Carbon Dioxide Transport and Sorption Behavior in Confined Coal Cores for Enhanced Coalbed Methane and CO$_2$ sequestration

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Abstract

Measurements of sorption isotherms and transport properties of CO$_2$ in coal cores are important for designing enhanced coalbed methane/CO$_2$ sequestration field projects. Sorption isotherms measured in the lab can provide the upper limit on the amount of CO$_2$ that might be sorbed in these projects.

Because sequestration sites will most likely be in unmineable coals, many of the coals will be deep and under considerable lithostatic and hydrostatic pressures. These lithostatic pressures may significantly reduce the sorption capacities and/or transport rates. Consequently, we have studied apparent sorption and diffusion in a coal core under confining pressure. A core from the important bituminous coal Pittsburgh #8 was kept under a constant, three-dimensional external stress; the sample was scanned by X-ray computer tomography (CT) before, then while it sorbed, CO$_2$. Increases in sample density due to sorption were calculated from the CT images. Moreover, density distributions for small volume elements inside the core were calculated and analyzed. Qualitatively, the computerized tomography showed that gas sorption advanced at different rates in different regions of the core, and that diffusion and sorption progressed slowly. The amounts of CO$_2$ sorbed were plotted vs. position (at fixed times) and vs. time (for various locations in the sample). The resulting sorption isotherms were compared to isotherms obtained from powdered coal from the same Pittsburgh #8 extended sample.

The results showed that for this single coal at specified times, the apparent sorption isotherms were dependent on position of the volume element in the core and the distance from the CO$_2$ source. Also, the calculated isotherms showed that less CO$_2$ was sorbed than by a powdered (and unconfined) sample of the coal. Changes in density distributions during the experiment were also observed. After desorption, the density distribution of calculated volume elements differed from the initial distribution, suggesting hysteresis and a possible rearrangement of coal structure due to CO$_2$ sorption.

1. INTRODUCTION

In the carbon dioxide enhanced coal bed methane production/sequestration process, CO$_2$ is injected into a coal seam to drive methane out of the bulk matrix. Carbon dioxide is sorbed by the coal$^1$; thus CO$_2$ injection will not only increase pressure gradients to drive methane flow, but it may also displace methane that would otherwise stay sorbed in the coal matrix, by decreasing the partial pressure of methane in the gas mixture. Because coal seams have proven to store
large quantities of sorbed gases for geologic time, they exhibit significant potential for sequestration of carbon dioxide for the indefinite future\(^2\).

There are two important parameters to consider when evaluating future ECBM/CO\(_2\) sequestration prospects: the amount of gas that the reservoir can store, and, the potential to transport large quantities throughout the reservoir.

Sorption isotherms relate the gas storage capacity of the coal as a function of pressure\(^3,4\). The information obtained from sorption isotherms can be used to predict the maximum possible volume of gas that can be stored in the coal during injection, or the amount of the gas that will be released from the coal during production.

Most of the sorption studies on coal have been done on coal powders\(^5,6\). These studies provide a relatively fast way of estimating the storage capacity of coal. However, for modeling purposes, one needs also an accurate time for the sorption equilibrium to be achieved when the sample is under the same confining pressure as in reservoir conditions\(^7\). Thus, the sorption isotherms and the rate of diffusion of CO\(_2\) in coal are important parameters for both technical and economic purposes\(^8,9\).

Only a few researchers have attempted the measurement of CO\(_2\) sorption under confining pressure. Karakan and Okandan\(^10\) observed gas transport and storage using coal samples mounted in X-ray transparent coreholders that enabled application of confining pressure during CO\(_2\) injection. The processed images from computer tomography were correlated with SEM (scanning electron microscopy) pictures at the same location, revealing different sorption rates for various lithotypes. Later, Karacan and Mitchell\(^11\) studied the behavior of different coal microlithotypes in a 2.5-cm-diameter Pittsburgh coal sample when exposed to CO\(_2\), while under a constant effective pressure of 1.36 MPa during gas uptake. The coal was exposed to CO\(_2\) at three different gas pressures: 1.7 MPa, 3.06 MPa, and 4.4 MPa (16.78, 30.2, and 43.42 atm). The CO\(_2\) storage process was observed using both qualitative and quantitative X-ray computerized tomography (CT) scanning. The core was examined petrographically after exposure to CO\(_2\) to identify those microlithotypes showing different sorption behavior. Results showed that different microlithotypes react differently when exposed to high pressure CO\(_2\), displaying different sorption and rate properties. Vitrinite was found to be a swelling maceral under CO\(_2\) sorption conditions. Clay and inertinite had the highest CO\(_2\) sorption rate, whereas the gas sorption rates of vitrinite and liptinite by comparison were low.

