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On the water content in CO₂ + CH₄ and CO₂-rich mixtures: experimental and modelling evaluation at temperatures from 233.15 to 288.15 K and pressures up to 15 MPa

Valderio de Oliveira Cavalcanti Filho^a, Antonin Chapoy^{a,b,1} and Rod Burgass^a

^a*Hydrates, Flow Assurance & Phase Equilibria Research Group, Institute of Petroleum Engineering, Heriot-Watt University, UK*

^b*MINES ParisTech, CTP-Centre Thermodynamique des Procédés, 35, Rue Saint Honoré, 77305 Fontainebleau, France*

Abstract

An experimental and modelling investigation of water content in CO₂+CH₄ and CO₂-rich mixtures in equilibrium with hydrates or liquid water was carried out at temperatures between 233.15 and 288.15 K at pressures up to 15 MPa. Some measurements were undertaken in two-phase region, in the presence of hydrates, with liquid and vapour compositions also reported. Predictions from cubic-plus association SRK, SRK incorporating NRTL with Huron-Vidal mixing rules and multiparametric EoS-CG/GERG equations of state were compared with experimental data. In comparison with pure carbon dioxide, the addition of small amounts of impurities (permanent gases and/or hydrocarbons) resulted in a significant reduction in the water content of the fluid phases present. Overall, sCPA showed good agreement with experimental data, although SRK-HV-NRTL gave better results for some cases, despite the use of only two adjustable parameters. By contrast, multiparametric EoS-CG yielded poor representation of the experimental data. In the two-phase region, no matter the equation of state used, a tendency to underestimate water content in the liquid phase was observed.

Keywords: water content, CO₂+CH₄, CO₂-rich mixtures, sCPA, SRK+HV+NRTL, EoS-CG

¹ Corresponding author. Tel.: + 44 131 451 3797
E-mail address: a.chapoy@hw.ac.uk

1. Introduction

Carbon capture and storage (CCS), enhanced oil recovery (EOR) using alternate gas and water (WAG) injection, sour-gas-associated oil reservoirs exploration, dense phase transportation and subsea processing are only a few cases where, nowadays, CO₂-rich systems play an important role. In most of these situations, fluid phases encounter low temperatures and high pressures, as new technologies made possible the exploration of remote areas, namely in ultradeep water, such as the Brazilian pre-salt. The development of the production in such new frontiers demands carbon dioxide separation, conditioning, compression and final disposal.

Specifications for water content for such systems depend on several aspects. According to John Carrol [1], in the United States, the value is usually 7 lb/MMCF (about 112 kg/MM std m³), whereas in Canada, it is 4 lb/MMCF (about 65 kg/MM std m³), which might represent dew points as low as 235 K, at atmospheric conditions, depending on the gas composition. In Europe, EASEE-gas has a limit of 265.15K for dew points, referenced to a pressure of 7 MPa. Due to the requirement for transportation through subsea pipelines, offshore drying units might have higher specifications in order to prevent water condensation and hydrate formation.

In Brazilian offshore scenario, for instance, Andrade et al. [2] mentioned a specification of 2 lb/MMCF (circa 32 kg / MM std m³) for the exported treated gas from standard FPSO units operating in the deep water Campos Basin. For the recently discovered pre-salt fields, however, the authors presented a more detailed description of surface facilities designed to handle inlet natural gas streams containing up to 30% of CO₂ which are dehydrated to a specified water content as low as 1 ppmv, using molecule sieves. In addition, the description also includes membrane systems that are used to remove carbon dioxide from natural gas streams (maximum 5% CO₂), en route to further processing in onshore liquid recovery plants, and produces a secondary stream with high CO₂ concentration (up to 90% mole/mole) destined to reservoir reinjection. These units, and other process facilities dealing with CO₂-rich mixtures, may experience low temperatures during pressure drop (e.g., through valves, restrictions or porous media flow during WAG

1 injection), blowdown (controlled emptying) or depressurisation (accidental rupture) events.
2 In all these occurrences, phase transitions can take place and the presence of water might
3 lead to hydrate formation, resulting in partial or total blockages.

4 Accurate water content predictions are of utmost importance for the above-
5 mentioned processes. Predictions are required in order to dictate design and operation of
6 dehydration units and/or inhibitor injection pumps. Despite recent advances, no reliable
7 model for both high pressure and low temperature involving associative molecules (such as
8 water and carbon dioxide) is available. In addition, there is a lack of experimental data
9 which contributes to a tendency to overlook the influence of minor impurities.

10 Measurements for water content in CO₂ rich-mixtures are mainly restricted to CH₄ –
11 CO₂. Table 1 details the majority of data published to date [3–18]. Composition for all fluid
12 phases present are rarely reported and, instead, material balance, correlation predictions or
13 dry basis data are commonly presented. The work of Al Ghafri et al [15] is the only
14 exception, where vapour-liquid-liquid (VLLE), vapour-liquid (VLE) and liquid-liquid (LLE)
15 equilibrium compositions have been reported. Data for the upper quadruple point (H-L_w-
16 L_{CO₂}-V) for a wide range of carbon dioxide concentration are also available [5,15,17,19].

17 CH₄-H₂O and CO₂- H₂O mutual solubilities have been investigated over a wide range of
18 temperature and pressures [20–65]. Essential for parameter fitting purposes, these data
19 have been correlated using different equations of state (EoS). Huron Vidal mixing rules [66]
20 with Soave-Redlich-Kwong (SRK) and NRTL Gibbs excess energy (G^{ex}) model (SRK-HV-NRTL)
21 was used by Pedersen et al. [67] and Austegard et al. [68]. While Pedersen et al. applied this
22 model to water – reservoir hydrocarbons equilibrium at high temperature (308.15 – 473.15
23 K) and pressures (70 – 100 MPa), Austegard et al. compared the SRK-HV-NRTL model with
24 the simplified Cubic plus Association model proposed by Kontogeorgis et al. [69,70]. The
25 latter used predictions for mutual solubilities between water, CO₂ and CH₄ and concluded
26 that the SRK-HV-NRTL model can produce acceptable results, while sCPA has been found
27 less accurate. Also, the Peng-Robinson coupled with the Wong-Sandler approach with NRTL
28 (PR-WS-NRTL) was tested by Valtz et al. [71] and Yang et al. [72]. High deviations between
29 CO₂ solubility and PR-WS-NRTL results were observed by Valtz and co-workers, particularly
30 at pressures above 8 MPa. Yang et al. (2019) have found compatible predictions from sCPA
31 and SRK-HV-NRTL models for gas condensate mixtures with water. Gernert and Span [73]

1 extended the original GERG-2008 model for humid and CO₂-rich gases. Later, a new version
2 extended to combustion gases (EoS-CG) was reported as accurate for VLE predictions for
3 mixtures including water, CO₂ and CH₄, covering a wide temperature and pressure range
4 [74].

5 Recently, a broader comparative study for CO₂ – water was performed by Aasen et al.
6 [75]. The authors have fitted and tested predictions from traditional Peng-Robinson (PR)
7 and Soave-Redlich-Kwong (SRK), using Twu et al. [76] alpha function, EoS/G^{ex} models (PR
8 and SRK with HV[77]/WS[78] using classic or TWU- α functions), equations that include
9 association terms (sCPA [69,70] , sCPA-PR [79] and PC-SAFT [80], all of them using different
10 association schemes for water), multiparametric EoS (GERG-2008 [81] and EoS-CG [73,74])
11 and predictive models (PR-UMR [82] and VTPR [83]). An extensive data selection and
12 evaluation was carried out and only accepted measurements between 273 – 478 K and at
13 pressures below 61 MPa were used. According to their findings, at least three fitting
14 parameters are required to represent the binary mixture within an accuracy of 10%.
15 Moreover, the PR/HV with the Twu alpha function and volume shift correction was reported
16 as being the most accurate model considering phase compositions and densities [75]. More
17 recently, Yang et al. [84] used PR and the Wong-Sandler (WS) mixing rules with NRTL to deal
18 with CO₂/water and oil in compositional flooding simulations. Despite the good results, a
19 very limited range of pressures and temperatures were studied.

Table 1. Data published for CH₄ – CO₂ – water.

Temperature (K)		Pressure (MPa)		CO ₂ % mole range (dry basis)		Equilibrium Conditions	Type of Data			Reference
min	max	min	max	min	max		Water content	Solubility Data	Gas in Hydrate Phase	
293.15	313.15	3	6	10	50	VL _w E	yes	no	no	[3]
	344.15	10	100	17.4	94.13	VL _w E	no	yes	No	[4]
324	376.2	10.5	50.6	34	62	VL _w E	yes ^(a)	yes	No	[11]
288.71	323.15	5.559	7.517	94.69		VL _w E Three Phase Locus	yes	no	no	[12]
243.1	288.4	0.11	6.05	20	70	VL _w E	yes ^(b)	no	no	[13]
304.26	473.15	3.45	103.42	10	70	VL _w E	yes	no	no	[14]
						VL _w LCO ₂ E				
285.15	300.5	4.963	20	49	99	VL _w E Four Phase Locus	yes	yes	no	[15]
						L _w LCO ₂ E				
273.6	284.2	1.51	7.19	22.5	76.1	HL _w V	no	yes	yes	[16]
283.32	285.76	4.412	7.251	79.74	100	Four Phase Locus	no	no	yes	[19]
	280.3	3.04	5.46	0	100	HL _w V	no	yes	no	[10]
274.02	280.05	1.66	4.03	28.3	61.2	HL _w V	no	yes	no	[18]
						HV				
273.16	283.26	1.5	5	0	100	HL _w V Four Phase Locus	no	no	yes	[17]
283.09	287.04	4.46	8.37	78	100	Four Phase Locus	no	no	no	[5]
	323.15	10		19	93	VL _w E	no	yes	no	[8]
310.23	344.67	6.99	13.89	11.32	20.22	VL _w E	yes	no	no	[7]
285.11	288.39	7.17	27.71	84.6	90	HL _w LCO ₂	no	yes	no	[9]

(a) Data calculated by model.

(b) Reported as dew point temperatures

1 No similar comparative study has been published for water content predictions for
2 CH₄+CO₂ systems. Predictions from PR-CPA [85], sCPA [85,86], PR and SRK with asymmetric
3 mixing rules [87], PC-SAFT [88,89], SAFT-VR [15,86], SAFT- γ Mie group contribution[90],
4 Peng-Robinson-Stryjek-Vera (PRSV) with WS mixing rules and NRTL[91,92] have been
5 individually analysed. Al Ghafri et al. [15] has found that the SAFT-VR model presented by
6 Míguez et al. [93] strongly underestimated solubilities in aqueous phase while water
7 content in vapour phase is overpredicted. Some models were only evaluated for
8 temperatures above 273.15 K, for instance, the PRSV-WS-NRTL with a linear composition
9 dependent function presented by Zhao and Lvov [91,92].

10 Moreover, in the case of water content in CO₂-rich multicomponent mixtures, the
11 capabilities of different models to predict phase changes based on parameters fitted from
12 binary data are rarely mentioned and hardly discussed. Perhaps partially because of the
13 absence of published data, partially due to a natural tendency to oversimplify such mixtures
14 as pure carbon dioxide or, at best, carbon dioxide – methane. In a previous study, Chapoy et
15 al. [94] have highlighted some problems with such approaches.

