

# Use glycerol to dehydrate supercritical carbon dioxide

## This technology increases hydrocarbon recovery

M. SWADENER, Denbury Resources Inc., Plano, Texas; J. LUNDEEN, K. FISHER and C. BEITLER, Trimeric Corporation, Buda, Texas

Twenty years after the first two glycerol dehydration units were built for supercritical carbon dioxide (CO<sub>2</sub>) use, Denbury Resources found a need to build new units to process the increasing amount of CO<sub>2</sub> production from the Jackson Dome area in Mississippi. Denbury has now completed and started up five new glycerol-based dehydration facilities in the past five years, with other units under construction. Glycerol is used to dehydrate CO<sub>2</sub> when high-pressures and non-idealities cause excessive vapor-phase glycol (i.e., ethylene, diethylene or triethylene glycol) losses, making normal glycol-based dehydration uneconomical in comparison with glycerol-based dehydration. Denbury uses the supercritical CO<sub>2</sub> primarily for enhanced oil recovery (EOR).

This article presents reasons for Denbury's recent surge in glycerol unit construction as well as types of other applications where glycerol dehydration may be well suited. Differences between using glycerol and glycol for dehydration with supercritical, dense-phase CO<sub>2</sub> gas streams will also be discussed. Since commercial process simulators are not well suited for designing this type of system, a model was developed to rigorously predict phase equilibria in the absorber and simulate other process equipment. The simulator and other data used to build the new glycerol dehydration units are reviewed along with presenting key design features of the dehydration plants.

### Denbury's new and existing CO<sub>2</sub> dehydration facilities.

Denbury is a growing independent oil and natural gas company. It's the largest oil and natural gas operator in Mississippi; it owns the largest reserves of CO<sub>2</sub> used for tertiary oil recovery east of the Mississippi River, and holds significant operating acreage in Louisiana, Alabama, Texas, Montana, North Dakota and Wyoming.

Denbury's CO<sub>2</sub> source field, Jackson Dome, located near Jackson, Mississippi, was discovered during the 1970s while being explored for hydrocarbons (Fig. 1). This significant source of CO<sub>2</sub> is the only one known of its kind in the US east of the Mississippi River. Mississippi's first EOR project began in the mid 1980s in the Little Creek Field following the installation of Shell Oil Company's Choctaw CO<sub>2</sub> pipeline. The 183-mile Choctaw Pipeline (now referred to as the NEJD Pipeline) transported CO<sub>2</sub> produced from Jackson Dome to the Little Creek Field. While the CO<sub>2</sub> flood proved successful in recovering significant oil amounts, commodity prices at that time made the project unattractive and Shell later sold its oil fields in this area, as well as the CO<sub>2</sub> source wells and pipelines.

Currently, more than 1.0 billion cu ft per day (Bscfd) of CO<sub>2</sub> is produced from Jackson Dome Area operations. The pipeline network has been expanded from the single NEJD Pipeline including the Free State Pipeline to East Mississippi; the Delta Pipeline going into northeast Louisiana; and the recent completion of the Green Pipeline that delivers CO<sub>2</sub> across the Gulf Coast Region from the end of the NEJD Pipeline to south of Houston. Source well operating pressures range from 800 psig to 4,000 psig. Jackson Dome is generally referred to as the CO<sub>2</sub> source, but it is also the name of the original central dehydration facility that uses a glycerol process to dehydrate supercritical CO<sub>2</sub>-rich streams. The Jackson Dome dehydration unit was built by Shell and based on research performed in the 1980s to understand the dynamics of dehydrating supercritical CO<sub>2</sub> with glycols and glycerols.<sup>1-5</sup> The research showed that glycerol can dehydrate supercritical CO<sub>2</sub> more effectively than glycols. Critical pressure and temperature for pure CO<sub>2</sub> are 1,070 psia and 87.8°F, respectively. It should be noted that a glycerol unit built 20 years ago and operated by a different company is believed to be no longer operational.<sup>6</sup>

In 2003, an expansion to the Jackson Dome dehydration plant was completed, adding internals to the inlet separator and contactor to almost double the capacity. Since then, Denbury has built several new glycerol-based CO<sub>2</sub> dehydration units to keep pace with its increase in CO<sub>2</sub> production. Since Denbury continues to find high-pressure source wells, it is advantageous to dehydrate the CO<sub>2</sub> with glycerol at pipeline conditions without compression. The new dehydration units are described briefly in Table 1. The gas composition is generally 99.4% CO<sub>2</sub>, 0.3% N<sub>2</sub> and 0.3% C<sub>1</sub>.



