib et al. (1988)) who found reduced foam strength in low-permeability rocks. The two different surfactants we tested here show similar foam strength, but the experiment with the AES surfactant shows indications of multiple steady states at very high foam qualities. As a result of this, our fitting approach to determine whether this surfactant is suitable for a SAG foam application cannot be applied accurately to that case, because we do not know where to place the shock on the fractional flow curve. For the experiment using the APG experiment we did not find these multiple steady states during the experiments so for this experiment we can use our fitting approach. We find that at the leading
edge of a foam front in a gas-injection cycle of a SAG process, this foam is just able to maintain mobility control over the displaced phase. It is comparable to the ‘weak foam’ formulation discussed in Boeije and Rossen (2015b). When crude oil is added to the injection process, the foam strength is reduced. In our experiments we found that the resulting steady-state pressure gradient is roughly halved in the presence of oil. Also we observed that the fluids at the outlet appear more like an emulsion than a foam. Still, some mobility reduction can be achieved with crude oil flowing along with the foam.

![Figure 18: Liquid saturation ($S_w$) inside the largest major vugs. Left figure of dry core indicates the vug of which the saturation profile is shown. Second figure from the left shows detailed view of the vug. Second figure from the right shows profile from the first scan that is taken. Rightmost figure shows the profile of the last scan that is taken. Profiles show separate regions for liquid (white color in the profile) and gas (black) in the vugs with gas being on top of the liquid. The interface between the two phases remains at roughly the same location throughout the experiment.](image)

CT visualization experiments in heterogeneous, vuggy rocks show that foaming is also possible in these rocks. Based on average saturation profiles we conclude that the influence of vugs is limited. Average saturation (on a scale of cm) is fairly uniform throughout the core, indicating that there is foam present. The detailed, non-averaged saturation profiles of the entire core show more-chaotic behaviour than the average values. At steady state, regions of high and low liquid saturation are present very close to each other. This might be the result of layering in the heterogeneous core. The core needs to be analysed in greater detail (e.g. through µCT imaging) to verify this conclusion. The influence of the vugs does not prevent the generation of foam on the core scale. We
find that there are two distinct regions inside the vugs. The bottom part is filled with liquid whereas the remainder is full of gas. This implies that foam segregation occurs in the vugs, but this effect is limited to the vugs. Foam still exists in the remainder of the core.

References