16 In the present paper, an experimental and modelling investigation into CH₄ - CO₂ and
17 multicomponent CO₂-rich mixtures in equilibrium with hydrates or liquid water is described.
18 Equations of state representing the most promising approaches to calculate water content
19 in natural gas components were evaluated. It includes association theory (sCPA), G^{ex}/EoS
20 mixing rules (SRK-HV-NRTL) and highly accurate multiparametric (EoS-CG GERG version)
21 thermodynamic models. A parametrization procedure that included data in hydrate region
22 was used to obtain new fitted parameters for SRK-HV-NRTL. Model predictions were
23 compared with experimental water content data measured for CO₂ + CH₄ (25/75, 50/50 and
24 75/25 initial ratio) and multicomponent mixtures (containing at least 38.65% of CO₂, light
25 hydrocarbons, up to i-C₅, and permanent gases, which included nitrogen, oxygen, argon and
26 hydrogen) carried out at temperatures between 233.15 and 288.15 K and pressures up to
27 15 MPa. Measurements in the two phase-region in the presence of hydrates were also
28 conducted and complete compositions are presented for the liquid and vapour phases.
29 Water content predictions in both phases were also compared with model predictions.

30

31

2. Experimental Methods

2.1. Materials

Carbon dioxide and methane used in these experiments were 99.99% pure, supplied by BOC (Table 2). Methane/Carbon Dioxide mixtures (1:3, 1:1 and 3:1) were gravimetrically prepared from pure components.

Table 2. Composition of the chemical used in this work

Chemical	Symbol	CASRN	Purity	Supplier
Carbon dioxide	CO ₂	124-38-9	99.99 vol%	BOC
Methane	CH ₄	74-82-8	99.99 vol%	BOC

All the CO₂-rich synthetic mixtures (referred to as MIX 1, 2, 3 and 4) were prepared by BOC and their compositions are given in Table 3. De-ionized water was used in all tests.

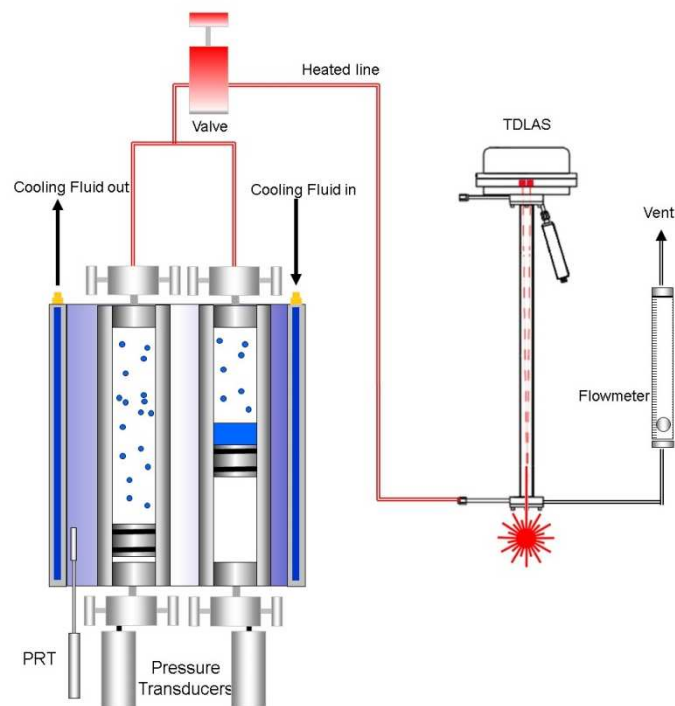
Table 3. Composition, mole% each component, of the multicomponent mixtures used in this work. Uncertainties are given in brackets.

Component	MIX 1	MIX 2	MIX 3	MIX 4
CO ₂	Balance (95.36)	Balance (38.65)	Balance (96.07)	Balance (69.3)
Methane	-	41.30 (±2.06)	-	26.2 (±0.5)
Ethane	-	-	-	0.93(±0.02)
Propane	-	-	-	0.290(±0.006)
n-Butane	-	-	-	0.070(±0.001)
i-Butane	-	-	-	0.070(±0.001)
n-Pentane	-	-	-	0.020(±0.0004)
i-Pentane	-	-	-	0.0300(±0.0006)
Nitrogen	3.0 (±0.1)	20.05 (±1.06)	1.92 (±0.04)	3.08 (±0.06)
Hydrogen	-	-	0.60 (±0.01)	-
Oxygen	10.6 ppm (±0.5)	-	0.83 (±0.02)	-
Argon	1.59 (±0.03)	-	0.58 (±0.01)	-

2.2. Water content measurement set-up

The equipment is comprised of an equilibrium cell and a set-up for measuring the water content of equilibrated fluids flowing out of the cell. A schematic of the set-up is shown in Figure 1. The equilibrium cell is a 300 ml, Titanium piston vessel rated to 69 MPa. The cell is surrounded by a jacket which is connected to a temperature-controlled circulator. The circulator can control the temperature of the fluid pumped through the jacket within ±0.1 K of the set-point and can be used at temperatures between 183.15 and 373.15 K. The

1 cell temperature is measured using a PRT (Platinum Resistance Thermometer) located in the
2 jacket. The cell pressure is measured using a strain gauge pressure transducer mounted on
3 the lower end of the cell. The difference between the temperature probe in the jacket and
4 the temperature inside the cell was checked against a platinum resistance probe that has a
5 certificate of calibration issued in accordance with NAMAS Accreditation Standard and
6 NAMAS Regulations. The pressure transducer is regularly checked for accuracy using a
7 Budenberg dead weight tester.



8

9 **Figure 1. Schematic diagram showing equilibrium cell and water content measurement**
10 **set-up arrangements.**

11

12 The moisture content set-up is comprised of a heated line, a tuneable diode laser
13 absorption spectrometer (TDLAS) and a flow meter. A TDLAS for accurate water content
14 measurements from Yokogawa (Figure 1) was used in this work. The unit is constructed of
15 polished Monel and thus can be used with corrosive gases such as H_2S . The set-up is such
16 that a tuned infrared source is passed through the test sample to a detector. The
17 measurement of water content is based upon true peak area and therefore, unlike other
18 units which use peak height, it is not influenced by changes in background gas. This is
19 subject to the manufacturer specifications regarding the range of concentrations of
20 different gases that will not interfere with the accuracy of the measurements. The unit has

1 two measurement ranges 0-100 ppmV and 0-3000 ppmV, both having a stated accuracy of
2 $\pm 1\%$ of full scale.

3 **2.3. Procedures**

4
5 At the start of a test around 10 ml of 0.1 mm glass beads are placed in a cup shaped
6 depression in the bottom of the piston. 2 ml of deionised water are then mixed with the
7 glass beads. The glass beads have been found to aid in formation and dissociation of
8 hydrates in previous work, helping to achieve equilibrium [95]. The cell is then closed, and
9 the temperature reduced to 263.15 K and evacuated before injecting the fluid. The cell
10 temperature and pressure are then adjusted to achieve the desired test conditions. The cell
11 temperature is then cycled to lower and higher temperatures than the set point over at
12 least 20 hours. This has been confirmed as being sufficient time for equilibrium to be
13 achieved by conducting water content measurements over a number of days in one test.

14 Once equilibrium had been achieved the valve at the top of the cell was opened in
15 order to fill the section of heated line up to the valve prior to the hygrometer at the same
16 time nitrogen was introduced into the base of the cell in order to maintain the pressure
17 constant. Following this, the valve prior (inlet) to the TDLAS was opened sufficiently to
18 achieve a flow rate of between 0.5 and 1 litre per minute through the spectrometer. The
19 water content reading from the hygrometer was then monitored until it was stable for at
20 least 10 minutes. This was then taken as the moisture content of the equilibrated fluid in
21 the cell (i.e., flowing out of the cell). During sampling the heated line was maintained at a
22 temperature of 463.15K. The overall estimated experimental accuracy is 4% of the reading,
23 for water content above 100ppmV, and ± 2 ppmV for values below.

24

25 **2.4. Two-phase region analysis**

26

27 In tests where both liquid (L) and vapour (V) phases were present in equilibrium, a
28 sample of the fluids coming from the test cell was collected in an evacuated cylinder and
29 subsequently analysed using GC. Details of the calibration procedure for the GC are shown
30 below.

31 The Flame Ionisation Detector (FID) was used to detect the hydrocarbons (methane,
32 ethane, propane, i-butane, n-butane, pentanes). For calibration, pure gases are simply

1 injected in the chromatograph via the injector with gas syringes of given volumes: 500- μ l
2 syringe for methane calibration and a 100- μ l syringe for ethane. For heavier hydrocarbons
3 (propane to pentanes), MIX 4 was used for calibration. Calibration curves for the different
4 hydrocarbons are obtained, that is a relationship between the response of the detector and
5 the injected quantity.

6 Comparison between injected quantities and calculated quantities (after adjustment
7 of the parameters of polynomial expressions) allows estimation of the calibration
8 uncertainty, which is in a range of $\pm 0.8\%$ for methane (second order polynomial
9 adjustment), of $\pm 1.2\%$ for ethane (first order polynomial adjustment) and of $\pm 2\%$ for
10 propane to pentanes (first order polynomial adjustment). NB: iso-pentane and n-pentane
11 were calibrated and analysed together as a single component.

12 The Thermal Conductivity Detector (TCD) was used to detect N_2 and CO_2 . For
13 calibration, the same procedure was used, the gases are simply injected in the
14 chromatograph via the injector with gas syringes of given volumes: 500- μ l syringe for CO_2
15 calibration and a 100- μ l syringe for Nitrogen. Comparison between injected quantities and
16 calculated quantities allows estimation of the calibration uncertainty, which is in a range of
17 $\pm 1\%$ for CO_2 (second order polynomial adjustment), of $\pm 0.8\%$ for Nitrogen (first order
18 polynomial adjustment).

19

20 **3. Thermodynamic Modelling**

21

22 A representative range of equation of states relevant to industrial applications were
23 used in this study. The choice included one association theory model (sCPA), one EoS with
24 G^{Ex} mixing rules and one highly accurate multiparametric GERG, extended to combustion
25 gases (EoS-CG).

26

27 **3.1. Simplified Cubic-plus Association SRK (sCPA)**

28

29 A widely used simplified version of Soave-Redlich-Kwong Cubic Plus Association
30 (sCPA), originally presented by Kontogerogis et al.[69,70] was used for this study. The
31 pressure-explicit expression for the model is given by:

1

$$P = \frac{RT}{v-b} - \frac{a}{v(v+b)} - \frac{1}{2} \frac{RT}{v} \left(1 + \rho \frac{\partial \ln g}{\partial \rho} \right) \sum_{i=1}^N x_i \sum_{A_i} (1 - X_{A_i}) \quad (1)$$

2

3 where x_i is mole fraction of the component i and X_{A_i} represents the mole fraction of
4 molecule i not bonded to the site A and expressed as:

5

$$X_{A_i} = \left(1 + \sum_j \sum_{B_j} x_j X^{B_j} \Delta^{A_i B_j} \right)^{-1} \quad (2)$$

6

7 where $\Delta^{A_i B_j}$ represents the association strength between site A on molecule i and site B on
8 molecule j and is defined as:

9

$$\Delta^{A_i B_j} = g(d) \left[\exp\left(\frac{\varepsilon^{AB}}{RT}\right) - 1 \right] b \beta^{A_i B_j} \quad (3)$$

10

11 where ε and β are the association energy and volume, respectively. Parameter values for
12 water have been previously published[95]. The simplified expression of the radial
13 distribution, $g(d)$, adopted by Kontogeorgis et al[70] was used:

$$g = \frac{1}{1 - 0.475 \frac{b}{v}} \quad (4)$$

14

15 In the present work, a four site (4C) association scheme was adopted for water. It
16 considers that hydrogen bonding can occur between the two hydrogen atoms and the two
17 lone pair of electrons in the oxygen atom. A cross-associative approach was adopted to the
18 carbon dioxide molecule.

