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Abstract

Responding to the need to reduce atmospheric emissions of carbon dioxide, Donald Brown (2000) proposed a novel enhanced geothermal systems (EGS) concept that would use CO₂ instead of water as heat transmission fluid, and would achieve geologic sequestration of CO₂ as an ancillary benefit. Following up on his suggestion, we have evaluated thermophysical properties and performed numerical simulations to explore the fluid dynamics and heat transfer issues in an engineered geothermal reservoir that would be operated with CO₂. We find that CO₂ is superior to water in its ability to mine heat from hot fractured rock. CO₂ also has certain advantages with respect to wellbore hydraulics, where larger compressibility and expansivity as compared to water would increase buoyancy forces and would reduce the parasitic power consumption of the fluid circulation system. While the thermal and hydraulic aspects of a CO₂-EGS system look promising, major uncertainties remain with regard to chemical interactions between fluids and rocks. An EGS system running on CO₂ has sufficiently attractive features to warrant further investigation.
INTRODUCTION

The resource base for geothermal energy is enormous, but commercial production of geothermal energy is currently limited to hydrothermal systems, in which naturally present fracture networks permit fluid circulation, and allow geothermal heat to be produced by tapping these hot fluids through wellbores. Most geothermal resources occur in rocks that lack fracture permeability and fluid circulation. The “enhanced” or “engineered” geothermal systems concept (EGS) aims to extract geothermal energy from these resources by (1) creating permeability through hydraulic stimulation or fracturing, which involves fluid injection through deep boreholes to activate existing rock fractures or create new ones, and (2) setting up and maintaining fluid circulation through these fracture networks by means of a system of injection and production boreholes, so that the thermal energy can be transmitted to the land surface for human use.

Previous attempts to develop EGS in the U.S., Japan, Europe and Australia have all employed water as a heat transmission fluid. Water has many properties that make it a favorable medium for this purpose, but it also has serious drawbacks. An unfavorable property of water is that it is a powerful solvent for many rock minerals, especially at elevated temperatures. Injecting water into hot rock fractures causes strong dissolution and precipitation effects that change fracture permeability and make it very difficult to operate an EGS reservoir in a stable manner (Xu and Pruess, 2004). Also, water is a sparse and valuable commodity in many areas, as e.g. the western U.S., and inevitable water losses during fluid circulation can be a severe economic liability.

Responding to the need to reduce atmospheric emissions of carbon dioxide, Donald Brown (2000) proposed a novel EGS concept that would use CO$_2$ instead of water as heat transmission fluid, and would achieve geologic storage of CO$_2$ as an ancillary benefit. Brown noted that CO$_2$ has certain physical and chemical properties that would be favorable for operation of an EGS system. Favorable properties of CO$_2$ emphasized by Brown include the following:

- large expansivity would generate large density differences between the cold CO$_2$ in the injection well and the hot CO$_2$ in the production well, and would provide buoyancy force that would reduce the power consumption of the fluid circulation system;
- lower viscosity would yield larger flow velocities for a given pressure gradient; and
• CO₂ would be much less effective as a solvent for rock minerals, which would reduce or eliminate scaling problems, such as silica dissolution and precipitation in water-based systems.

Brown also noted the lower mass heat capacity of CO₂ as an unfavorable property, but pointed out that this would be partially compensated by the greater flow capacity of CO₂ due to lower viscosity. Fouillac et al. (2004) suggested that an EGS using CO₂ as heat transmission fluid could have favorable geochemical properties, as CO₂ uptake and sequestration by rock minerals would be quite rapid. A preliminary evaluation of CO₂ as a working fluid for EGS was presented by Pruess and Azaroual (2006). Their findings suggest that CO₂ is roughly comparable to water as a heat transmission fluid, while offering distinct advantages for wellbore hydraulics.

