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Density of Aqueous Solutions of CO_2 *

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Abstract

In this report, we present a numerical representation for the partial molar volume of CO_2 in water and the calculation of the corresponding aqueous solution density. The motivation behind this work is related to the importance of having accurate representations for aqueous phase properties in the numerical simulation of carbon dioxide disposal into aquifers as well as in geothermal applications. According to reported experimental data the density of aqueous solutions of CO_2 can be as much as 2-3 % higher than pure water density. This density variation might produce an influence on the groundwater flow regime. For instance, in geologic sequestration of CO_2 , convective transport mixing might occur when, several years after injection of carbon dioxide has stopped, the CO₂rich gas phase is concentrated at the top of the formation, just below an overlaying caprock. In this particular case the heavier CO_2 saturated water will flow downward and will be replaced by water with a lesser CO_2 content.

1 Introduction

Disposal of CO_2 into geological formations requires, among other conditions, the presence of a regional caprock for confinement. In order to avoid two-phase conditions in the injection line, geological disposal of CO_2 would be made at supercritical conditions. Under such conditions, CO_2 is less dense than water and would have a tendency to migrate to the top of the formation (Pruess and García, 2001). After a relatively long time (decades after ending injection) almost all gas would be stored at the top, underlying the caprock. Because the density of CO_2 saturated water is slightly higher than the density of pure water, density driven flow will occur enhancing the dissolution process by convective mixing. In such flow problems, numerical simulation capabilities that take into account variable density brines are crucial (Oldenburg and Pruess, 1995).

In this report we present a new correlation for the partial molar volume of CO_2 in water and the calculation of the corresponding aqueous solution density. This aqueous phase density correlation is then incorporated into the TOUGH2 Equation-of-State (EOS) module for carbon dioxide ECO2 (under development) (Pruess and García, 2001; Pruess et al., 1999). Currently, the brine density correlation considers only salinity effects with no CO_2 dependency (Battistelli et al., 1997). Figure 1 shows the computed brine density from 10 to 110 °C at various salt mass fractions (XNaCl=0.00, 0.05, 0.10 and 0.25) as implemented inside EWASG. The density increase due to salinity is up to almost 20%.

Experimental data for the system H_2O-CO_2 show that the carbon dioxide content produce an increase in aqueous phase density on the order of 2 to 3 %. Because the salinity effect is considerable higher, the dependency on CO_2 content is often ignored. This assumption is perfectly acceptable in geothermal applications. Nevertheless, as discussed before, it can conceal key processes in fluid flow dynamics of carbon dioxide sequestration.

Based on thermodynamic theory (Söhnel and Novotný, 1985; Anderson and Crerar, 1993), the density of aqueous solutions of CO_2 may be expressed as:

$$\rho_{aq} = \rho_1 + M_2 \cdot c - c \cdot \rho_1 \cdot V_\phi \tag{1}$$

where V_{ϕ} is the apparent molar volume of dissolved CO₂; M_2 is the molecular weight of CO₂; ρ_1 is the density of pure water and c is the CO₂ concentration

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Figure 1: Computed density and density increase of NaCl solutions for salt mass fraction (XNaCl) from 0 to 0.25 using the EWASG correlation.

expressed by the number of moles of solute in 1 m^3 of solution (for details see Appendix A).

2 Experimental Data and Preliminary Correlations

A large number of data exist on the solubility of CO_2 in water (Scharlin (1996) and Drummond (1981)). In spite of their significance, there is limited experimental data on densities of the aqueous system H₂O-CO₂-NaCl. Usually, there is two sets of data, one for the system H₂O-NaCl and another for the system H₂O-CO₂. Because the effect of salinity on density is considerably higher than the effect of carbon dioxide, the latter is often ignored. The most significant investigations of partial molar volume of CO₂ and density of aqueous solutions of CO₂ along with experimental conditions are listed in Table 1.

The data collected indicate that CO_2 saturated water is heavier than pure water below 300 °C and down to at least 5 °C. Some measurements were done at CO_2 saturation, while others were for very low mole fraction. Apparent molar volumes for data sets re-

Table 1: Experimental data for partial molar volume and density of aqueous solutions for the system CO_2 - H_2O .

NP *	T $[^{o}C]$	P[MPa]	Reference
24	5-20	6.4 - 29.5	Teng et al. (1997)
14	25 - 300	20-35	Hnědkovský et al. (1996)
5	274 - 297	19.6-29.4	Malinin (1974)
8	25 - 100	1.96	Ellis and McFadden (1972)
1	5 - 40	NR **	Parkinson and Nevers (1969)
1	25	NR **	Moore et al. (1982)

* Number of points used in this study ** Not reported

porting only densities and solubilities were calculated using the theory presented in Appendix A.