In another investigation using a confined high-volatile bituminous Pittsburgh seam coal, Karacan\(^12\) pressurized a 2.5-cm-diameter core that was 2.5 cm long with CO\(_2\) at 298 K (25 °C) at various pressures and various confining pressures. Changes in the sorbed gas concentration in the coal and matrix properties were simultaneously investigated by means of dual-energy X-ray computed tomography. Application of this technique allowed measurement of the amount of gas sorbed in various microlithotypes, observation of changes in the coal matrix, and estimation of the kinetics of the complex heterogeneous processes. Different parts of the coal reacted differently. Vitrinite, liptinite, and clarite swelled as a result of imbibition of CO\(_2\). Clay and inertinite were compressed. Vitrinite first swelled, then after reaching a maximum volume, expelled some CO\(_2\). This was called “breathing” and was thought to be the result of rearrangement of the macromolecular structure. The largest amounts of CO\(_2\) was stored in the inertinite and clay regions, which imbibed 0.08 g CO\(_2\) per gram of coal. Vitrinite adsorbed 0.06 g
CO₂ per gram of coal. Clay-rich sections were compressed to compensate for the swelling in vitrinite regions. In vitrinite, liptinite, and clarite regions of the coal the amount of gas sorption increased substantially, followed by a falloff where some of the sorbed gas was expelled due to rearrangement of the macromolecular network to a structure that has decreased gas solubility.

In a recent work\(^{13}\) amounts of sorption and diffusion rates were each measured for unconfined coal cores by means of an ambient-pressure gravimetric technique; the sorptions were compared with results obtained using computer tomography. The results showed that CT can provide not only qualitative measurements of spatial variations of CO₂ concentrations within the core, but quantitatively accurate measurements of CO₂ sorption.

In this work we present a unique sorption and desorption experiment under three-dimensional confinement of a cylindrical coal core exposed to CO₂ only at one side. In this way we studied the storage capacity of coal to CO₂, the rate of sorption through the coal at different pressure ranges, and the sorption kinetics of two regions with different lithotypes.

2. METHODOLOGY

Mineral samples, such as coals, are heterogeneous and contain a spectrum of lower and higher atomic number elements. When a cylindrical core is scanned, the attenuation of the X-rays is measured, and digital images are produced. These images map the numerical values of the attenuation coefficient within a trans-axial plane (i.e., "slice") of the investigated object. The attenuations depend on the electron densities, but usually these can be directly related by means of appropriate calibrations to the conventional weight densities. If only one energy level is used, the only quantitative information obtained refers to the local value of the linear attenuation coefficient, which may be insufficient for a detailed analysis\(^{14}\). Consequently, a dual energy scanning technique was used during this experiment. This method, known as dual energy X-ray absorptiometry (DXA), is widely used in the medical field as a means of measuring bone mineral density. Two X-ray beams with differing energy levels are aimed at the object to be studied. In the high-energy case the Compton scattering is predominant, while in the low case energy the photoelectric absorption prevails. These two effects have a known dependence on the atomic number and electron density\(^{15}\).

Most medical scanner units provide X-ray attenuation data in terms of CT numbers (CTN in Hounsfield units)\(^{14}\). The density of a scanned material is a linear function of the low energy \([\text{CT}_{\text{low}}]\) and high energy \([\text{CT}_{\text{high}}]\) CT numbers, as expressed by the following relationship\(^{16}\):

\[
\rho = A [\text{CT}_{\text{low}}] + B [\text{CT}_{\text{high}}] + C
\]  

The coefficients, A, B, and C can be calculated using the CT numbers for the three reference materials, by solving a system of equations with three unknowns. With these coefficients, one can calculate the density of the core under vacuum and the density of the coal after the CO₂ sorption process has begun. Having the values of density at each time and pressure step, and knowing the volume and porosity of the sample, one can calculate the amount of CO₂ sorption.
3. EXPERIMENTAL

A sample of Pittsburgh #8 coal was obtained from the Emerald Mine near Wayneburg, PA. A cylinder of diameter 25.0 mm was cored from the sample and scanned in the CT instrument for an initial examination of its density heterogeneities. This core was then cut trans-axially into two shorter cores: one for further CO₂ injection and CT scanning experiments and another for a manometric test of CO₂ sorption. The CT scans revealed no significant differences between the two cores. The core used in the manometric experiment was then powdered and its sorption of CO₂ was measured at 30 °C by means of a conventional manometric technique and literature values for the CO₂ density.5

Figure 1 presents a schematic of the experimental setup for the CT experiments. The core with length 25.5 mm was placed in the sleeve (Buna rubber) of a tri-axial aluminum coreholder. This coreholder allowed application of a radial confining pressure over the length of the core, and injection of CO₂ at either (or both) ends of the core. In this experiment CO₂ pressure was applied only at one end. This arrangement allowed observation of the kinetics of CO₂ sorption over the entire length of the core.