The present paper compares thermophysical properties of CO₂ and water, and examines pressure and temperature conditions for flow of CO₂ in wellbores as well as in reservoirs with predominant fracture permeability. Comparisons are made with the flow behavior of water, in order to identify favorable as well as unfavorable characteristics of CO₂ as an EGS working fluid. We also present preliminary considerations on chemical aspects of a CO₂-EGS system.

**THERMOPHYSICAL PROPERTIES**

Fig. 1 shows the phase diagram for CO₂ in the range of temperature and pressure conditions that are of interest for injection into and production from enhanced geothermal systems. The critical point of CO₂ is at $T_{\text{crit}} = 31.04 \degree \text{C}$, $P_{\text{crit}} = 73.82 \text{ bar}$ (Vargaftik, 1975). At lower (subcritical) temperatures and/or pressures, CO₂ can exist in two different phases, a liquid and a gaseous state, as well as two-phase mixtures of these states (Fig. 1). Supercritical CO₂ forms a phase that is distinct from the aqueous phase and can change continuously into either gaseous or liquid CO₂ with no phase boundaries.
Fluid mass flow rates for a given driving force are proportional to the ratio of density to viscosity, \( m = \frac{\rho}{\mu} \). The sensible heat carried by mass flow is proportional to the specific enthalpy of the fluid. Additional parameters that are important for mass flow and heat transfer behavior include compressibility \( c = \left(1/\rho\right)\left(\partial\rho/\partial P\right) \) and thermal expansivity \( \varepsilon = -(1/\rho)\left(\partial\rho/\partial T\right) \).

Figs. 2 and 3 present thermophysical properties of CO\(_2\) and water. CO\(_2\) properties were calculated from the correlations of Altunin (Altunin, 1975; Pruess and García, 2002). We began using Altunin's correlations in 1999 when a computer program implementing them was conveniently made available to us by Victor Malkovsky of the Institute of Geology of Ore Deposits, Petrography, Mineralogy and Geochemistry (IGEM) of the Russian Academy of Sciences, Moscow. Altunin's correlations were subsequently extensively cross-checked against experimental data and alternative PVT formulations, such as Span and Wagner (1996), and were found to be very accurate (García, 2003). Water properties were obtained from the steam table equations as given by the International Formulation Committee (IFC, 1967).

The ratio of density to viscosity is generally larger for CO\(_2\) than for water, and dependence on temperature and pressure conditions is very different for the two fluids (Fig. 2). For water this ratio is mostly a function of temperature, with only weak dependence on pressure, reflecting the
primary dependence of both water density and viscosity on temperature. For CO$_2$, density and viscosity have significant dependence on both temperature and pressure. The variations are such that $(\rho/\mu)$ attains maximum values in a region that is emanating from the CO$_2$ saturation line, becoming smaller for liquid-like CO$_2$ (low T, high P) and for gas-like CO$_2$ (high T, low P). For (T, P)-conditions relevant for fluid injection, T \leq 50 ^\circ C, (\rho/\mu) for CO$_2$ is larger than for water by factors of 4-10. For temperatures near 200 ^\circ C, (\rho/\mu) for CO$_2$ is larger than for water by approximately a factor 2 at high pressures, while at pressures below 150 bar, water has the larger (\rho/\mu).

Fig. 3 compares the specific enthalpies for CO$_2$ and water. In both cases the reference state (zero enthalpy) was chosen as (T, P) = (20 ^\circ C, 100 bar). At high pressures near 500 bar, the increase of specific enthalpy with temperature for CO$_2$ is less than half of the increase for water, indicating that more than twice the CO$_2$ mass flow rate would be needed to achieve the same rate of sensible heat transport. Specific enthalpy of liquid water depends primarily on temperature, with only a weak pressure dependence. For CO$_2$ the pressure dependence is weak for liquid-like conditions, but becomes increasingly strong at lower pressures and higher temperatures. For adiabatic (thermally insulated) decompression, thermodynamic conditions will move along isenthalps (lines of constant specific enthalpy). Accordingly, decompression of hot, high-pressure CO$_2$ will be accompanied by substantial temperature decline, while for liquid water there would be a small temperature increase.