Experimental results show that the partial molar volume (approximated here as the apparent molar volume) is weakly dependent on CO_2 mole fraction (Parkinson and Nevers, 1969) and, for temperatures below $300^{\circ}C$ is independent of pressure (Malinin, 1974). Salinity effects on partial molar volume of CO_2 in water are not considered in this study. There are limited data that consider salinity. Malinin (1974) presented several data points for solutions of 1 M NaCl at high temperatures and pressures. His findings showed that the salinity effects are weak and within experimental uncertainty.

2.1 Previous Work

There are correlations available for the partial molar volume as a function of temperature. Most of these representations consider a polynomial fit as follows:

$$V_{\phi} = a + bT + cT^2 + dT^3 + eT^4 \tag{2}$$

Table 2 summarizes the different fits for Equation 2 found in the literature as well as other types of correlations.

2.2 Novel Correlation

The fit to Equation 2 proposed in this study considers 53 data points (See Figure 2). The following equation adequately represents the data with an $R^2 = 0.98$.

$$V_{\phi} = 37.51 - 9.585 * 10^{-2}T + 8.740 * 10^{-4}T^2 - 5.044 * 10^{-7}T^3$$
(3)

Equation 3 and 1 were used to compute densities of aqueous solutions of CO_2 at a pressure of 100 bar

a	b	с	d	е	Reference
37.36 1799.36 35.663 37.51	$-7.109 * 10^{-2} \\ -17.8218 \\ -5.960 * 10^{-2} \\ -9.585 * 10^{-2}$	$-3.812 * 10^{-5} \\ 0.0659297 \\ +6.308 * 10^{-4} \\ +8.740 * 10^{-4}$	$+3.296 * 10^{-6}$ $-1.0579 * 10^{-4}$ $-5.044 * 10^{-7}$	$-3.702 * 10^{-9} \\ +6.200 * 10^{-8}$	Andersen et al. (1992) * Enick and Klara (1990) ** Jonathan Ennis-King ¹ * This work *
$V_{\phi} = e^{154}$	$4.7881 - \frac{3582.452}{T} - 26.$	Iglesias and Moya (1992) ***			

Table 2: Correlations used by other authors for partial molar volume of CO₂ in water.

* Equation 2 V_{ϕ} is in $cm^3/mole$ and T is in degrees Celsius

** Equation 2 V_{ϕ} is in $cm^3/mole$ and T is in degrees Kelvin

*** V_{ϕ} is in $cm^3/mole$ and T is in degrees Kelvin

¹ Personal Communication



Figure 2: Fit of Partial Molar Volume Vs. Temperature. (a) Teng et al Teng et al. (1997), (b) Hnedkosky et al Hnědkovský et al. (1996), (c)Malinin Malinin (1974), (d) Ellis and McFadden Ellis and McFadden (1972), (e)Moore et al Moore et al. (1982)

for mole fractions (Xmole) of 0.02 and 0.05 (Figure 3). A maximum density increase of 2.5 % is obtained for a solution with a CO_2 mole fraction of 0.05.

As expected according Equation 1, the density variation of the aqueous solutions of CO_2 is nearly linear with respect to CO_2 mole fraction (See Figure 4). Figure 5 and 6 show a comparison between the



Figure 3: Computed density of aqueous solutions of CO_2 (P=100 bar)

measured and predicted values of the partial molar volume of CO_2 in water and the density of aqueous solutions of CO_2 , respectively. Predicted values using Equation 3 agree very well with the measured data.



Figure 4: Computed density of aqueous solutions of CO_2 (P=100 bar)



Figure 5: Comparison between measured and predicted partial molar volume of CO_2 in water



Figure 6: Comparison between measured and predicted density of aqueous solution of CO_2

3 TOUGH2-ECO2 Implementation

The proposed new formulation for the partial molar volume of CO_2 in water and the corresponding aqueous solution density is incorporated into the TOUGH2 Equation-of-State module ECO2 through the new subroutine COCO2. Only few changes are required in the rest of the module. Calls to subroutine COCO2 require the following arguments: aqueous phase temperature and pressure, CO_2 and Salt mass fractions, and brine density:

CALL COCO2(TX,PL,XG,XSA,DB)

The CO_2 dependency on brine density can be turned on through parameter specification in data block SELEC. A description of additional **SELEC**tion options is provided in Pruess et al. (1999).

- - 0: Brine density independent of CO_2 content
 - 1: Full dependence

The beta version of subroutine COCO2 is presented in Appendix B.