<table>
<thead>
<tr>
<th>Reference Sample</th>
<th>CT number at 130 kV</th>
<th>CT number at 80 kV</th>
</tr>
</thead>
<tbody>
<tr>
<td>Water</td>
<td>-95</td>
<td>11.5</td>
</tr>
<tr>
<td>Fused silica</td>
<td>1162</td>
<td>1755</td>
</tr>
<tr>
<td>Aluminum</td>
<td>1780</td>
<td>2725</td>
</tr>
</tbody>
</table>

The assembled core holder was mounted in the chucks of a modified medical CT scanner (Universal System, HG 350). The scanner is a fourth-generation medical scanner, with one rotating X-ray source and 1200 detectors, with a pixel resolution of 0.25 mm. The core was scanned in contiguous 2 mm thick slices; hence each measured CT number represented a voxel of 0.25 mm x 0.25 mm x 2.0 mm. Fifteen scans were taken along the axial length; these included the metallic end plugs as well as the core. However, for analysis only 11 slices were used, representing only coal. Two energy levels, 130kV and 80 kV at 100 mA were used. The scanner was calibrated with reference materials of known density: aluminum (Al), fused Silica (SiO₂), and water (H₂O). These were placed inside the coreholder.

The confining pressure was supplied from a nitrogen bottle via a pressure regulator. The pressure was kept constant without problems during various pressure steps of the experiment. That was possible due to the very large nitrogen bottle volume compared to the inner volume between the sleeve and the exterior aluminum cylinder.

The CO₂ was delivered to the core from a CO₂ bottle. Due to the small volume of the core, the CO₂ volume in the cylinder was comparatively extremely large; hence a constant CO₂ pressure was maintained at the injection port, even if there were any slow leaks in the system.

The evacuated core was initially scanned at the two energy levels, 130 kV and 80 kV, at a confining pressure of 200 psig (1.36 Mpa). In all of the following steps, the pressure differential...
between confining pressure and inlet pressure was kept constant at 200 psig. The first CO\textsubscript{2} sorption step was performed at a CO\textsubscript{2} injection pressure of 195 psig (hence a confining pressure of 395 psig). To measure the CO\textsubscript{2} uptake, the core was scanned repeatedly at predetermined, frequent intervals and at each energy level. The intervals between scans were increased as the elapsed time for sorption became longer, and the scanning for the first pressure step was stopped when the change in X-ray sorption appeared to be negligible. After injection for 165 hours at 200 psig, the process was repeated at CO\textsubscript{2} injection pressures of 325 psig, 450 psig, and 600 psig, with corresponding increases in the confining pressure. The injection was continued for 392 hours at 325 psig, for 282 hours at 450 psig, and for 226 hours at 600 psig, for a total of 1065 hours of injection.

After the injection at 600 psig had been stopped, the CO\textsubscript{2} was desorbed from the core at 400 psig (for 180 hours), 200 psig (for 144 hours), and atmospheric pressure (for 278 hours), respectively, for a total experimental time (sorption plus desorption) of 1667 hours; during desorption the same 200 psig pressure differential between the confining pressure and the CO\textsubscript{2} outlet pressure was maintained. The temperature in the laboratory was kept approximately constant at 21 °C, during all phases of the CT experiment.

The average CT number for each slice at each energy level was calculated for use in equation (1). The average (bulk) density of each slice at each time was calculated from its average CT number, equation (1), and the values of A, B, and C obtained from the calibration measurements. From each of these slice densities the average density of the corresponding slice under vacuum was subtracted to give the density increase of the slice due to its CO\textsubscript{2} uptake. These slice-average CO\textsubscript{2} densities (in units of mass CO\textsubscript{2} per unit volume of slice) were converted to slice-average CO\textsubscript{2} concentrations in units of mass CO\textsubscript{2} per unit volume of slice by means of the average density of each slice as obtained from the CT measurements on the evacuated core. As a check on the accuracy of the densities determined from the CT data, the average (mass) density of the core was calculated from the CT data and compared to the sample density as obtained from its weight and volume.