Table 1 shows that CO$_2$ is substantially more compressible than water, and has larger expansivity as well, especially at lower temperatures. Fluid densities will therefore vary much more strongly for CO$_2$ than for water as functions of pressure and temperature changes.
Figure 2. Ratio of fluid density to viscosity in units of $10^6$ sm$^{-2}$ for CO$_2$ (left) and water (right).

Figure 3. Specific enthalpy of CO$_2$ (left) and water (right) in units of kJ/kg, as function of temperature and pressure.
Table 1. Density, compressibility, and expansivity of CO₂ and water at selected (T, P)-conditions.

<table>
<thead>
<tr>
<th>T (°C)</th>
<th>P (bar)</th>
<th>CO₂</th>
<th>compressibility (1/Pa)</th>
<th>expansivity (1/°C)</th>
<th>water</th>
<th>compressibility (1/Pa)</th>
<th>expansivity (1/°C)</th>
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</thead>
<tbody>
<tr>
<td>20</td>
<td>100</td>
<td>856.251</td>
<td>1.490e-8</td>
<td>8.607e-3</td>
<td>1001.76</td>
<td>3.489e-10</td>
<td>1.944e-4</td>
</tr>
<tr>
<td></td>
<td>500</td>
<td>1048.77</td>
<td>2.484e-9</td>
<td>2.696e-3</td>
<td>1015.94</td>
<td>3.538e-10</td>
<td>1.448e-4</td>
</tr>
<tr>
<td>200</td>
<td>100</td>
<td>122.184</td>
<td>1.076e-7</td>
<td>3.036e-3</td>
<td>870.798</td>
<td>8.377e-10</td>
<td>1.321e-3</td>
</tr>
<tr>
<td></td>
<td>500</td>
<td>581.322</td>
<td>1.274e-8</td>
<td>3.172e-3</td>
<td>900.990</td>
<td>8.668e-10</td>
<td>1.077e-3</td>
</tr>
</tbody>
</table>

**WELLBORE FLOW**

The pressure gradient in a flowing well can be represented by a superposition of gravity, frictional, and acceleration terms (Brill and Mukherjee, 1999),

\[
\nabla p = (\nabla p)_{\text{grav}} + (\nabla p)_{\text{fric}} + (\nabla p)_{\text{acc}}
\]

For most applications of interest, the gravitational contribution to the pressure gradient is by far the dominant term, with frictional and inertial pressure gradients contributing typically a few percent or less. In a first effort to evaluate the pressures in EGS injection and production wells, we consider only the dominant gravitational gradient.

In an injection well, temperatures will increase with depth, primarily because of heat transfer from the surrounding rocks (Ramey, 1962). Additional temperature changes are expected as a consequence of fluid compression due to pressure increase. The latter effect is very small for water, but could be more significant for the highly compressible CO₂. Similarly, in a production well the temperature of a fluid parcel flowing upward will decrease due to heat loss to the surrounding formations. Additional temperature change will occur from decompression, which is expected to be a small effect for water, but could be significant for CO₂.
The temperature effects from heat exchange with the surroundings are of a transient nature and will diminish over time. In order to obtain a basic comparison between the pressure behavior of water and CO$_2$ wells, we start from the simplest possible approximation to temperature conditions: we consider an injection well that is entirely at the injection (wellhead) temperature $T_{\text{inj}}$, and a production well that is entirely at the production (downhole) temperature $T_{\text{pro}}$. In order to calculate the static pressure profile in a well, the well depth $z$ is divided into $N$ increments, $\Delta z = z/N$. The calculation proceeds recursively from level $n\Delta z$ to $(n+1)\Delta z$ by assigning $P_{n+1} = P_n + \rho_n g \Delta z$, where $\rho_n = \rho(T_n, P_n)$, and $g = 9.81$ m$^2$/s is gravitational acceleration. For definiteness, we adopt parameters applicable to the European EGS experiment at Soultz, and take a well depth of 5,000 m, with a downhole temperature of $T_{\text{pro}} = 200$ °C (Baria et al., 2005; Dezayes et al., 2005). Injection temperature is set at $T_{\text{inj}} = 20$ °C.