19 The energy parameter (a) of the SRK-CPA is defined using a classical Soave-type
20 temperature expression:

21

$$a = a_0 [1 + c_1 (1 + \sqrt{T_r})] \quad (5)$$

22

23 where parameters a_0 and c_1 , in the case of associative molecules, were adjusted for single
24 component using vapour pressure data obtained from open literature and previously
25 published[95]. Co-volume is assumed as temperature independent and is also adjusted for
26 single component using saturated liquid volume data for associative molecules. For the non-

1 associative ones, traditional SRK expression for a_i and b_i based on critical point coordinates
2 were used.

3 The extension to mixtures is made using the classical van der Waals mixing rules
4 expressions modified to include binary interaction parameters (BIPs). These were correlated
5 to experimental data. Cross association parameters are obtained from CR-1 combining rules.

6

7 **3.2. Soave-Redlich-Kwong / Huron-Vidal / NRTL model (SRK/HV/NRTL)**

8

9 The Huron-Vidal mixing rules were applied to incorporate the NRTL Gibbs excess
10 energy to the Soave-Redlich-Kwong equation[66] . The starting point is given by:

$$a = b \left[\sum_{i=1}^c x_i \frac{a_i}{b_i} - \frac{G^{ex}_{model}}{q_{eos}} \right] \quad (6)$$

11 where $q_{eos} = \ln 2$ for SRK. The traditional expression is employed to \mathbf{b} :

$$b = \sum \sum x_i x_j \frac{b_{ii} + b_{jj}}{2} \quad (7)$$

12

13 For an improved pure component vapour pressure prediction, Mathias-Copeman-like
14 alpha functions[96] were fitted for water and carbon dioxide, according to Equation 8:

15

$$\alpha = \left[1 + c_1(1 - \sqrt{T_r}) + c_2(1 - \sqrt{T_r})^2 + c_3(1 - \sqrt{T_r})^3 \right] \quad (8)$$

16

17 Values for c_1 , c_2 and c_3 are reproduced in Table 4.

18

19 **Table 4. Parameters for Mathias-Copeman alpha function, Equation 8, used in this work.**

	Carbon Dioxide		Water
	$T_r \leq 1$	$T_r > 1$	
C_1	0.880929	0.880929	1.09442
C_2	-0.879632	0	-0.67481
C_3	3.326455	0	0.691994

20

21 Originally, Huron and Vidal[66] used a version of NRTL using local composition as
22 corrected volume fractions, which leads to the introduction of the co-volume in the
23 calculation of c_{ij} :

$$\frac{G^{ex}}{RT} = \sum_{i=1}^c \frac{\sum_{j=1}^c x_j \tau_{ji} c_{ji}}{\sum_{k=1}^c x_k c_{ji}} \quad (9)$$

1
2 where,

$$c_{ji} = b_j e^{(-\alpha_{ji} \tau_{ji})} \quad (10)$$

3
4 and $\alpha_{ii} = \tau_{ii} = 0$.

5 One of the main advantages of such choice for the g^E model is that an exact reduction
6 to the classical mixing and combining rules can easily be obtained by setting:

$$\alpha_{ji} = 0 \quad (11)$$

$$\tau_{ji} = \frac{q_{eos}}{RT} \left[-2 \frac{\sqrt{b_i b_j}}{b_i + b_j} \sqrt{\frac{a_i a_j}{b_i b_j}} (1 - k_{ij}) \right] \quad (12)$$

9
10 Apart from CO₂/water, CH₄/water and CO₂/CH₄, to which τ_{ij} and τ_{ji} were fitted using
11 data available on the literature, Equations 11 and 12 were applied to recover traditional the
12 SRK model. For this case, binary interaction parameters were taken from Jaubert and co-
13 workers Group Contribution approach [97] which was extended to SRK in [98].

14 15 **3.3. GERG-2008 /EOS-CG**

16
17 The original GERG-2008 wide-range equation of state for natural gases was published
18 by Kunz and Wagner[81] for 21 components including methane and carbon dioxide. The
19 general structure of this multiparametric model is explicitly expressed in term of
20 dimensionless Helmholtz energy ($\alpha = a/RT$) split into ideal (α^o) and residual (α^r)
21 contribution terms:

$$\alpha(\delta, \tau, x) = \alpha^o(\rho, T, x) + \alpha^r(\delta, \tau, x) \quad (13)$$

23

1 for which, reduced mixture density (δ) and inverse reduced mixture temperature (τ) are
 2 defined as:

$$\delta = \frac{\rho}{\rho_r} \quad (14)$$

$$\tau = \frac{T_r}{T} \quad (15)$$

3
 4
 5 Reduced properties ρ_r and T_r are composition dependent and given by, respectively:

$$\frac{1}{\rho_r(x)} = \sum_{i=1}^c x_i^2 \frac{1}{\rho_{c,i}} + \sum_{i=1}^{c-1} \sum_{j=i+1}^c 2x_i x_j \beta_{v,ij} \gamma_{v,ij} \frac{x_i + x_j}{\beta_{v,ij}^2 x_i + x_j} \frac{1}{8} \left(\frac{1}{\rho_{c,i}^{1/3}} + \frac{1}{\rho_{c,j}^{1/3}} \right)^3 \quad (16)$$

$$T_r(x) = \sum_{i=1}^c x_i^2 T_{c,i} + \sum_{i=1}^{c-1} \sum_{j=i+1}^c 2x_i x_j \beta_{T,ij} \gamma_{T,ij} \frac{x_i + x_j}{\beta_{T,ij}^2 x_i + x_j} (T_{c,i} T_{c,j})^{1/2} \quad (17)$$

6
 7
 8
 9 where $\beta_{v,ij}$, $\gamma_{v,ij}$, $\beta_{T,ij}$ and $\gamma_{T,ij}$ are binary parameters adjusted to experimental data and
 10 obey the following relations:

$$\beta_{v,ji} = \frac{1}{\beta_{v,ij}} \quad (18)$$

$$\beta_{T,ji} = \frac{1}{\beta_{T,ij}} \quad (19)$$

$$\gamma_{v,ji} = \gamma_{v,ij} \quad (20)$$

$$\gamma_{T,ji} = \gamma_{T,ij} \quad (21)$$

11
 12
 13
 14
 15 The ideal representation of Helmholtz energy is given in terms of ideal gas mixture:

$$\alpha^o(\rho, T, x) = \sum_i^N x_i [\alpha_{oi}^o(\rho, T) + \ln x_i] \quad (22)$$

16
 17

1 A dimensionless form of the Helmholtz ideal-gas state, $\alpha_{oi}^0(\rho, T)$, is presented by Kunz
 2 and Wagner[81]. Additionally, the residual part is split into a general ideal gas residual
 3 contribution, $\alpha_{oi}^r(\delta, \tau)$, and a binary specific residual function, $\alpha_{ij}^r(\rho, \tau)$, as follows:

$$\alpha^r(\rho, T, x) = \sum_i^N x_i \alpha_{oi}^r(\delta, \tau) + \sum_i^{N+1} \sum_{j=i+1}^N x_i x_j F_{ij} \alpha_{ij}^r(\rho, \tau) \quad (23)$$

4
 5 Binary specific residual functions are normally expressed as a polynomial expression
 6 that also includes exponential terms. Different formats have been introduced, and a
 7 generalized form is presented by Herrig [74]. Any specific function was originally included in
 8 the GERG-2008 formulation for water binary mixtures. Later, Gernert and Span [99]
 9 introduced an extended form for humid gases and carbon capture and storage (CCS)
 10 mixtures. This extension, named EoS/CG, included specific residual functions for CO₂/ water
 11 (validated for 251-623 K and 1 – 350 MPa range for VLE data). Recently, Herrig [74]
 12 developed a binary specific departure function for methane/water. For model tuning, the
 13 author used data for methane solubility, water content and validation were performed in
 14 the temperature range of 250 and 600 K and pressure up to 68.8 MPa. Thus, for the aim of
 15 this work, values for $\beta_{v,ij}$, $\gamma_{v,ij}$, $\beta_{T,ij}$ and $\gamma_{T,ij}$ as well as binary specific functions
 16 expressions were taken from the works of Kunz and Wagner [81], Gernert and Span [99] and
 17 Herrig [74].

18 Hydrate phase and hydrate forming conditions are modelled by the solid solution
 19 theory of van der Waals and Platteeuw, as implemented by Parrish and Prausnitz. A detailed
 20 description of the thermodynamic methodology as well as parameter values was previously
 21 presented [95].

22 **4. Results**

24 **4.1. Adjustment for SRK/HV Model**

25
 26 A new fitting was carried out for SRK-HV-NRTL model, for the temperature interval
 27 between 243.15 and 423.15 K and pressures up to 100 MPa. For this purpose, data for the
 28 binary systems CO₂/CH₄, CO₂/H₂O and CH₄/H₂O from several authors, including LL and VL
 29 including hydrates region, were collected, deemed and selected to avoid off-trend and
 30 inconsistent measurements[20–65]. The lack of reliable data limited the lower temperature

1 to 243.15 K, although, in the following sections, comparisons at 233.15 K are performed.
 2 However, such extrapolations are regarded as an important step for model evaluation, to
 3 investigate its ability to provide reasonable predictions outside fitting temperature and
 4 pressure ranges.

5 For methane and carbon dioxide, the following temperature independent parameter
 6 values were obtained: $\alpha = 0.4779$, $\frac{\tau_{ij}}{RT} = 0.8868$ and $\frac{\tau_{ji}}{RT} = 0.6136$. For the other pairs
 7 involving water, temperature dependent τ_{ij} were fitted and 3rd order polynomial
 8 expressions are reported in Table 5. Parameters for any component not mentioned in Table
 9 5 was described using Equations 11 and 12 to reduce the model to the original SRK cubic
 10 equation.