Color images were generated that illustrated the spatial distribution of density in each slice either before or after sorption had occurred. Images also were generated that illustrated the spatial distribution of CO\textsubscript{2} within the sample after a chosen injection time. The analyses were performed with VoxelCalc\textsuperscript{17} and Image J\textsuperscript{18}, software processing packages.

After the end of the CO\textsubscript{2} sorption/desorption experiment the coal core was recovered, set in epoxy, and polished. Using the CT images obtained, the orientation of the core was determined, and a petrographic examination of the macerals and minerals in the one of the polished surfaces of the coal core was performed.

4. RESULTS and DISCUSSION

CT Sorption

From the calibration measurements and equation (1) we obtained the following set of three equations for water, silica, and aluminum, respectively:
1.00 g/cm$^3$ = A [11.5 CTN] + B [-95 CTN] + C \hspace{1cm} (2a)

2.20 g/cm$^3$ = A [1755 CTN] + B [1162 CTN] + C \hspace{1cm} (2b)

2.699 g/cm$^3$ = A [2725 CTN] + B [1780 CTN] + C \hspace{1cm} (2c)

Equations (2) gave the following values of the parameters A, B, and C for calculation of (bulk) mass densities from equation (1) and the CT numbers measured at 80 kV and 130 kV: A= 2.07 g/CTN cm$^3$, B=-0.806 g/CTN cm$^3$, and C=1206.23 g/cm$^3$.

Figure 2 illustrates the density variations in each trans-axial cross section of the coal sample. The scans were obtained with the core under vacuum. The light blue color represents the lower density coal matrix, while the colors toward red represent higher density intrusions in the coal. It can be seen that the middle-lower left area of each slice shows a higher density than other regions. The average bulk density of the sample calculated from the CT data was $\rho = 1.20$ g/cm$^3$, which was within 1.5% of the density ($\rho = 1.22$ g/cm$^3$) calculated from the mass and volume measurements.

Figure 3a illustrates the spatial distribution of sorbed CO$_2$ after 165 hours of injection at 195 psig. It can be seen that slices further away from the injection port contained less CO$_2$. Also, one can observe that the slices did not sorb CO$_2$ uniformly, and that the lower-left portion of the each slice sorbed very little CO$_2$.

Figure 3b is a plot of the average CO$_2$ concentrations in various slices vs. time of injection at 195 psig, up to 165 hours of injection. (Slice 1 is at the injection port; slice 2 was contiguous to it; and each slice was 2 mm thick.) It is clear that the diffusion-sorption process had not reached equilibrium over a 2 cm distance even after 165 hours.

Figures 4a, 4b, and 4c show CO$_2$ sorption images for each of the slices at injection pressures 325 psig, 450 psig, and 600 psig, respectively. Careful visual observation of the images confirms that slices close to the injection point had sorbed more CO$_2$ than slices not as close to it, even after the end of the fourth pressure step after a total CO$_2$ exposure time (for all injection pressures) of 1065 hours (6 weeks).

The average sorption for each of the (contiguous) slices near the end of the injection period for each CO$_2$ pressure is plotted in Figure 5. The sorption seems to decrease almost monotonically from the CO$_2$ inlet toward the opposite end of the core.

Figure 6 is a plot of the “apparent sorption isotherms” for various slices as obtained from the CT data. For slice 1, with the shortest diffusion length, the CO$_2$ sorption at the end of step 3 (400 psig, 839 hours total injection time) and that at the end of step 4 (600 psig, 1005 hours) are very nearly the same, indicating that for this slice sorption equilibrium may have been achieved. Sorption equilibrium in the other slices was not obtained, as indicated by the monotonic decrease of CO$_2$ concentration with distance from the injection inlet.

From Figures 3-6 one can see that the sorption of CO$_2$ was non-uniform in each slice, and also along the length of the core. It also was observed that the sorption seemed to be more pronounced in the upper right regions of the slices and less in the lower left regions.
Consequently, two rhomboidal “regions of interest” (ROI’s) of approximately the same size were defined, one in each of these regions. Figure 7a illustrates the locations of these regions. The density before sorption was calculated for each of these two regions in each slice of the sample. In Figure 7b these densities, along with the average density for the entire slice, are plotted vs. length along the core from the injection end. As expected, the coal densities in the ROI’s are different from the average slice density. In region 1 the density is higher than the average density of the core; however, its trend along the length of the core tends to follow that of the average density. The density in region 2 is always lower than the density in region 1; however, the variations are much larger, and the density is sometimes larger, sometimes smaller, than the curve for the average core density.