For both water and CO$_2$ we start from an injection wellhead pressure of 57.4 bar, slightly in excess of the CO$_2$ saturation pressure at injection temperature ($P_{\text{sat,CO}_2} = 57.36$ bar at $T_{\text{inj}} = 20$ °C). Corresponding static downhole pressures at 5,000 m depth are 528.7 bar for CO$_2$ and 553.4 bar for water (Fig. 4). Using these downhole pressures as starting values, we then obtain static pressures in the production well by integrating upwards at $T = 200$ °C. This results in production wellhead pressures of 288.1 bar for CO$_2$ and 118.6 bar for water. The difference in wellhead pressures between production and injection wells is 230.7 bar for CO$_2$ and 61.2 bar for water, indicating that a CO$_2$ circulation system would have far stronger buoyant drive, and would require less power to operate.

A more realistic outlook on longer-term P,T-conditions in flowing injection and production wells can be obtained by approximating fluid flow in the wellbore as isenthalpic. This approximation is often referred to as "adiabatic;" it ignores heat transfer between the wellbore fluid and the surroundings, which is appropriate for longer-term flow behavior. The isenthalpic flow approximation accounts for temperature changes that arise from (de-)compression of fluids, the so-called Joule-Thomson effect (Katz and Lee, 1990).
Figure 4. Static pressure profiles in CO$_2$ and water wells for constant temperatures of 20 and 200 °C, respectively.

We have calculated pressure and temperature conditions in static, isenthalpic columns of CO$_2$, using a similar recursion as for the constant-temperature wells considered above, except that now we need to consider temperature variation with depth as well, based on constant specific enthalpy. For an injection well, we perform a "top down" calculation starting from wellhead conditions of (T$_0$, P$_0$), corresponding to a specific enthalpy of h$_0$ = h(T$_0$, P$_0$). At depth level n we have conditions of (T$_n$, P$_n$), from which we obtain $\rho_n = \rho(T_n, P_n)$ and $P_{n+1} = P_n + \rho_n g \Delta z$, just as before. The temperature at level n+1 is obtained as $T_{n+1} = T(P_{n+1}, h_0)$; the required inversion of the $h = h(T, P)$ relationship is accomplished by Newtonian iteration, using $T_n$ as a starting guess. Fig. 5 shows (T, P)-profiles in a 5000 m deep injection well for several different wellhead temperatures and pressures. Fig. 6 shows (T, P)-profiles in a 5000 m deep production well for different downhole conditions.

Fig. 5 indicates that temperatures will tend to increase as CO$_2$ is flowing down the injection well and attaining increasing pressures. The difference between downhole and wellhead temperatures...
is larger for smaller wellhead pressures, and increases strongly when wellhead temperatures are increased. For the lower wellhead temperature cases shown in Fig. 5, temperature changes versus depth are non-monotonic, with significant temperature decline at greater depth, especially when wellhead temperature is low and/or wellhead pressure is large.

![Temperature-pressure conditions for isenthalpic flow of CO₂ in a 5000 m deep injection well, for different wellhead temperatures and pressures.](image)

Figure 5. Temperature-pressure conditions for isenthalpic flow of CO₂ in a 5000 m deep injection well, for different wellhead temperatures and pressures.

These features can be understood from the dependence of specific enthalpy of CO₂ on temperature and pressure, see Fig. 3. For adiabatic (de-)compression processes, thermodynamic conditions will move along lines of constant enthalpy, indicating that compression starting from modest pressures and/or elevated temperatures will be accompanied by strong temperature increases. At temperatures below 50 °C, however, the isenthalps shown in Fig. 3 curve slightly backwards towards lower temperatures at high pressures, indicating that in this region isenthalpic compression will be accompanied by a temperature decline.
Figure 6. Temperature-pressure conditions for isenthalpic flow of CO₂ in a 5000 m deep production well, for different downhole temperatures and pressures.