11

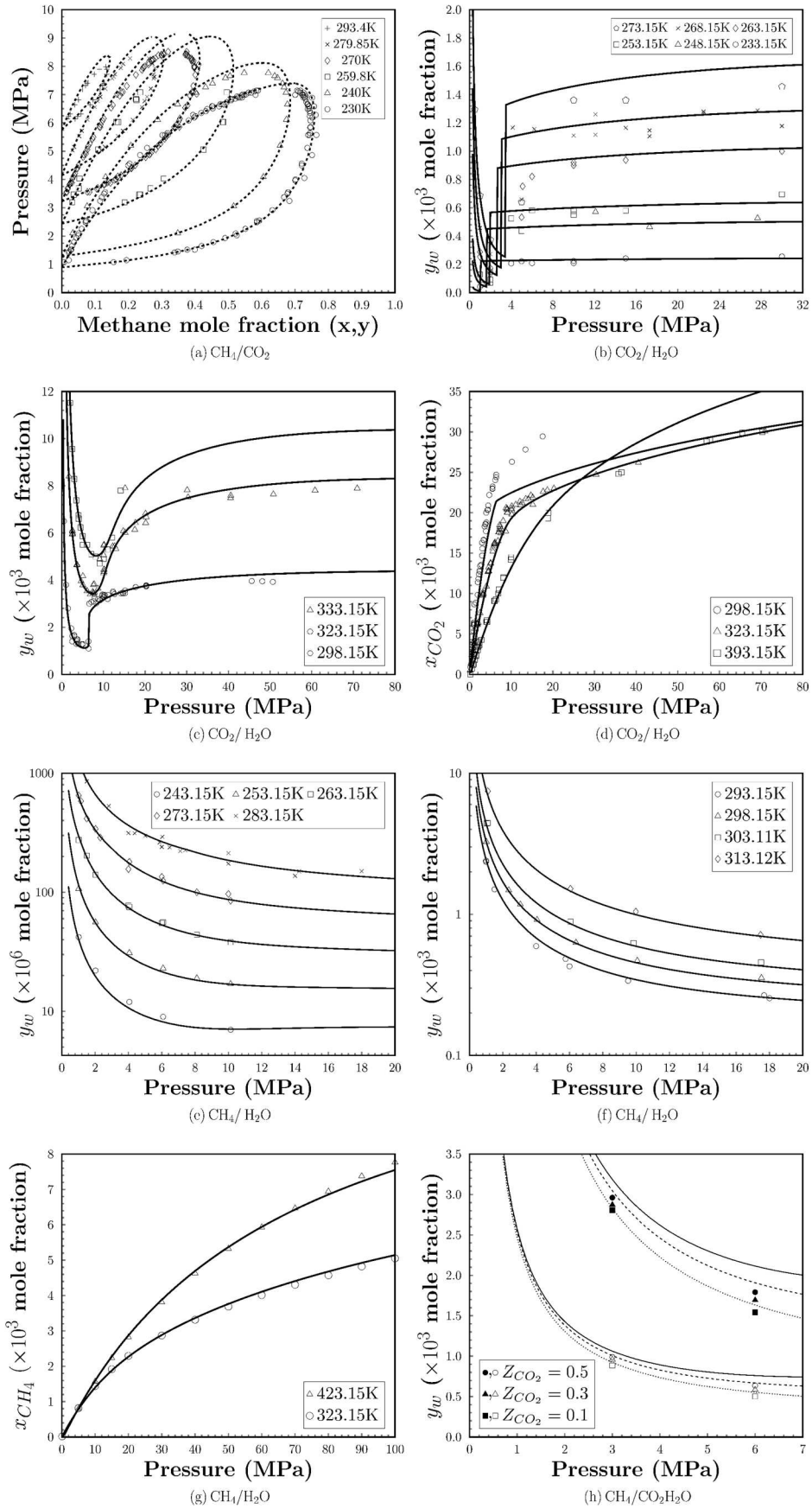
12 **Table 5. Adjusted parameters for SRK-HV-NRTL for water pair with methane or CO₂.**

		Water(j)
		$\alpha = 0.05$
Methane(i)	$\frac{\tau_{ij}}{RT}$	$= -8.171 \cdot 10^{-7}T^3 + 9.994 \cdot 10^{-4}T^2 - 0.4485T + 81.36$
	$\frac{\tau_{ji}}{RT}$	$= 4.861 \cdot 10^{-7}T^3 - 5.656 \cdot 10^{-4}T^2 + 0.2347T - 37.68$
		$\alpha = 0.03$
Carbon Dioxide(i)	$\frac{\tau_{ij}}{RT}$	$= -1.559 \cdot 10^{-6}T^3 + 1.829 \cdot 10^{-3}T^2 - 0.7493T + 118.657$
	$\frac{\tau_{ji}}{RT}$	$= 8.444 \cdot 10^{-7}T^3 - 9.624 \cdot 10^{-4}T^2 + 0.3858T + 62.534$

13

14 The adjusted model was then compared with experimental data, as depicted in Figure
 15 2 a – h. In general, a good agreement between the SRK-HV-NRTL predictions and
 16 experimental data was found for the CH₄/CO₂ binary system (Figure 2a), water content in
 17 liquid and vapour CO₂ (Figure 2b – 2c), CO₂ solubility in water (Figure 2d), water content in
 18 vapour CH₄ (Figure 2f) and CH₄ solubility in water (2g). Figures 2b and 2e show satisfactory
 19 results for fluid phase equilibrium with hydrates. Note that some limitations in dealing with
 20 liquid-liquid CO₂-water equilibrium in the vicinity of carbon dioxide critical point ($T_c =$
 21 304.13K) were also observed.

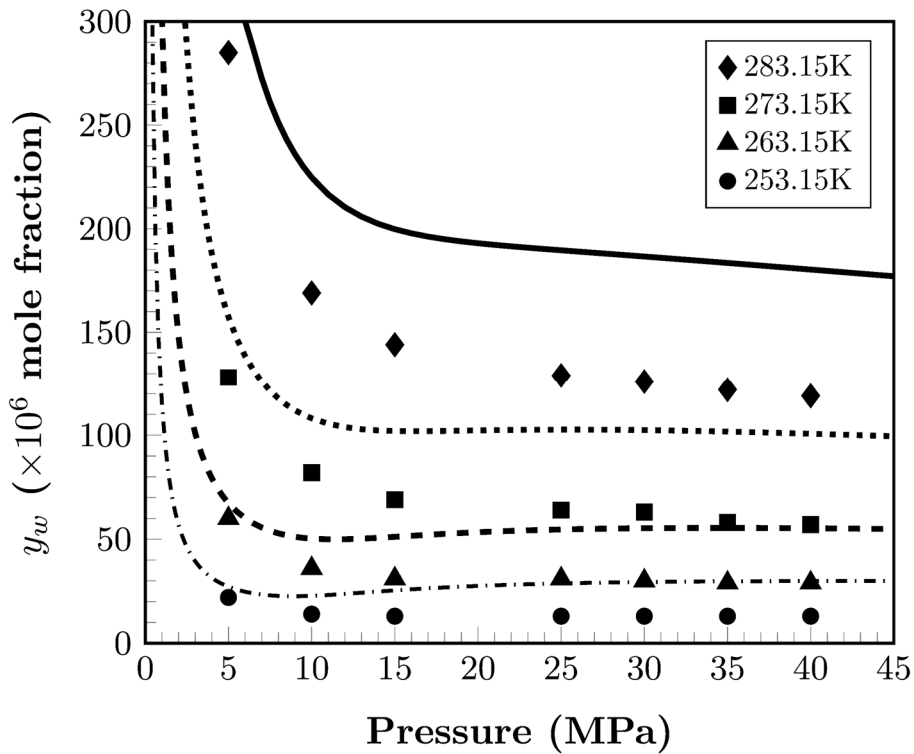
1 Despite the results obtained for binary systems, when the model was extrapolated to
2 predict water content for CO₂ – CH₄ mixtures, the results were unsatisfactory, see Figure 2h.
3 It was particularly unexpected since neither high carbon dioxide concentration ($z_{CO_2} = 10 -$
4 50% mole/mole) nor extreme conditions (3 – 6 MPa and 293.15 and 313.15 K) are used
5 (data from Chapoy et al. [3]).



1

2 **Figure 2. Predictions from adjusted SRK/HV/NRTL model. Data from:(a) [26,43,54,61–65]**
 3 **(b) [21,22,32], (c) [27–31,33–35,55] (d) [15,27,31,33,36–38,40,41,48–53,59,71];(e)**
 4 **[23,24,56], (f) [23,24,56–58] (g) Duan and Mao [60,100] (h) Chapoy et al. [3].**

1 Further evaluations were carried out using a different set of data published by Burgass
 2 et al. [101] for a 9% CO₂/CH₄ mixture, at low temperatures, in equilibrium with hydrates. As
 3 plotted in Figure 3, the model was unable to reproduce water content measurements using
 4 the fitted parameters obtained for the binary corresponding mixtures. In fact, it showed a
 5 clear trend to overestimate values.



6
 7 **Figure 3. Water content predictions for 9% CO₂/Methane mixture at several temperatures.**
 8 **Data from Burgass et al. [101]. For this set of data, AAD = 65%.**

9
 10 One might attribute these limitations to self or cross-associations and the inability of
 11 SRK-HV-NRTL to properly describe them. To clarify this, predictions and experimental data
 12 for a 9% C₂H₆/CH₄ mixture (for which there is no cross-association), at the same conditions,
 13 were compared. Prior to this, a fitting for the ethane – water system was conducted, and
 14 the polynomial expression is shown in Table 6.

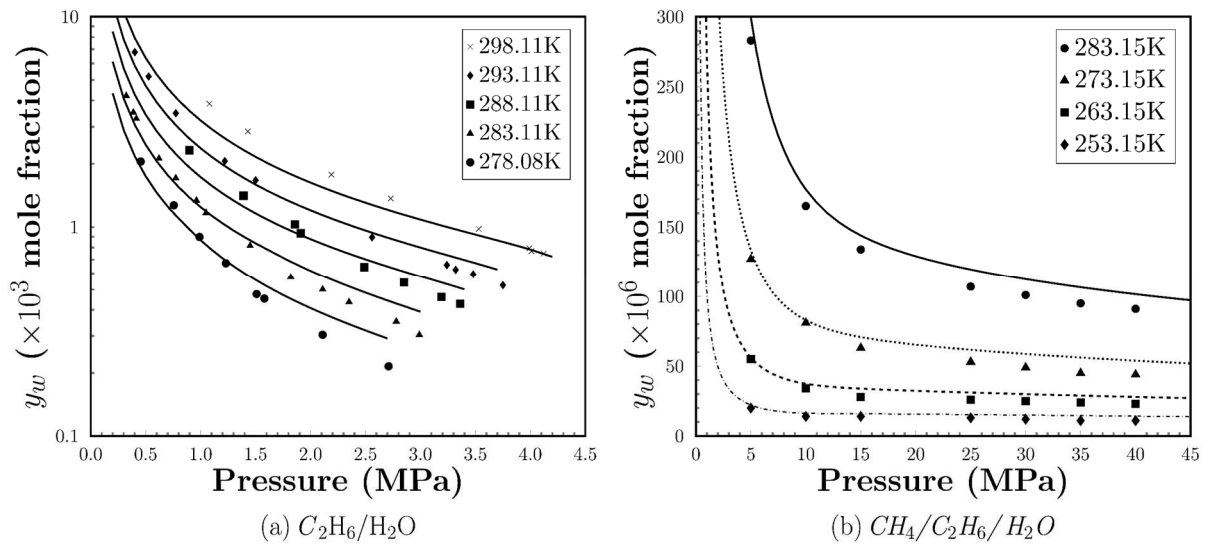
15
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 17
 18
 19

1 **Table 6. Adjusted parameters for Ethane/Water using data from Chapoy et al. [99].**

Water(j)	
$\alpha = 0.05$	
Ethane (i)	$\frac{\tau_{ij}}{RT} = 1.807 \cdot 10^{-4}T^2 - 0.18T + 53.237$
	$\frac{\tau_{ji}}{RT} = -5.717 \cdot 10^{-5}T^2 + 0.06244T - 21.564$

2

3 Water content in ethane was well represented using the parameters from Table 6
 4 (Figure 4a). When extrapolated to the ternary mixture, although over predictions were
 5 observed at higher pressures, (Figure 4b), average deviations were considerably lower (ADD
 6 = 15% against 65% for 9%CO₂/CH₄). It could indicate that Huron-Vidal mixing rules fail to
 7 account for water – carbon dioxide cross-association.