4 Nomenclature

- c molarity of solution (moles/m³)
- M molecular weight (kg/mol)
- n_i number of moles of the *i*-th component in the mixture (mol)
- P pressure (Pa)
- T temperature (°C)
- V volume (m³)
- \overline{V}_i partial molar volume of the i-th component (m³/mol)
- V_m molar volume (m³/mol)
- V_{ϕ} apparent molar volume of dissolved CO₂ (m³/mol)
- x_i molar fraction
- ρ density of the solution (kg/m³)
- ρ_1 density of water (kg/m³)

Subscripts

aq	aqueous phase
s	mixture
1	solvent (H_2O)
2	solute (CO_2)

Superscripts

o at infinite dilution

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A Density of Binary Aqueous Solutions

The partial molar volume is an important thermodynamic quantity since: (1) its value may be used to correct Henry's constant when the gas pressure is large (Prausnitz et al., 1986) and; (2) the knowledge of its value can be used to estimate the density of the solution. In this appendix we present an overview of theory related to partial molar volumes and estimation of the corresponding binary aqueous solutions density. For additional details the reader is referred to Prausnitz et al. (1986) and Anderson and Crerar (1993).

The definition of a partial molar property is applicable only to extensive properties differentiated at constant temperature and pressure. The total volume of a system with j-components can be expressed by the relation:

$$V = \sum_{i=1}^{j} n_i \overline{V_i} \tag{4}$$

where

$$\overline{V}_i = \left(\frac{\partial V}{\partial n_i}\right)_{T,P,n_{j\neq i}} \tag{5}$$

is the partial molar volume of the i-th component.

In practice, the apparent molar quantities are more easy to be determined, and the partial molar quantities are calculated from these. For binary solutions the apparent molar volume, V_{ϕ} , is defined as:

$$V_{\phi} = \frac{V - n_1 V_{m,1}}{n_2} \tag{6}$$

As illustrated in Figure 7, the apparent molar volume is the volume that should be attributed to the solute in the solution if one assumes that the solvent contributes the same volume it has in its pure state. Alternatively,

$$V = n_1 V_{m,1} + n_2 V_{\phi} \tag{7}$$



Moles of solute, n_2

Figure 7: Total volume of a solution as a function of solute concentration showing the difference between partial molar volume and apparent molar volume. Adapted from Anderson and Crerar (1993).

Dividing Equation 7 by $(n_1 + n_2)$ we obtain:

$$V_m = x_1 V_{m,1} + x_2 V_\phi \tag{8}$$

The apparent molar volume V_{ϕ} can be seen (Figure 8) to be the intercept on the solute axis.



Figure 8: Molar volume of a binary solution versus mole fraction showing the difference between partial molar volume and apparent molar volume for solution at composition A. Adapted from Anderson and Crerar (1993)

The partial molar volume of solute and solvent in a binary solution can be obtained from V_{ϕ} as:

$$\overline{V}_2 = \left(\frac{\partial V}{\partial n_2}\right)_{T,P,n_1} = V_\phi + n_2 \left(\frac{\partial V_\phi}{\partial n_2}\right)_{T,P,n_1} \tag{9}$$

at infinite dilution (i.e. $n_2 \rightarrow 0$) the following relation holds

$$\overline{V}_2^o = V_\phi^o \tag{10}$$

i.e. the partial molar volume of a solute at infinite dilution is equal to its apparent molar volume at infinite dilution.

The apparent partial molar volume can be expressed in terms of densities and molecular weights (Teng et al., 1997).

$$V_{\phi} = \frac{1}{n_2} \left(\frac{n_1 M_1 + n_2 M_2}{\rho} - n_1 \frac{M_1}{\rho_1} \right)$$
(11)

If the composition is expressed by the number of moles of solute in 1 kg of solvent (i.e. $n_2 = m n_1 = 1/M_1$), we obtain:

$$V_{\phi} = \frac{\rho_1 - \rho}{m\rho\rho_1} + \frac{M_2}{\rho} \tag{12}$$

Or, if the composition is expressed by the number of moles of solute in 1 m^3 of solution (i.e. $n_2 = c n_1 = (\rho - cM_2)/M_1$):

$$V_{\phi} = \frac{\rho_1 - \rho}{c\rho_1} + \frac{M_2}{\rho_1}$$
(13)