Increased downhole temperatures in the injection well are favorable from the viewpoint of reservoir heat extraction, but they also reduce the pressure increase with depth in the injection well. This will reduce the buoyant pressure drive available for pushing CO₂ through the EGS reservoir, and will increase the power requirements for maintaining fluid circulation.

Analogous considerations apply to temperature and pressure behavior in production wells (Fig. 6). Here the isenthalpic decompression will cause CO₂ temperatures to decline as it flows up the well. The temperature drop along the well becomes stronger for smaller downhole pressures; at $T_{WB} = 200$ °C the temperature changes range from $\Delta T = -22.6$ °C for $P_{wb} = 550$ bar to -25.7 °C at $P_{wb} = 500$ bar and -28.7 °C at $P_{wb} = 450$ bar. Temperature declines become smaller for increased downhole temperature. In the production well it is of course desirable to reduce temperature decline during fluid upflow as much as possible. This can be achieved by increasing downhole pressures, which however will require increased power consumption in the fluid circulation systems.
From this discussion it is apparent that optimal operation of a CO$_2$-EGS system will involve complex tradeoffs between reservoir heat extraction and power consumption in the fluid circulation system.

Fig. 7 presents static pressure profiles in CO$_2$ injection wells for 80 bar wellhead pressure and different wellhead temperatures. Downhole pressures decrease with increasing wellhead temperatures, and more so for adiabatic than for constant temperature conditions. This is because for adiabatic conditions wellbore temperatures are larger, and accordingly fluid densities are smaller. The differences in downhole pressures range from 5.3 to 33.3 bar (Table 2).

Table 2. Downhole pressures in CO$_2$ injection wells for 80 bar wellhead pressure and different wellhead temperatures, for isothermal and adiabatic conditions.

<table>
<thead>
<tr>
<th>T (°C)</th>
<th>$P_{ad}$ (bar)</th>
<th>$P_{isoth}$ (bar)</th>
<th>$\Delta P$ (bar)</th>
</tr>
</thead>
<tbody>
<tr>
<td>10</td>
<td>574.09</td>
<td>579.42</td>
<td>5.33</td>
</tr>
<tr>
<td>20</td>
<td>545.57</td>
<td>558.58</td>
<td>13.01</td>
</tr>
<tr>
<td>30</td>
<td>500.79</td>
<td>534.13</td>
<td>33.34</td>
</tr>
</tbody>
</table>
Figure 7. Static pressure profiles in CO\textsubscript{2} injection wells for 80 bar wellhead pressure and different wellhead temperatures. Profiles are shown for isothermal as well as adiabatic conditions.

**RESERVOIR HEAT EXTRACTION**

**FIVE-SPOT FRACTURED RESERVOIR PROBLEM**

In order to compare CO\textsubscript{2} and water as heat transmission fluids, we consider an idealized fractured reservoir problem whose parameters were loosely patterned after conditions at the European EGS site at Soultz (see Table 3; Baria et al., 2005; Dezayes et al., 2005). Instead of an injector-producer doublet, we consider a five-spot well configuration with a basic pattern area of 1 km\textsuperscript{2} (production-injection well distance of 707.1 m; Fig. 8). This geometry is often chosen for fundamental studies of reservoir behavior, because the high degree of symmetry makes it possible to limit the model domain, which greatly simplifies the modeling problem (Sanyal and Butler, 2005). As seen in Fig. 8, the computational grid needs to cover only 1/8 of the domain,
Figure 8. Five-spot well pattern with computational grid for modeling a 1/8 symmetry domain.

Table 3. Parameters for five-spot fractured reservoir problem.