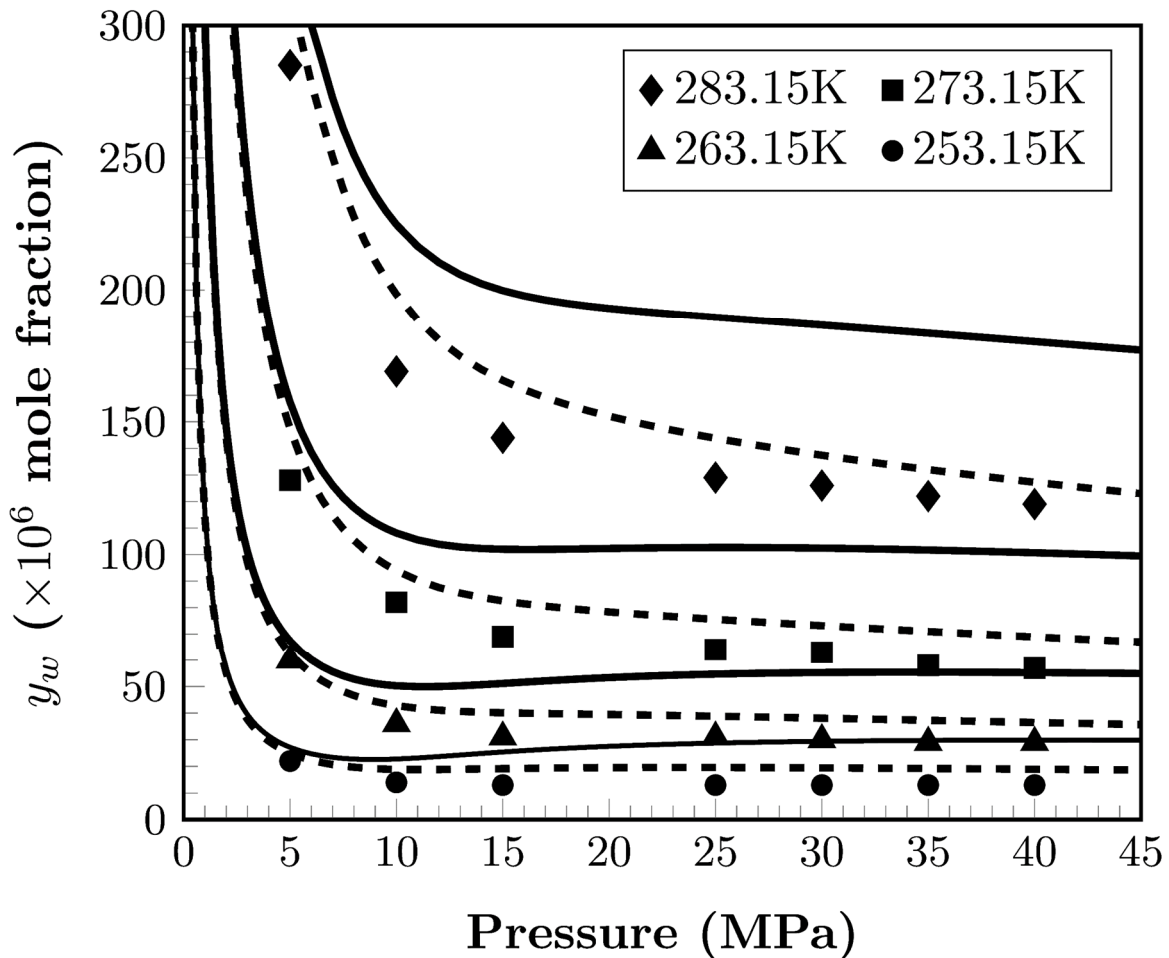
8

9 **Figure 4. Predictions from SRK/HV/NRTL, data from (a) Chapoy et al. [99]. and (b) Burgass**
 10 **et al. [101].**

11

12 At this point, the fact that no adjustment was performed to methane-ethane (instead,
 13 original SRK was used) and yet good predictions were obtained suggested that the same
 14 should be used for methane-carbon dioxide. Thus, as an alternative approach, Equations 11
 15 and 12 were adopted to recover original SRK model for CH₄/CO₂ pair, instead of employing
 16 the fitted α , τ_{ij} and τ_{ji} . Contrary to expectations, this shift gave better predictions, as seen
 17 in Figure 5. As can be seen the average absolute deviation (AAD) reduced very significantly
 18 from 65% to 23%. This is a considerable improvement and indicates that using Huron-Vidal

1 approach only to binary involving water is enough to guarantee the best SRK-HV-NRTL
 2 performance. Therefore, this approach was adopted for the remaining sections of this work.



3
 4 **Figure 5. Comparison between predictions from two different SRK-HV-NRTL approaches:**
 5 **(a) Solid lines represent those obtained using $\alpha = 0.4779$, $\frac{\tau_{ij}}{RT} = 0.8868$ and $\frac{\tau_{ji}}{RT} =$**
 6 **0.6136 for carbon dioxide – methane; (b) Dashed lines represent those obtained using**
 7 **Equations (11) and (12) approach to carbon dioxide – methane. Data from Burgass et al.**
 8 **[101].**

9
 10 **4.2. Water content measurements for methane/carbon dioxide mixtures**
 11

12 New measurements for water content in methane/carbon dioxide mixtures were
 13 carried out at temperatures between 233.15 and 288.15 K at 15 MPa. In this work,
 14 evaluations were focused in the supercritical and liquid region for which no data are
 15 available. Results are presented in Table 7.

16 All measurements for the 75 mole% CH₄ systems were carried above the critical point
 17 in the supercritical region. As expected, the water content is increasing with the CO₂

1 concentration in the feed gas, reducing with CH₄ content, i.e., 92%, 81 % and 55% average
 2 reduction compared to pure CO₂ in the 0.75 CH₄ + 0.25 CO₂, 0.50 CH₄ +0.50 CO₂ and 0.25
 3 CH₄ + 0.75 CO₂ systems, respectively. For all cases, water content in liquid/supercritical
 4 phases reduces as temperature decreases, as normally observed in the vapour phase, for
 5 instance see Figure 2(f) and 3.

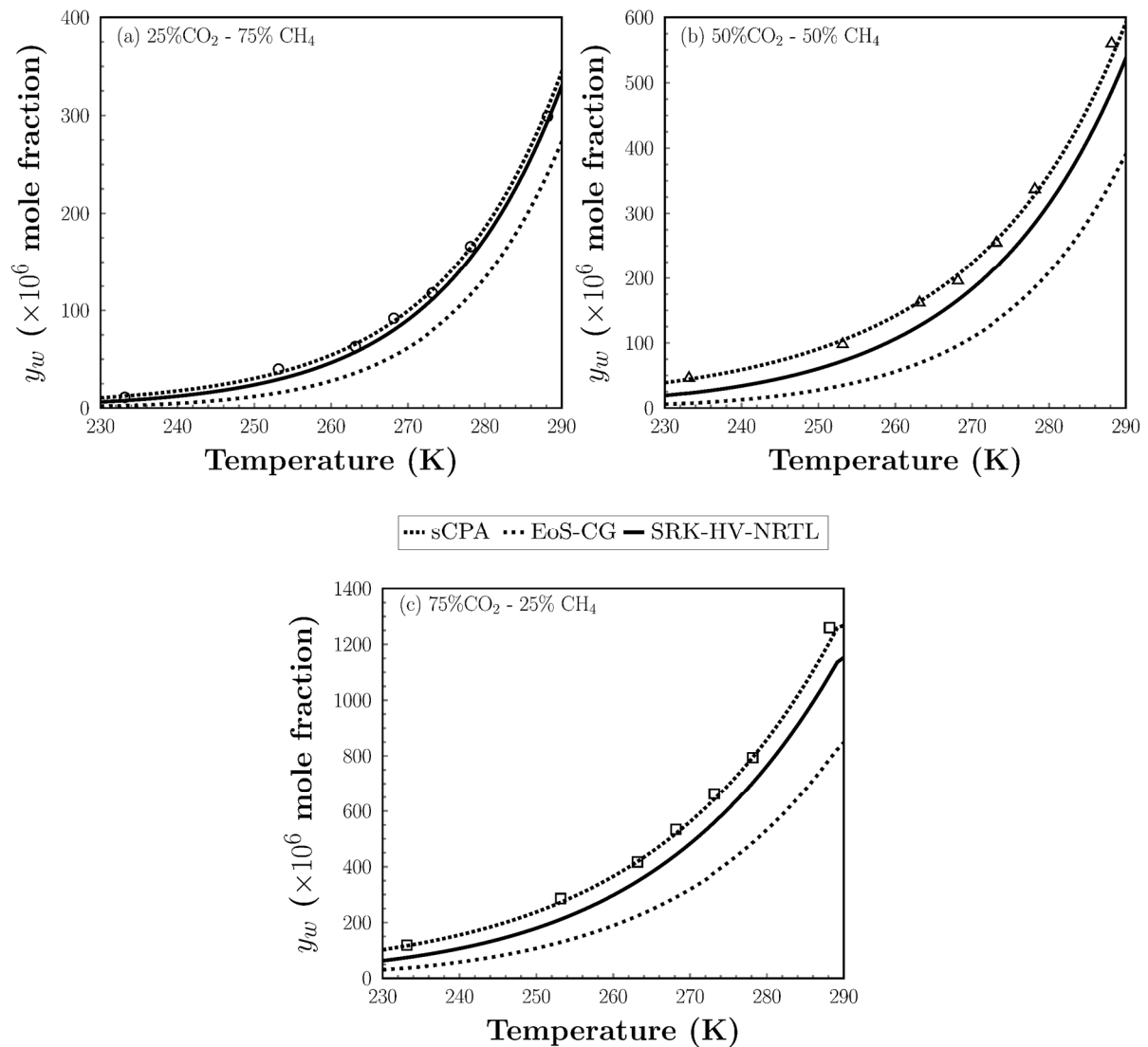
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7 **Table 7. Experimental water content at 15 MPa in the carbon dioxide + methane system in**
 8 **equilibrium with hydrates (L: liquid region; SC: supercritical region).**

T/ K	Water Content / ppm		
	Z _{CO2} =0.25	Z _{CO2} =0.50	Z _{CO2} =0.75
233.15	11 ^{SC}	46 ^L	119 ^L
253.15	40 ^{SC}	98 ^L	286 ^L
263.15	63 ^{SC}	162 ^{SC}	417 ^L
268.15	92 ^{SC}	196 ^{SC}	534 ^L
273.15	118 ^{SC}	255 ^{SC}	663 ^L
278.15	166 ^{SC}	337 ^{SC}	794 ^L
288.15	299 ^{SC}	560 ^{SC}	1260 ^{SC}

9

10 Comparison between sCPA, SRK-HV-NRTL and EoS-CG predictions and the
 11 measurements from Table 7 are depicted in Figure 6 (a)-(c). It is evident that sCPA provides
 12 the better representation for water content in liquid and supercritical region. A considerable
 13 variation was observed, in terms of absolute average deviations (AAD), between the
 14 different equations of state (4, 18 and 49% to sCPA, SRK-HV-NRTL and EoS-CG, respectively).