FIG. 1 Dehydration facility map.

The post-expansion process performance at Jackson Dome was used to validate modeling results. Improvements were also made to the new dehydration plants based on Denbury's Jackson Dome experience and advancements with process technology.

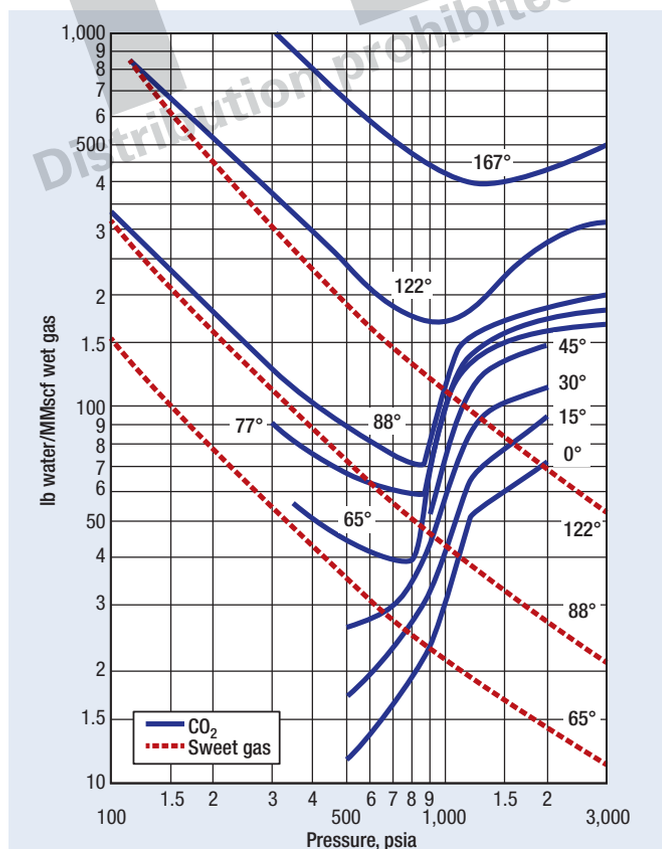
## Dehydration of supercritical CO<sub>2</sub> streams with glycerol.

The properties of supercritical CO<sub>2</sub> impact the dehydration system design in two ways:

- Mutual solubilities of the desiccant and CO<sub>2</sub>
- Saturated water content of CO<sub>2</sub> with temperature and pressure.

Data taken in a laboratory setting, and from field tests, concluded that glycerol had a much lower solubility in supercritical CO<sub>2</sub> than did ethylene glycol (EG), diethylene glycol (DEG) and triethylene glycol (TEG). Table 2 illustrates that TEG is 10–200 times more soluble than glycerol in CO<sub>2</sub> over a pressure range of 1,200 psig–2,000 psig. This means that TEG would have excessive vapor losses and makeup requirements, making normal glycol-based dehydration uneconomical. Operational data from Jackson Dome supports the glycerol solubility data from these sources. A gross material balance of the glycerol added to the system indicates 4 ppmv–5 ppmv of glycerol is soluble at 1,300 psig and 115°F.

Table 2 also shows that, unlike the glycols, glycerol solubility in CO<sub>2</sub> did not increase significantly with lower CO<sub>2</sub> temperatures. However, due to the high glycerol viscosity and rapidly increasing supercritical CO<sub>2</sub> density as the temperature decreases, glycerol would probably not be used to dehydrate CO<sub>2</sub> below 100°F. In addition, the CO<sub>2</sub> solubility in glycerol was also determined to be only one-tenth of the solubility in glycols. This results in lower flashing losses.



**FIG. 2** Solubility limit for water in pure CO<sub>2</sub>.<sup>7</sup> (Used with permission from GPSA).

Water solubility in pure CO<sub>2</sub> is well documented in several sources.<sup>7</sup> Fig. 2 shows the maximum water amount that can be dissolved into a homogeneous CO<sub>2</sub> phase—for the purpose of this article, it is called the “solubility limit” for water in CO<sub>2</sub>. As shown in Fig. 2, at a fixed temperature, the solubility limit for water decreases with increasing pressure up to near the critical pressure. As the pressure increases beyond the critical pressure (1,070 psia), the solubility limit for water increases. However, small concentrations of other gases can significantly impact the water-holding capacity of CO<sub>2</sub>.<sup>4</sup> Knowing the correct moisture content of the supercritical CO<sub>2</sub> ensures that the absorber and regeneration equipment are not undersized.