<table>
<thead>
<tr>
<th>Formation</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>thickness</td>
<td>305 m</td>
</tr>
<tr>
<td>fracture spacing</td>
<td>50 m</td>
</tr>
<tr>
<td>permeable volume fraction</td>
<td>2%</td>
</tr>
<tr>
<td>permeability</td>
<td></td>
</tr>
<tr>
<td>porosity in permeable domain*</td>
<td></td>
</tr>
<tr>
<td>rock grain density</td>
<td>2650 kg/m$^3$</td>
</tr>
<tr>
<td>rock specific heat</td>
<td>1000 J/kg/°C</td>
</tr>
<tr>
<td>rock thermal conductivity</td>
<td>2.1 W/m/°C</td>
</tr>
</tbody>
</table>

| Initial Conditions            |                                |
| reservoir fluid              | all CO$_2$, or all water       |
| temperature                  | 200 °C                         |
| pressure                     | 500 bar                        |

| Production/Injection          |                                |
| pattern area                  | 1 km$^2$                       |
| injector-producer distance   | 707.1 m                        |
| injection temperature        | 20 °C                          |
| injection pressure (downhole) | 510 bar                        |
| production pressure (downhole)| 490 bar                        |

* we include some wall rock in the definition of the fracture domain
but results will be given on a “full well” basis. We use a two-dimensional 5-point grid of 36 square blocks with 70.71 m side length. Formation parameters, initial conditions, and production-injection specifications are given in Table 3.

We assume three orthogonal fracture sets with 50 m spacing, so that the matrix blocks enclosed by the fractures are cubes with (slightly less than) 50 m side length. The matrix blocks are assumed impermeable and are sub-gridded into 5 continua using the MINC approach (Pruess and Narasimhan, 1985). As colder injected fluid is migrating through the fracture system, it is heated by conduction from the matrix blocks. The problem is run in two variations, assuming the reservoir fluid to be either (liquid) water or supercritical CO$_2$. The simulations use our TOUGH2 code with a special fluid property module "EOSM" for mixtures of water and CO$_2$ (Pruess, 2004a, 2004b). Net heat extraction rates were calculated as $G_i = F_i h_i - (F_i h_i)_{inj}$ ($i = $ CO$_2$, water), where $F_i$ and $h_i$ are mass flow rate and produced specific enthalpy, respectively, and $h_{i,inj}$ is injection enthalpy, evaluated at downhole conditions of $(T, P) = (20 \, ^\circ C, 510 \, \text{bar})$. Results for heat transfer rates and cumulative heat extraction are shown in Fig. 9.

For the system with CO$_2$ as heat transmission fluid, heat extraction rates are seen to be approximately 50% larger than for water, which represents a very substantial acceleration of energy recovery. Fig. 10 shows a comparison of mass flow rates between the CO$_2$ and water cases. CO$_2$ mass flow rates are larger than those for water by a factor of approximately 3.7 initially, and decrease less over time than water flow rates. The initial factor of 3.7 seems surprisingly large, in view of the fact that the parameter group ($\rho/\mu$) is only about a factor 1.9 larger for CO$_2$ than for water at initial reservoir conditions (see Fig. 2). However, for lower temperatures water viscosity increases much more than the viscosity of CO$_2$, giving CO$_2$ an additional advantage for flow in the vicinity of the injection well. Profiles of fluid pressures along a line from production to injection well show that for water much of the total pressure drop available for fluid circulation occurs in the cooled region near the injection well, while for CO$_2$ pressure gradients are only moderately stronger near the injection well than near the production well (Fig. 11).
Fig. 9. Rate of net heat extraction (top) and cumulative heat produced (bottom) for the five-spot fractured reservoir problem (full well basis). The top frame also shows the ratio of heat extraction rates for the CO$_2$ and water systems.
Figure 10. Mass flow rates for the five-spot problem. The ratio of flow rates in the CO₂ and water systems is also shown.

Figure 11. Pressure and temperature profiles along a line from production to injection well after a simulation time of 25 years.