1

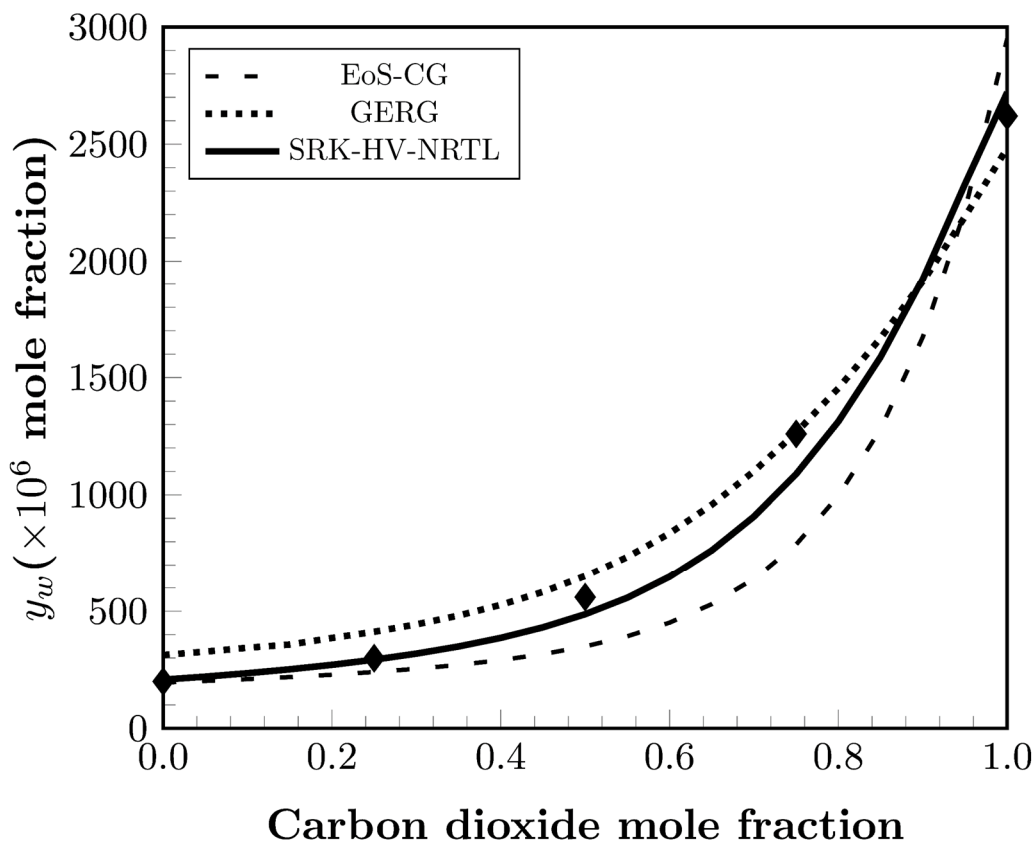
2 **Figure 6. Predictions for water content in carbon dioxide – methane mixtures with**
 3 **different compositions.**

4

5 It is somewhat surprising that the multiparametric approach yields such poor water
 6 content predictions (with AAD = 49%), see dotted lines in Figure 6a-c, although it is regarded
 7 as highly accurate in describing natural and combustion gas compounds. Aasen and co-
 8 workers also reported lower AAD for a PR-HV-NRTL, in comparison with EoS-CG, in their
 9 investigations for CO₂-water mutual solubilities. The authors have attributed this
 10 unexpected output to the fact that the multiparametric model was not fitted directly to the
 11 selected data used in their work. Moreover, they highlighted that this type of model is
 12 normally fitted to more properties than just phase compositions [75].

13 Given the above, the original GERG-2008 (which includes a binary specific departure
 14 function only for carbon dioxide - methane) and EoS-CG predictions were compared, as

1 given in Figure 7, as a function of initial carbon dioxide concentration. It demonstrated that
 2 the combustion gas version only gives good results when $z_{CO_2} \rightarrow 0$. In fact, considering the
 3 overall CO_2 -mole fraction range, GERG-2008 provided better average absolute deviation
 4 (23% against 31%). Since EoS-CG includes specific departure functions for water – methane
 5 and water – carbon dioxide, this observation was unexpected. This suggests that this
 6 multiparametric approach might have limitations in either the specific residual function for
 7 carbon dioxide-water or in the mixing rules and extrapolation procedure to multicomponent
 8 mixtures.



9

10 **Figure 7. Effect of carbon dioxide content in the water content in the vapour phase at 15**
 11 **MPa and 288.15K. Measurement for pure carbon dioxide were taken from [72].**
 12 **Estimation for pure methane is taken from sCPA which fits very well measurements at**
 13 **288.11K published in [67].**

14

15 In contrast, SRK – HV – NRTL results are in good agreement with experimental data,
 16 although, adherence is better at $z_{CO_2} \leq 0.25$. It might be caused by the limitations
 17 described in the adjustment for carbon dioxide – water pair in the vicinity of critical point
 18 and supercritical region, see Figure 2d. Regardless the use of only two parameters and the
 19 absence of a specific approach to associative molecules, its overall performance can be

1 considered satisfactory, particularly for those conditions where the model was validated
 2 (see previous section).

3

4 **4.3. Water Content in CO₂-rich mixtures**

5

6 Measurements have been made for mixtures 1, 2, 3 and 4, for which compositions are
 7 given in Table 2, in equilibrium with water or hydrates at a range of temperatures between
 8 233.15 to 288.15 K, at pressures up to 15 MPa. The measurements at lower pressures were
 9 made at conditions avoiding coming close to the two-phase region (see Figure 8 and 10 for
 10 experimental conditions), except for the measurements undertaken for MIX 4 (see Figure
 11 14).

12 Table 8 summarises average absolute deviations for each mixture individually. sCPA
 13 shows better overall predictions, although SRK-HV-NRTL presented better performance for
 14 MIX 2 and 4. Each mixture is briefly discussed in the following sections.

15

16 **Table 8. Average absolute deviations (ADD) for the models evaluated accounting for each**
 17 **specific mixture. *correspond to the data measured in the liquid vapour region, according**
 18 **to Table 11, for which deviations in the vapour phase at 233.15 K were not considered.**

	MIX 1	MIX 2	MIX 3	MIX 4	MIX 4 LV(*)	Average
sCPA	6.0%	10.2%	6.9%	11.8%	12.0%	9.4%
SRK-HV-NRTL	24.2%	4.9%	26.7%	4.9%	13.2%	15.4%
EoS-CG	19.8%	18.4%	20.3%	38.7%	21.6%	23.7%

19

20 **Mixture 1 (MIX 1):** Results for water content in MIX 1 at different temperature and
 21 pressures are presented in Table 9. Measurements were carried out in the presence of
 22 liquid water or hydrates; T and P condition coordinates are plotted in Figure 8. In addition,
 23 Figure 9 compares model predictions and experimental data.

24

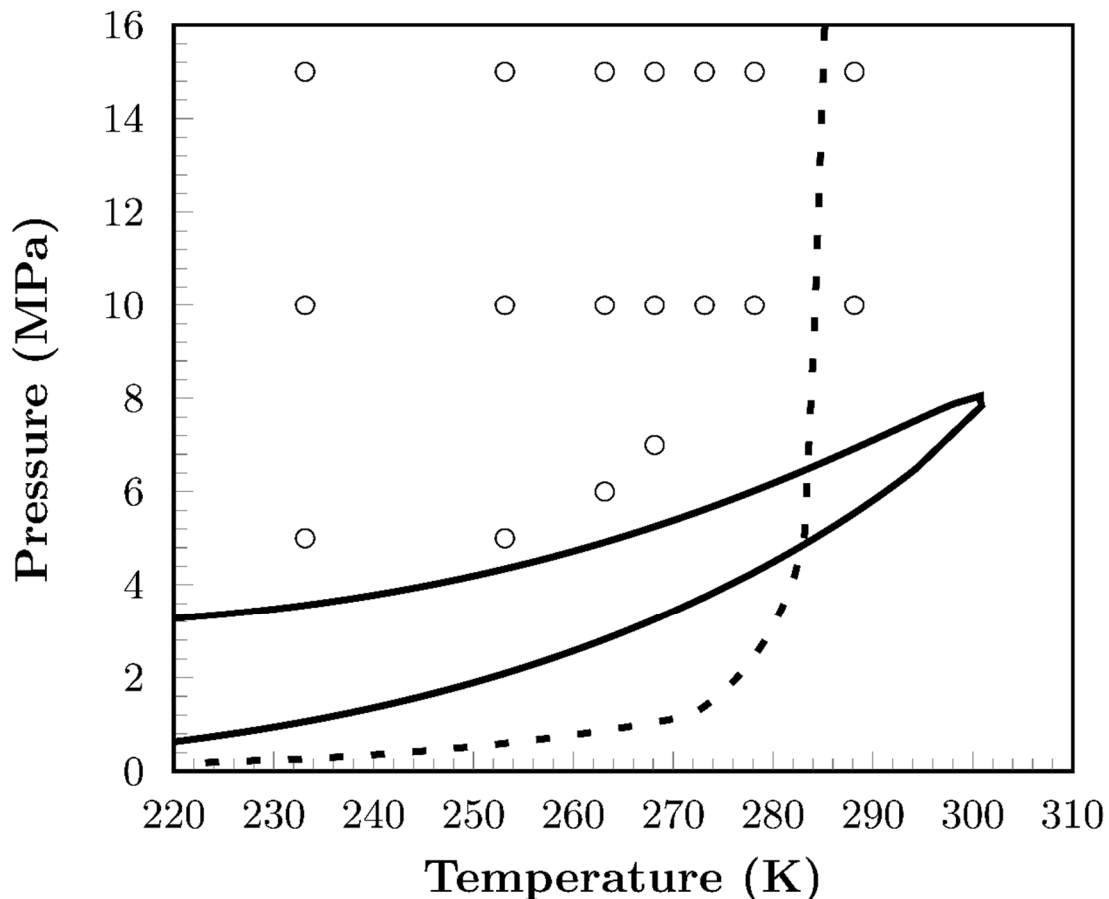
25 **Table 9. Water contents (10⁶ mole fraction) for MIX 1 in equilibrium with hydrates or**
 26 **water.**

Pressure (MPa)	288.15 K	278.15 K	273.15 K	268.15 K	263.15 K	253.15 K	233.15 K
15	2082	1438	1179	950	763	486	177
10	1945	1388	1134	918	745	477	178
7	-	-	-	882	-	-	-