Other parameters that are important are the density of CO<sub>2</sub>, which can change greatly in the presence of small amounts of inert gases and hydrocarbons, and the viscosity of glycerol, which is significantly different than glycol (1,490 cps and 47.8 cps, respectively, at 20°C).<sup>4</sup>

**Potential uses of glycerol-based dehydration.** Dehydration was originally developed for essentially pure supercritical CO<sub>2</sub>—streams containing high CO<sub>2</sub> content that need to be dehydrated at high pressure. An example to use this technology is the stream dehydrated using glycerol in Hungary.<sup>6</sup> The Hungarian stream had 81% CO<sub>2</sub> with the remainder consisting of methane—dehydrated at a pressure near 2,000 psi. Other potential applications of the glycerol technology may exist where a combination of high CO<sub>2</sub> content and a need to dehydrate at high pressure create the operating conditions and compositions where glycerol may have lower solubility losses than glycols and better overall economics.

**Glycerol dehydration simulation development.** A simulation was developed specifically for supercritical dehydration of CO<sub>2</sub> with glycerol. This is because the standard property sets in most process simulators perform poorly when modeling fluid properties and phase equilibrium at near-critical conditions. Careful review of the thermodynamic parameters in the process simulator is required, with adjustments made to match reliable data. Key modeling aspects will be discussed shortly.

**TABLE 1.** Denbury's new glycerol-based CO<sub>2</sub> dehydration units

Plant	Gluckstadt	Barksdale North	Barksdale South	Trace	Gluckstadt	DRI Dock
Operation date	2006	2007	2008	2010	2010	2011
Flow, MMscfd	100	300	300	150	300	300
Operating pressure, psig	1,550	1,500	1,500	1,500	1,550	TBD
Operating temperature, °F	120	110	110	110	115	TBD
Water content in, lb/MMscf	201	201	201	201	201	201
Water content out, lb/MMscf	20	20	20	20	20	20
Reboiler duty, MMBtu/hr	2.05	6	6	2.75	6	6

**TABLE 2.** Glycol and glycerol solubility in CO<sub>2</sub> (ppmv)<sup>1</sup>

	1,200 psi		2,000 psi	
	TEG	Glycerol	TEG	Glycerol
73°F	1,800	12.3	2,800	30
95°F	390	8.2	—	37
115°F	30	3.3	1,900	42

**Absorber K-values.** An important aspect of designing glycerol-based dehydration systems for supercritical CO<sub>2</sub> is to have a good estimate of the K-value (i.e., phase equilibrium distribution coefficient) for water at the conditions in the absorber. Unreliable CO<sub>2</sub>-water equilibrium data can produce erroneous results in the absorber that adversely affect predicting operations in downstream vessels. Absorber K-values for water were calculated using the thermodynamic relationship:

$$K = y/x = \gamma P^{\text{sat}} \Phi^{\text{sat}} \eta / \Phi P \tag{1}$$

where:

K = K-value, phase equilibrium distribution coefficient

y = Mole fraction water in the dense CO<sub>2</sub> phase

x = Mole fraction water in the liquid phase

γ = Activity coefficient for water (from vapor-liquid equilibrium data discussed in the following subsection)

P<sup>sat</sup> = Vapor pressure of pure water at the system temperature (from steam tables)

Φ<sup>sat</sup> = Fugacity coefficient for pure water vapor at the system temperature and P<sup>sat</sup> (assumed to be unity)

η = Poynting correction factor for water (from the component partial molar volumes)

Φ = Fugacity coefficient of water for the vapor phase

P = System operating pressure (known).

Most of the parameters in Eq. 1 are readily known or accessible from literature as indicated previously. However, it is the calculation of the vapor-phase fugacity of water in dense-phase CO<sub>2</sub> near the critical conditions that is more difficult.

Field data were used to validate the accuracy of the simulation in predicting K-values and resulting water contents over a range of actual operating conditions. Table 3 shows that the treated gas water contents are in very close agreement with the predicted equilibrium water content for the lean glycerol solution at the absorber conditions (i.e., within 5%–17% of the measured data). This indicates that the method discussed is reliable for predicting K-values and dehydrator outlet water contents.

**Water-glycerol phase behavior.** The vapor-liquid equilibrium (VLE) of water in glycerol is important because it is used to:

- Predict the K-values in the absorber (activity coefficient of water)
- Determine the lean glycerol water content from the reboiler and extra stripping column.