SENSITIVITY STUDIES
Additional simulation studies were performed for reservoir parameters that were identical to the case considered above, except that different initial reservoir temperatures were used. Fig. 12 shows simulated ratios of net heat extraction rates for CO₂ and water for reservoir temperatures of 240, 200, 160, and 120 °C. (The 200 °C case is the one discussed above.) The comparison
shows that the acceleration of heat recovery when using CO\(_2\) as compared to water is greater at lower temperatures. This suggests that serious consideration should be given to using CO\(_2\) as heat extraction fluid not only for high-temperature resources that would be used for electricity generation, but also for lower temperature systems that would be produced for direct heat applications.

Fig. 13 compares the reference case with two alternatives, (1) injection temperature increased from 20 to 40 °C, and (2) reservoir pressure decreased from 500 to 400 bar. For higher injection temperature, the advantage of CO\(_2\) as heat extraction fluid is reduced, while for lower reservoir pressure it is increased.

Figure 12. Ratios of net heat extraction rates (CO\(_2\) vs. water) for different initial reservoir temperatures.
CO₂ STORAGE

Operating with CO₂ as heat extraction fluid, the reference case considered above generates approximately Q = 75 MW of thermal power, which for an assumed heat rejection temperature of 20 °C corresponds to a potential rate of mechanical work of $W = (1 - T_{\text{rej}}/T_{\text{pro}})Q$. For $T_{\text{rej}} = 293.15$ K and $T_{\text{pro}} = 473.15$ K, the potential mechanical work is $W = 0.38Q = 28.5$ MW. Assuming a utilization efficiency of 0.45 (Sanyal and Butler, 2005), this translates into an electric power generation of $G = 0.45W = 12.83$ MW. The average mass production rate is approximately 280 kg/s, from which we deduce a fluid circulation requirement of approximately $280/12.83 = 21.8$ kg of CO₂ per MW electric power. No information is currently available on the fraction of CO₂ mass flow that would be lost in the reservoir due to different mechanisms; the loss rate likely depends on the site-specific permeability, porosity, water chemistry and mineralogy of the EGS reservoir. Long-term (almost one year) water circulation tests that were conducted at an experimental site at Fenton Hill, New Mexico, incurred water losses of 7-12 % of injected rates (Duchane, 1993). As fluid losses generally decrease over time, it appears reasonable for a first rough estimate to assume that long-term operation of a CO₂-based EGS would incur fluid losses on the order of 5 % of injected rate. From this figure, CO₂ losses can be estimated at approximately 1 kg/s per MW electric power, or 1 tonne per second per 1,000 MW.
To gain a perspective on this number, note that a large coal-fired power plant of 1,000 MW electric capacity generates approximately 30,000 tonnes of CO$_2$ per day (Hitchon, 1996). Our estimate then suggests that 1,000 MWe of installed CO$_2$-EGS could achieve geologic storage of the CO$_2$ generated by almost 3,000 MWe of coal-fired power generation. While these estimates are very rough and preliminary, they suggest that CO$_2$-EGS could provide a very large potential for geologic storage.

**CHEMICAL ISSUES**

A CO$_2$-based EGS is expected to comprise three zones (Fouillac et al., 2004; Ueda et al., 2005):

- **Zone 1** - The inner zone or "core" of the system, from which all water has been removed by dissolution into the flowing CO$_2$ stream, so that the fluid consists of a single super-critical CO$_2$ phase. This is the main volume from which thermal energy is extracted by the flowing CO$_2$.

- **Zone 2** - Surrounding the inner zone is an intermediate region that contains a two-phase mixture of CO$_2$ and aqueous fluid.

- **Zone 3** - The outer region affected by the EGS activities. The fluid is a single aqueous phase with dissolved and chemically active CO$_2$.