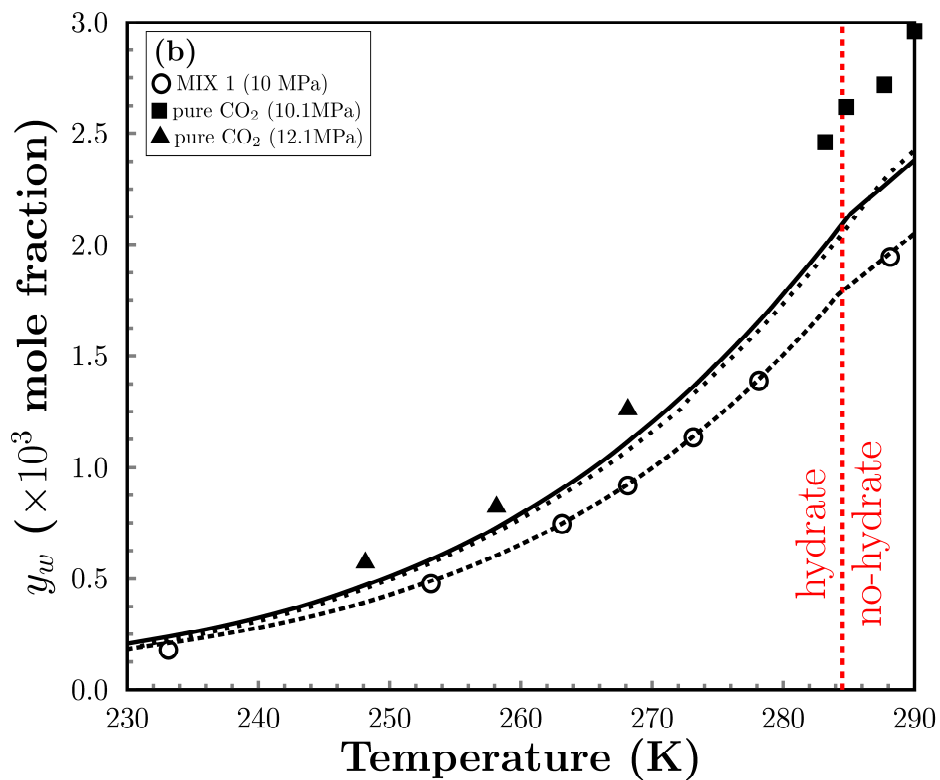
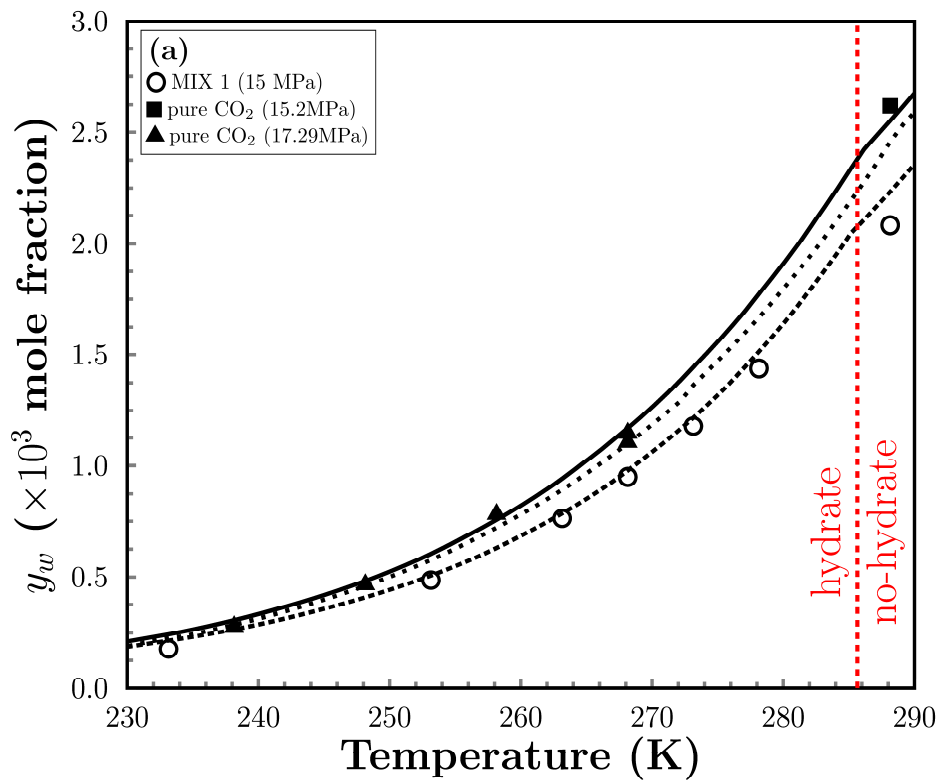
6	-	-	-	-	715	-	-
5	-	-	-	-	-	464	177

1 As seen in Figure 9, the water content in MIX 1 is very similar to the behaviour in pure
2 CO₂, however because of the impurities the water content is significantly lower. Consider,
3 for instance, the result at 288.15 K and 15 MPa. When compared with data published by
4 King et al. [55] at very similar conditions, one can conclude that less than 5% impurity
5 content results in an 20% lowering in water concentration in CO₂-rich phase. This finding is
6 in accordance with a previous study [102], and indicates that even a small amount of inert
7 gases might play an important role in water content predictions. Interestingly, these effects
8 have been widely overlooked or even completely ignored, although it might have a
9 considerable impact on dehydration requirements and hydrate prevention (overdesign or
10 underestimation).

11 Comparing the different modelling approaches, sCPA showed superior capabilities in
12 predicting water content with 6% average deviation. In contrast, EoS-CG and SRK-HV-NRTL
13 basically presented the same deviations (ADD = 20 and 25%, respectively) and were found
14 less accurate, although they can follow the main trend. In general, all the models exhibited
15 higher deviations as temperature decreases.



1 **Figure 8. Experimental Conditions for Water Content in MIX 1 in Equilibrium with**
 2 **Hydrates. Marks represent coordinates for data in Table 9 and dashed line, hydrate**
 3 **dissociation conditions.**



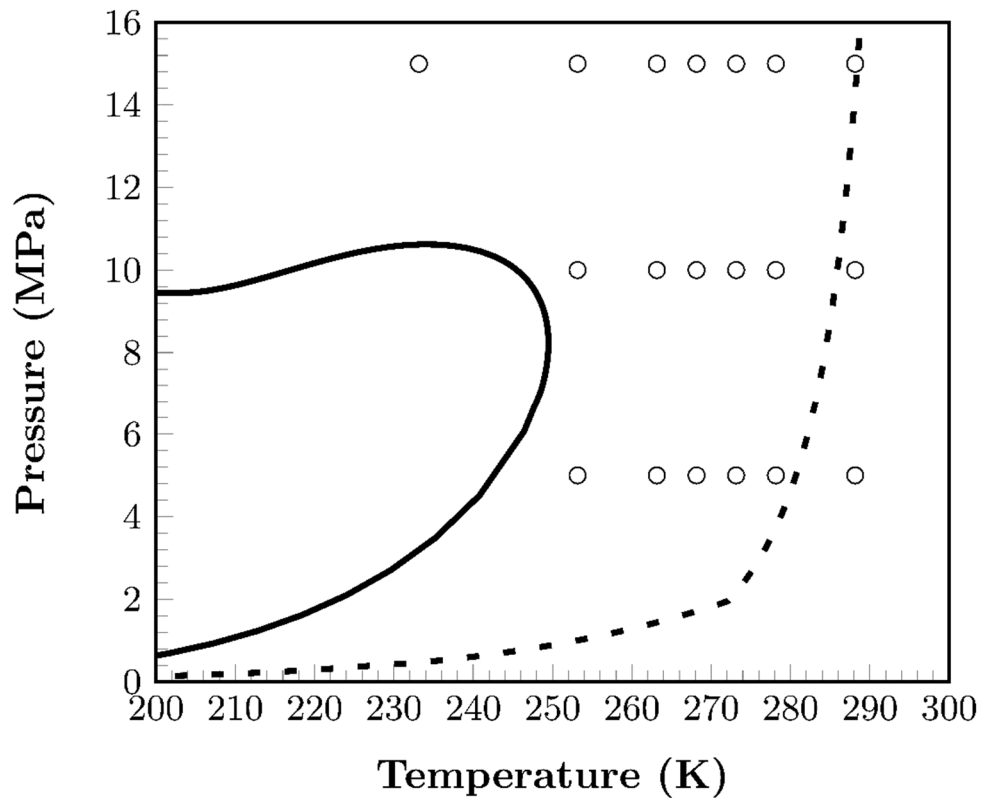
1 **Figure 9. Experimental and predicted water contents (10^6 mole fraction) for pure CO₂ and**
2 **MIX 1 in equilibrium with hydrates or water. Data for pure CO₂ was taken from King et al.**
3 **[72] and Jasperson et al. [63].**

4 **Mixture 2 (MIX 2):** Results for this mixture are presented in Table 9, for which behaviour is
5 more complex, as the water content exhibits a minimum. This characteristic has already
6 been reported for CO₂/methane mixtures by different authors [12,14] and it is attributed to
7 a phase change. Most of the measurements took place in the presence of hydrates, as
8 depicted in Figure 10. Experimental and predicted water content is compared in Figure 11
9 where the effect of pressure is pointed out.

10 In this case, SRK-HV-NRTL yields better predictions, with an AAD = 5%, while 10%
11 average deviation was obtained with sCPA. It is also noted that EoS-CG is unable to predict
12 this minimum in pressure. In fact, above 7 MPa, model outcomes exhibited remarkably poor
13 agreement with experimental data, in an almost ideal gas fashion. In spite of this, average
14 deviation maintained the same magnitude observed for the previous mixture,
15 approximately 20%. It is observed that EoS-CG predictions are considerably worse at higher
16 pressures and low temperatures, while the other equations of state compared in this study
17 showed no clear trend.

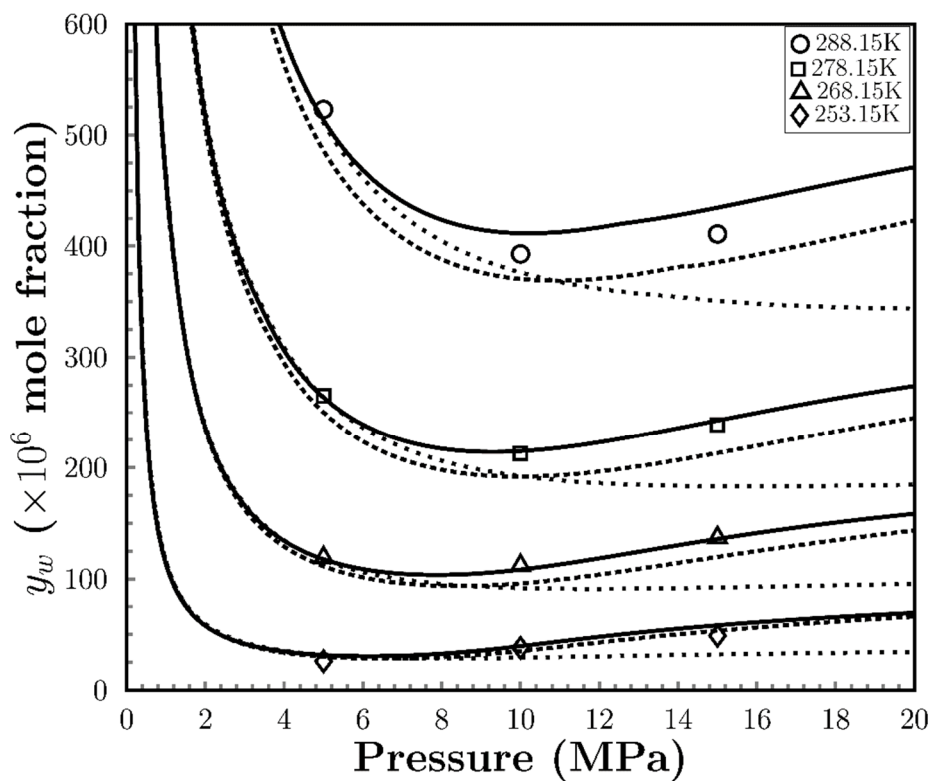
18 **Table 10. Water contents (10^6 mole fraction) for MIX 2 in equilibrium with hydrates or**
19 **water.**