VLE data for water and glycerol (in the concentration and temperature ranges in the absorber and reboiler) were obtained and used to regress the adjustable parameters in the Wilson activity coefficient model.

**CO<sub>2</sub> solubility in glycerol.** It is important to know the CO<sub>2</sub> solubility in glycerol for several reasons. First, it provides a means

**TABLE 3. Comparison to known field data**

Treated gas measured, lb/MMscf	Predicted equilibrium limit, lb/MMscf	% difference
7.0	6.7	-4.3
9.1	7.9	-13.2
7.2	7.8	8.3
9.8	10.9	11.2
13.2	15.5	17.4
21.4	25.1	17.3

to determine how much CO<sub>2</sub> is picked up in the absorber and will evolve in the flash tank. Second, the CO<sub>2</sub> solubility in glycerol is important in predicting the potential for two-phase flow through the heat exchangers in the process as the temperatures and pressures of the streams vary. Equilibrium data were used with the Wilson activity coefficient model to predict CO<sub>2</sub> solubility in glycerol.

**Viscosity.** The viscosity estimation method in the process simulator that most closely predicted viscosities found in the literature was selected. Viscosity data were used to calculate equipment pressure drop and model absorber hydraulics.

**Thermal conductivity.** A method that accurately predicted the thermal conductivity for both 50-50 glycerol-water mixtures and high-purity glycerol streams was selected.

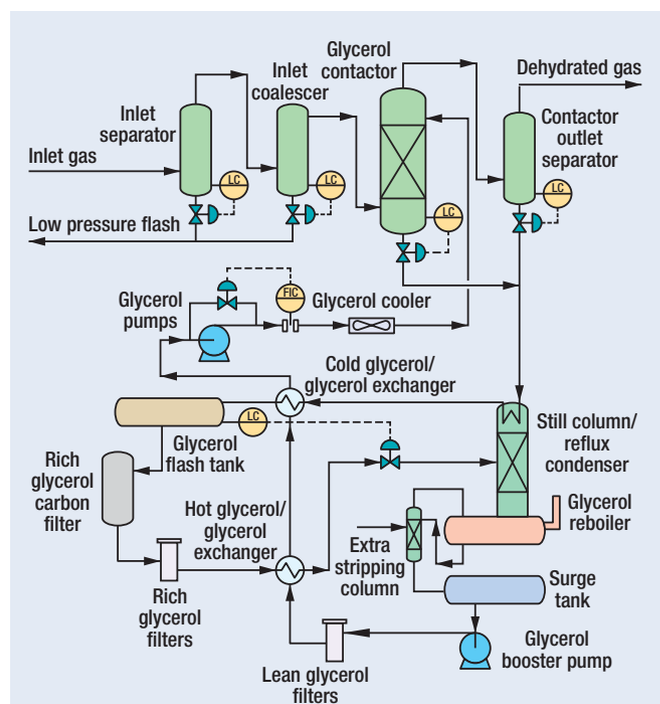
**Density.** The densities of the inlet and outlet CO<sub>2</sub> streams from the absorber were calculated using the Peng-Robinson equation of state. The calculated densities compared reasonably well to those obtained from other sources.

**Glycerol solubility in dense-phase CO<sub>2</sub>.** The glycerol solubility in dense-phase CO<sub>2</sub> has been reported in the literature and was used to estimate glycerol losses from the absorber.<sup>1</sup>

**Glycerol-based dehydration design and research.**

Fig. 3 illustrates a typical process flow diagram of the glycerol-based CO<sub>2</sub> dehydration units for Denbury's new facilities. The glycerol process operates much in the same manner as a TEG unit so a detailed process description is not provided.<sup>7</sup>

Operational experience at Jackson Dome and advancements in process technology influenced the new glycerol process designs. Construction material for all process piping and vessels at the original Jackson Dome facility is 316/316L SS. In general, many of the same material choices used for glycol are appropriate for glycerol. The following materials choices were different in the new units compared to the original Jackson Dome plant:



**FIG. 3** Process flow diagram for Denbury's glycerol-based CO<sub>2</sub> dehydration unit.



**FIG. 4** New Denbury glycerol unit.

- The contactor was built with 316L SS in the lower section and a carbon steel shell above the bottom 2 ft. of packing. This was done since the gas is sub-saturated as it moves counter-current to the glycerol through the packed section. This concept was successfully proven elsewhere.