Process behavior and issues are expected to be quite different in the different zones. This is especially true for chemical interactions. The outer reservoir zone would likely experience a combination of dissolution and precipitation effects that could impact reservoir growth and longevity. At the elevated temperatures of EGS, fluid-mineral reactions would be quite fast, providing a very favorable potential for rapid sequestration of CO$_2$ in the form of solid minerals (Fouillac et al., 2004; Ueda et al., 2005). The long-term behavior of the outer zone will be crucial for sustaining energy recovery, for estimating CO$_2$ loss rates, and for figuring tradeoffs between power generation and geologic storage of CO$_2$. The absence of water in the inner zone poses unique questions, as little is presently known about the geochemistry of non-aqueous systems. However, it seems clear that CO$_2$ would be a far less effective solvent than water. This would reduce the potential for dissolution and subsequent reprecipitation of minerals, and avoid
problems of scaling and formation plugging (Brown, 2000). It appears likely that prolonged exposure to supercritical CO\textsubscript{2} will cause dehydration reactions that would remove loosely bound water from rock minerals. Such reactions may reduce the molar volume of the minerals involved, which would increase porosity and permeability of the formations, and may promote reservoir growth (Pruess and Azaroual, 2006).

Aqueous solutions of CO\textsubscript{2} can be quite corrosive, and can dissolve different rock minerals, as well as attacking steel liners and casings used in well construction (Xu et al., 2005). However, aqueous fluids initially present in an EGS reservoir would be quickly removed by dissolution (evaporation) into the flowing CO\textsubscript{2} stream. Continuous operation of a CO\textsubscript{2}-EGS would be expected to produce a rather dry CO\textsubscript{2} stream that would not pose corrosion problems for production wells.

**CONCLUDING REMARKS**
At typical temperature and pressure conditions anticipated for EGS - approximately 200 °C and a few hundred bars - CO\textsubscript{2} is a supercritical fluid with liquid-like density and gas-like viscosity. Its thermophysical properties make it quite attractive as a heat transmission fluid. Our exploratory studies suggest that CO\textsubscript{2} is superior to water in its ability to mine heat from an EGS reservoir. CO\textsubscript{2} appears to offer advantages for wellbore hydraulics as well, which may lead to reduced power consumption for maintaining fluid circulation. The geochemistry of supercritical CO\textsubscript{2}, as opposed to aqueous solutions of CO\textsubscript{2}, is not well characterized, and needs to be understood so that long-term behavior of reservoir porosity and permeability may be predicted.

Specific results of our modeling studies can be summarized as follows.

- Due to much larger expansivity and compressibility as compared to water, supercritical CO\textsubscript{2} will generate much stronger buoyancy forces between injection and production wells. This will reduce power consumption for the fluid circulation system, and may possibly allow adequate fluid circulation without external pumping.
- The compression of CO\textsubscript{2} flowing down an injection well and the expansion flowing up a production well will give rise to substantial temperature changes of as much as 10–25 °C.
Specifically, CO$_2$ will heat upon compression and cool upon expansion. These effects are in addition to temperature changes due to heat exchange between the wellbore and its surroundings and will tend to reduce the buoyant drive.

- For a given total pressure drop between injection and production wells, CO$_2$ will generate on the order of 4 times larger mass flows and 50% larger net heat extraction rates as compared to water.

- The advantages of CO$_2$ in terms of accelerated energy extraction become larger for lower reservoir temperatures and pressures. CO$_2$ may be an attractive heat transmission fluid not only for high-temperature resources that would be used for electricity generation, but may offer even greater advantages for direct heat applications of lower-temperature resources.

- Based on fluid losses observed during long-term flow tests of the Fenton Hill hot dry rock system, fluid losses for CO$_2$-driven EGS are estimated as on the order of 1 tonne per second per 1,000 MW electric capacity. This means that 1,000 MWe of EGS-CO$_2$ could store all the CO$_2$ generated by 3,000 MWe of coal-fired power plants.

Fluid losses are an unavoidable aspect of engineered geothermal systems. Whereas the loss of water in a "conventional" EGS operation would be unfavorable and costly, fluid loss in an EGS system running with CO$_2$ would offer geologic storage of CO$_2$. Such storage may provide economic benefits and incentives in future carbon management scenarios where atmospheric emissions of CO$_2$ would be taxed, and avoidance of emissions could provide an additional revenue stream that would improve the economics of EGS.

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REFERENCES