Figure 7c is a plot of sorption vs. length along the core for region 1, region 2, and the entire slice, each measured after 1065 total hours of injection. These curves illustrate the inhomogeneity of the sorption in the core. The region 1 sorption curve is much lower than the average sorption, confirming the visual observations from the pictures, while the region 2 sorption is much higher than that of region 1, and generally higher than the average sorption. It can be also observed that, in general, the higher the coal density, the lower the sorption. (See, also, Figure 11.) For the region 1 the calculated sorption seems to dip to negative values (slices 4 and 11). It is unclear whether the negative values are erroneous, or due to changes in density in these regions, from swelling or other effects.

In the last part of the experiment, we gradually desorbed the CO$_2$ in three steps: a) the 600 CO$_2$ inlet pressure was reduced to 400 psig, with a corresponding decrease of confining pressure (from 800 psig to 600 psig); b) the CO$_2$ inlet pressure was reduced from 400 psig to 200 psig (confining pressure 400 psig); and c) the inlet pressure was reduced to atmospheric pressure (200 psig confining pressure). As shown above, during the injection phase the CO$_2$ concentration exhibited the expected decrease from inlet to far end of the core. And, as plotted in Figure 8, near the end of the last stage of the desorption the concentration exhibited the expected decrease from far end of the core to the outlet. The behavior indicated by the upper two curves in Figure 8 is not so intuitive, and their interpretation has not yet been completed.

**Density histograms**

Figures 2-7 illustrate the heterogeneities in the coal sample and its sorption properties. In principle, the CT instrument may be able to detect heterogeneities on a scale as small as that of individual voxels. There were *circa* 9900 voxels in each slice, each voxel representing a parallelepiped of (measured) dimensions 0.21 mm x 0.21 mm x 2.0 mm.

Figure 9 represents the histogram for the density of slice 1 of the evacuated core (before gas injection). The average density for the slice is 1197.9 mg/cm$^3$. The average skewness factor (measure of the asymmetry) of the distribution is +0.67. (For a perfectly symmetrical distribution the skewness factor is 0.)

The density distribution in Figure 10 represents the voxel densities in slice 1 after 1065 total hours of exposure to CO$_2$, at the end of 600 psig sorption cycle. It shows that the average voxel density increased to 1231 mg/cm$^3$, and skewness to 0.88. Figure 11 illustrates the relationship between the density difference caused by sorption (density after sorption minus initial density) and initial coal density. The results suggest that lower density macerals sorbed more CO$_2$ than
did higher density macerals. This conclusion is consistent with a comparison of the density and concentration plots in Figures 7b and 7c for regions 1 and 2.

Figure 12 shows the density distribution in slice 1 at elapsed time t=1667 hours (10 weeks), after the core had desorbed CO₂ (in three stages) for 602 hours. The average density in the slice was 1198.9 mg/cm³, vs. 1197.9 mg/cm³ initially. This very small difference (0.08%) seems to indicate that, at most, a very small amount of CO₂ remained in the core. However, while the average density of the slice after degasification returned to almost its pre-injection value, the frequency distribution of the voxel densities in the slice did not. (The calculated average skewness of the distribution was 0.82 after degasification, 0.67 before injection of CO₂; comparison of Figures 9 and 12 reveals obvious changes in the frequency distribution.) The latter change suggests a structural re-arrangement in the slice may have occurred due to CO₂ sorption and/or the confining pressure.

**Petrographic observations**

As it was mentioned earlier, the lower left “corner” (see rhomboid) of the core did not sorb CO₂ as well as most of the other regions of the core. The density calculations prior to CO₂ injection showed relatively higher density constituents in broken bands in the core. Some of these bands were in the low sorption region. These bands were interpreted prior to petrographic examination to indicate the direction of the bedding planes in this sample. In actuality these bands are composed of higher density inertinite macerals (fusinite, semifusinite, inertodetrinite) that were often associated with clay filling of cell lumens and do indeed indicate bedding plane direction. These bands are fairly consistent and appear in other areas of the core.