Either Equation 12 or 13 can be used to obtain an expression for aqueous density. From equation 13

$$\rho_{aq} = \rho_1 + M_2 \cdot c - c \cdot \rho_1 \cdot V_\phi \tag{14}$$

Or from equation 12:

$$\rho = \frac{1 + mM_2}{mV_{\phi} + \frac{1}{\rho_1}} \tag{15}$$

where m can be expressed in terms of mol fractions (x_1, x_2) as:

$$m = \frac{1}{M_1 \left(\frac{1}{x_2} - 1\right)} = \frac{x_2}{M_1 x_1} \tag{16}$$

$$x_2 = \frac{mM_1}{1 + mM_1} \tag{17}$$

Replacing equation 16 into equation 15

$$\rho = \frac{1 + \frac{x_2 M_2}{M_1 x_1}}{\frac{x_2 V_{\phi}}{M_1 x_1} + \frac{1}{\rho_1}}$$
(18)

After further simplification we obtain an alternative equation for the aqueous phase density

$$\frac{1}{\rho} = \frac{x_2 V_{\phi}}{M_T} + \frac{M_1 x_1}{\rho_1 M_T}$$
(19)

where $M_T = M_1 x_1 + M_2 x_2$

A.1 Ideal Solutions

For an *ideal solution*, the partial molar volumes of the species in solution are equal to the molar volumes of pure species at the same temperature and pressure. In this event, the total volume becomes:

$$V = n_1 V_{m,1} + n_2 V_{m,2} \tag{20}$$

Expressing the molar volumes in terms of densities $(V_{m,i} = \frac{M_i}{\rho_i})$:

$$V = n_1 \frac{M_1}{\rho_1} + n_2 \frac{M_2}{\rho_2} \tag{21}$$

$$\frac{n_1 M_1 + n_2 M_2}{\rho} = n_1 \frac{M_1}{\rho_1} + n_2 \frac{M_2}{\rho_2}$$
(22)

Dividing Equation 22 by the total mass:

$$\frac{1}{\rho} = \frac{X_1}{\rho_1} + \frac{X_2}{\rho_2} \tag{23}$$

Where $X_i = \frac{n_1 M_i}{n_1 M_1 + n_2 M_2}$ is the mass fraction of specie i.

B Subroutine COCO2

(inc)(line)	\bigcirc
01715	— c
01716	SUBROUTINE COCO2(TX, PX, XG, XSA, DBJ)
01717	- C IMPLICIT DOUBLE PRECISION (A-H,O-Z)
01718	— COMMON/GASLAW/R, AMS, AMA, CVNCG
01719	- COMMON/EOSG/ZA, AMSALT, GAST
01720	- COMMON/EOSEL/TE(16).FE(512)
01721	- COMMON/VISWAT/DW0
01722	COMMON/KONIT/KON, DELT, IGOOD
01723	- SAVE ICALL
01724	— DATA ICALL/0/
01725	- ICALL=ICALL+1
01726	IF(ICALL.EO.1) WRITE(11,899)
01727	899 FORMAT(6X, 'COCO2 0.9 19 September 2001', 6X,
01728	X'DENSITY OF AQUEOUS PHASE AS FUNCTION OF T, P',
01729	X' AND CO2 MOLE FRACTION')
01730	— C
01731	WHEN IE(9)>1 INCLUDE CO2 MOLE FRACTION DEPENDENCY.
01732	— C
01733	— DBC=DBJ
01734	— C Compute effective molecular weight
01735	<pre>EMW=(1.0-XG)/((1.0-XG-XSA)/AMS+XSA/AMSALT)</pre>
01736	- C Compute mole fraction.
01737	<pre> XMG=XG/(XG+AMA*(1XG)/EMW)</pre>
01738	— C Correlation for partial molar volume JULIO GARCIA (2001)
01739	IF(IE(9).EQ.1) THEN
01740	VPH=37.51-9.585E-2*TX+8.740E-4*TX*TX-5.044E-7*TX*TX*TX
01741	END IF
01742	- C Unit Conversion
01743	
01744	TOTM=(1.0-XMG)*EMW+XMG*AMA
01745	<pre>DBINV=(XMG*VPHI/TOTM)+(EMW*(1.0-XMG))/(DBC*TOTM)</pre>
01746	— DBJ=1./DBINV
01747	- CIDEAL SOLUTIONS
01748	- C Partial molar volumes of species in solution are
01749	- c equal to molar volumes of pure species.
01750	- c To be included only for testing purposes
01751	- C Get pure CO2 density at saturated conditions
01752	- c Calculate TOTI and TOT2 and evaluate density of aqueous solution.
01753 01754	C DBINVI=TOTI/DBC+TOT2/DCO2
01754	$- C \qquad DBZ=1./DBINVI$
01755	IF (IGOOD.NE.U) RETURN
01750 01757	CVISCOSILY OF COZ SATURATED AQUEOUS PHASE
UL/J/ 017E0	C Increate rew data points for this. To be included in a
01750	DETTION
01760	
01/00	