- Carbon steel was used for all glycerol piping except for downstream of the glycerol flash tank. In this area, 316L SS was used instead to provide low temperature integrity if CO<sub>2</sub> vented through the control valve if the liquid level control in the flash absorber was lost.

- Carbon steel was used for the rich/lean exchanger shells, the glycerol air cooler, the reboiler shell and filter vessels.

Design elements of note from the Jackson Dome process that were incorporated into the new glycerol units will be described further. Isolation valves for the hot and cold glycerol exchangers were included for periodic cleaning/maintenance due to fouling from upset conditions or high skin temperatures. Due to high glycerol viscosity, glycerol booster pumps were used to ensure continuous flow from the reboiler through the hot and cold glycerol exchangers. A fire tube liner was included in the reboiler to moderate the skin temperature in contact with the glycerol, thus, reducing the degradation rate. The glycerol flash tank and still were constructed of 316L SS due to the wet CO<sub>2</sub> in those areas.

Other key design aspects include using low-maintenance and low-cost horizontal submersible pumps or gear pumps for glycerol circulation and incorporating an extra stripping column for enhanced water removal. A provision for a filter coalescer upstream of the glycerol contactors for removal of water with high salt and chloride content as the well declines was also included in the design. Jackson Dome data was used to size the absorbers in the new units.

As work with supercritical CO<sub>2</sub> continues, there have been some areas of research and innovation. A series of tests were conducted to quantify the performance of liquid and solid hydrogen sulfide (H<sub>2</sub>S) scavengers at process conditions when 50 ppm–100 ppm of H<sub>2</sub>S was detected in one of the source wells. The test results indicated a reduced efficiency of these scavenging products for H<sub>2</sub>S removal compared to the typical performance on natural gas.<sup>8</sup> Another area of study was glycerol solubility in CO<sub>2</sub>. Although the solubility is much less than glycols, the price of glycerol was over \$1/lb in 2008 when oil prices were at their peak. Additives were identified to further reduce glycerol solubility in CO<sub>2</sub> and a full-scale test is planned. Fig. 4 shows a new glycerol unit.

**Summary.** The rigorous review of thermodynamic properties associated with the dehydration of supercritical CO<sub>2</sub> with

glycerol was required for the development of a reliable process-simulation tool to design new glycerol processes. Denbury's operational experience at Jackson Dome was important in constructing high-performing glycerol-based units for dehydration of supercritical CO<sub>2</sub>. Using glycerol dehydrators allows Denbury to process more CO<sub>2</sub> for increased hydrocarbon recovery. The information provided can be applied for designing other high-CO<sub>2</sub>, high-pressure applications, as well, even if the stream is not essentially pure CO<sub>2</sub>. **HP**

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**Mark Swadener** is a reservoir engineer at Denbury Resources in Plano, Texas. He has seven years of experience in facilities, production/operations and reservoir engineering for West Texas and Texas Gulf Coast oil fields. Most recently, Mr. Swadener was the operations engineer at Jackson Dome, responsible for the completion, production and maintenance of CO<sub>2</sub> source wells and facilities. He received his BS degree in mechanical engineering from Southern Methodist University in Dallas.



**Joe Lundeen** is a principal engineer at Trimeric Corp. in Buda, Texas. He has 21 years of experience in process engineering, process troubleshooting and facility installation for oil and gas production and CO<sub>2</sub> processing clients. Mr. Lundeen's recent experience has focused on dehydration, contaminant removal and transport of super-critical CO<sub>2</sub>. He earned BS and MS degrees in chemical engineering from the University of Missouri—Rolla and Texas A&M—College Station, respectively.



**Kevin Fisher** is a principal engineer at Trimeric Corp. in Buda, Texas. He has over 20 years of experience in process engineering, research and development, and troubleshooting for oil and gas production and oil refining clients, as well as for private and government-sponsored research programs. Mr. Fisher specializes in the areas of acid-gas removal, CO<sub>2</sub> capture, gas processing and gas dehydration. He earned an MS degree in chemical engineering from the University of Texas at Austin, and BS degrees in chemical engineering and chemistry from Texas A&M—College Station and Sam Houston State University, respectively.



**Carrie Beitler** is a senior engineer at Trimeric Corp. in Buda, Texas. She has over 15 years of experience in process engineering, process modeling and optimization of unit operating in the natural gas, petroleum refining and CO<sub>2</sub> processing areas. She also specializes in the development of process design packages for the fabrication of open-art technology such as caustic scrubbers, acid-gas injection units, glycol dehydrators and amine treaters. She earned a BS degree in chemical engineering from Purdue University.