Specifically, in the low sorption region, it was noted under petrographic examination that this area was composed mainly of trimaceral material (vitrinite, liptinite, inertinite), as shown in Figure 13. One observation of this region of the core was that the macerals were tightly intermixed and that liptinite (sporinite) and inertinite (fusinite, inertodetrinite) dominated the region as consistent bands. Moving further along into the interior of the core it was noticed, as seen in Figure 14, that there were consistent bands of vitrinite that alternated with mixed bands of intertinite and liptinite. These mixed bands were different than the trimaceral bands in the low sorption region in that there was not a high degree of tight intermixture as seen in the low sorption region.

**Isotherm Comparison: Confined Coal vs. Powder**

Figure 15 compares the sorption in slice 1 of the confined coal core as measured by CT at 21°C with the sorption obtained from the manometric experiment with the unconfined powder at 30°C. Because the sorption should be slightly smaller at the higher temperature, the difference between the two isotherms is actually somewhat larger than the Figure implies. It is impossible from these exploratory experiments to separate the effects of experimental method (CT vs. manometry), sample size (powder vs. core), confining pressure (none vs. several hundred psig), and degree of attainment of equilibrium on the difference between the two isotherms. However, it remains clear that many more measurements on confined and unconfined cores are needed for the attainment of physical property data needed for enhanced coal bed methane/sequestration projects.
5. CONCLUSIONS

Noninvasive, dual energy X-ray computed tomography provides an accurate way to calculate density variations and CO$_2$ concentrations, sorptions, and transport rates in coal cores at reservoir confining pressures. Average slice density increased during the CO$_2$ intake periods. Also, the slice-average CO$_2$ concentrations increased with time and with increasing CO$_2$ injection pressure. The CO$_2$ concentrations were higher for slices close to the injection point, and less as the distance from intake increased. This indicates that diffusion kinetics can play a very important role in a sequestration process, even for small cleat spacing. Desorption experiments further illustrated the importance of sorption kinetics for the sequestration process.

When small regions of interest inside the core were chosen for analysis, very high density variations were observed compared to the sample average. The calculated CO$_2$ sorption for a region with higher density showed markedly lower sorption values than the slice averages, while the region with lower density showed a higher sorption capacity. This finding was also confirmed by analysis of data for individual voxels (0.21 mm x 0.21 mm x 2.0 mm). Changes in the density histograms from before CO$_2$ injection to after virtually complete desorption suggest an internal structural rearrangement in the coal core.

Petrographic work implied that the regions of lower sorption were the higher- density inertinite macerals (fusinite, semifusinite, inertodetrinite), which were often associated with clay filling of cell lumens.

CO$_2$ sorption seems to be reduced by application of a confining pressure. However, further studies need to be performed, especially on confined coals, to address differences in experimental methods, and possible changes in coal structure due to confinement.

6. ACKNOWLEDGEMENTS

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7. REFERENCES


Figure 1. Experimental set-up.

Figure 2. Density variations in the scanned core.
Figure 3a. Changes of sample density caused by CO2 injection at the end of the first injection pressure step (195 psig).

Figure 3b. CO2 concentrations in various slices vs. time of injection at 195 psig.
Figure 4a. Changes of sample density caused by CO2 injection at the end of the second injection pressure step (325 psig).

Figure 4b. Changes of sample density caused by CO2 injection at the end of the third injection pressure step (450 psig).
Figure 4c. Changes of sample density caused by CO2 injection at the end of the fourth injection pressure step (600 psig).

Figure 5. CO2 concentrations vs. diffusion distance.
Figure 6. Apparent sorption isotherms for various slices.

Figure 7a. Locations of rhomboid regions of interest (ROI) 1 and 2.
Figure 7b. Density vs. core length for the complete slices and for the two regions of interest.

Figure 7c. CO2 concentrations vs. core length for the average slice and for the two regions of interest.
Figure 8. CO2 concentrations during the desorption experiment vs. distance from the gas outlet.

Figure 9. Histogram for the voxel densities of slice 1 of the coal sample before injection of CO2.
Figure 10. Histogram for the voxel densities of slice 1 after CO$_2$ sorption at 600 psig (total elapsed sorption time 1065 hours).

Figure 11. Increase of voxel density due to injection vs. voxel density before injection.
Figure 12. Histogram for the voxel densities after the core was degassed.

Figure 13. Trimaceral region in the low sorption region. The dark wavy material is liptinite (sporinite). The light material is intertinite. The gray material is vitrinite.
Figure 14. Inertinite (fusinite) band surrounded by vitrinite bands.

Figure 15. Comparison of sorption isotherms for the coal in powdered (unconfined) form and slice 1 of the confined core.