Pressure (MPa)	288.15K	278.15K	273.15K	268.15K	263.15K	253.15K	233.15K
15	411	239	173	137	102	49	18
10	393	213	151	112	78	38	-
5	523	265	184	119	82	26	-



1

2 **Figure 10. Experimental Conditions for Water Content in MIX 2 in Equilibrium with**
 3 **Hydrates. Marks represent coordinates for data in Table 10 and dashed line, hydrate**
 4 **dissociation conditions.**



.....sCPA — SRK-HV-NRTL ... EoS-CG

5

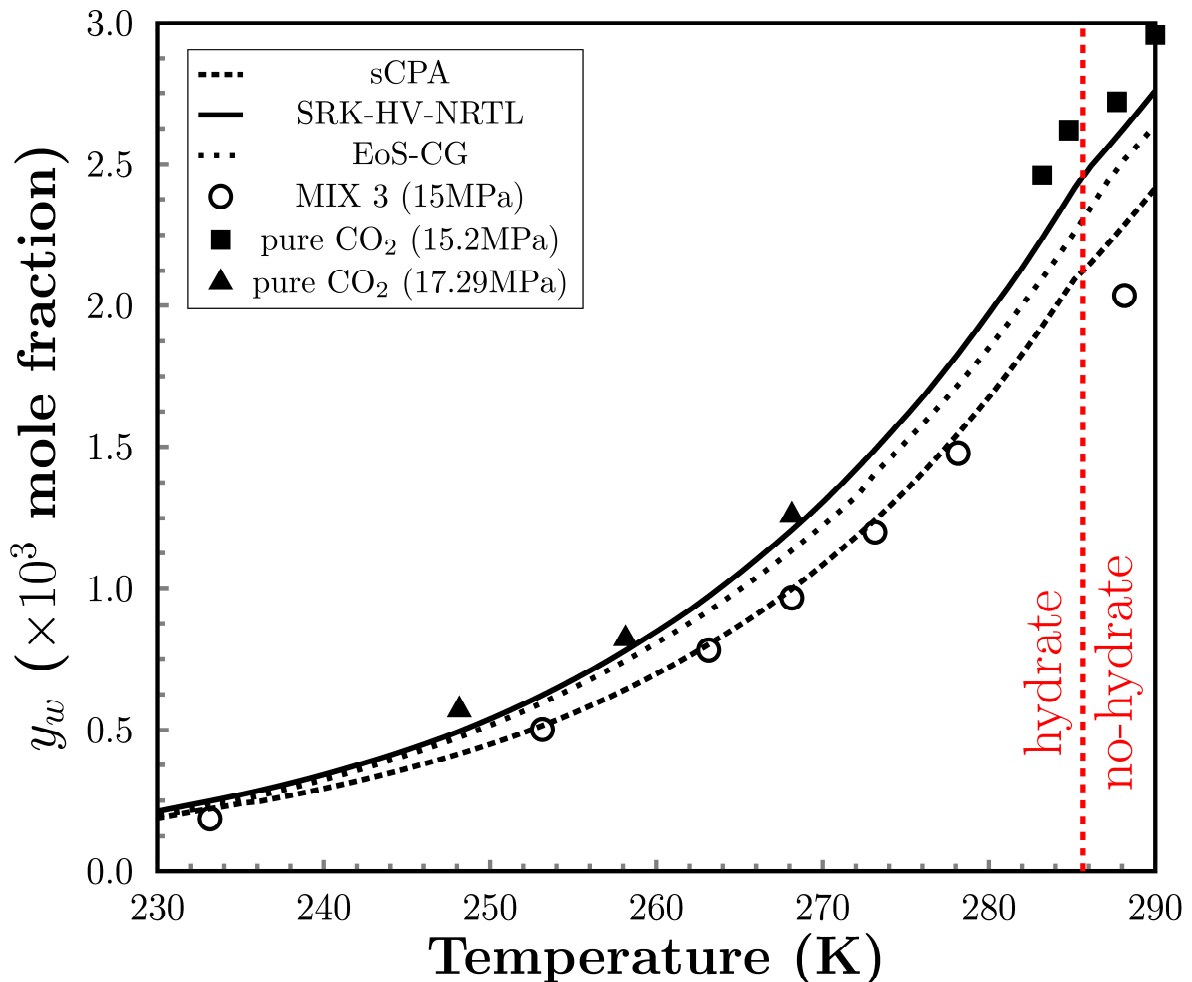
6 **Figure 11. Experimental and predicted water contents (10^6 mole fraction) for MIX 2 in**
 7 **equilibrium with hydrates or water – Pressure effect.**

1 **Mixture 3 (MIX 3):** Measurements were made at 15 MPa and are presented in Table 11 and
 2 compared with predictions in Figure 12. As observed for MIX 1, less than 5% (oxygen,
 3 nitrogen, hydrogen and argon) inert gases considerably reduced water content in
 4 comparison with pure carbon dioxide. Again, at 288.15 K and 15 MPa, this reduction
 5 reached more than 20%. Such results reinforce that small amounts of impurity play an
 6 important role in water content for carbon dioxide rich mixtures.

7
 8 **Table 11. Water contents (10^6 mole fraction) for MIX 3 in equilibrium with hydrates or**
 9 **water***

Pressure (MPa)	288.15K	278.15K	273.15K	268.15K	263.15K	253.15K	233.15K
15	2035*	1478	1199	967	781	502	186

10



11
 12 **Figure 12. Experimental and predicted water contents (10^6 mole fraction) for MIX 3 in**
 13 **equilibrium with hydrates or water. Data for pure CO₂ was taken from King et al. [72] and**
 14 **Jasperson et al. [63].**

15

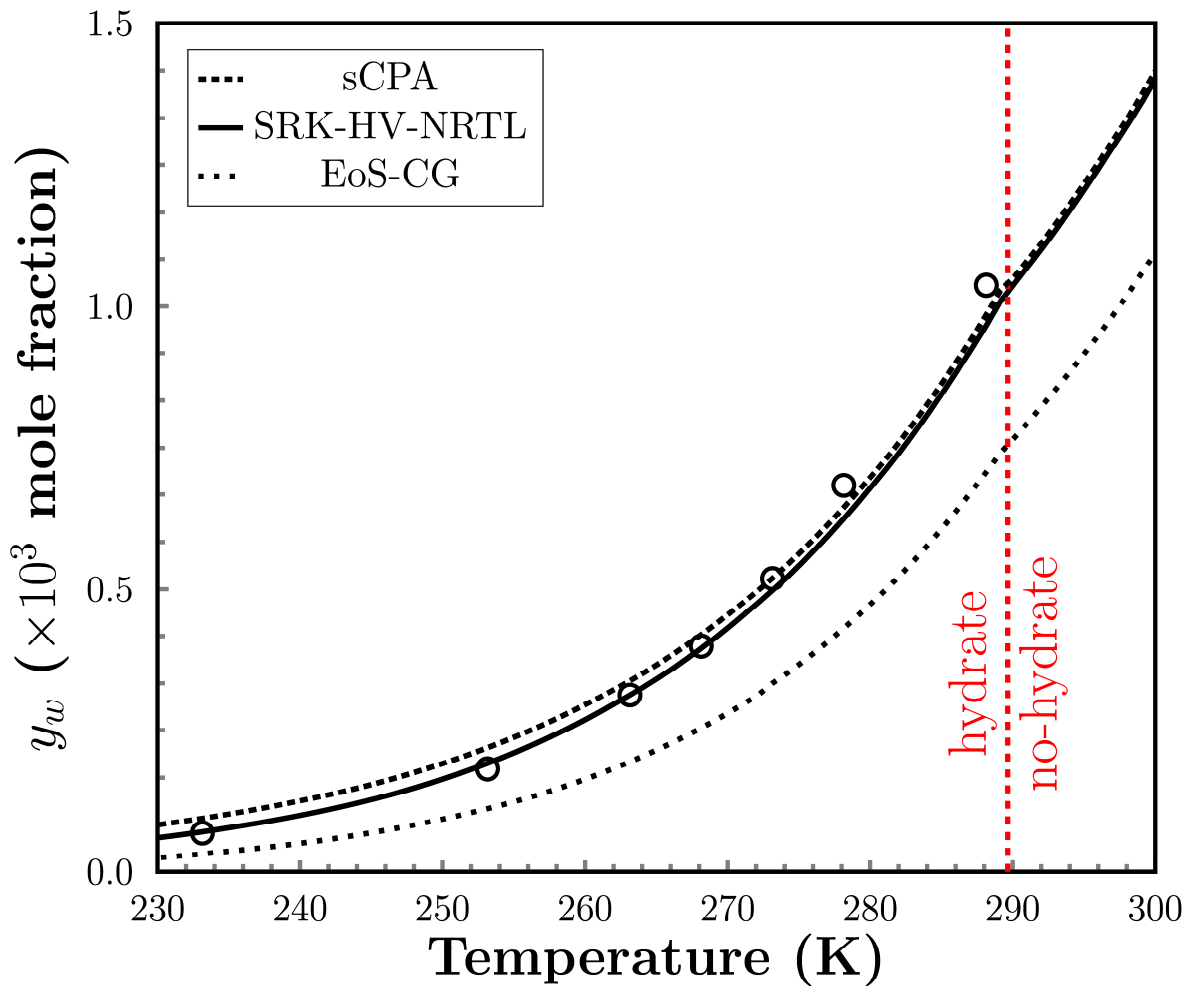
1 Improved predictions for MIX 3 were obtained from sCPA for which ADD was found as
2 low as 7%. Larger deviations were observed for EoS-CG and SRK-HV-NRTL models (20 and
3 26%, respectively). These values are similar to those achieved for MIX 1.

4 **Mixture 4 (MIX 4):** This is a CO₂-rich mixture containing light hydrocarbons, particularly
5 methane, and nitrogen. Results for water content in equilibrium with hydrates or liquid
6 water are shown in Table 12. Experimental data and model predictions are presented in
7 Figure 13. In general, MIX 4 showed the same pattern observed for MIX 1 and 3, regarding
8 the effect of temperature. Moreover, although the mixture has almost 70 % carbon dioxide,
9 reduction in water content, when compared with pure carbon dioxide, corresponds to 60 %
10 at 288.15 K. Again, it highlights the considerable effect of inert gases and hydrocarbons in
11 terms of water content reduction.

12
13 **Table 12. Water contents (10⁶ mole fraction) for MIX 4 in equilibrium with hydrates or**
14 **water***

Pressure (MPa)	288.15K	278.15K	273.15K	268.15K	263.15K	253.15K	233.15K
15	1037	683	519	399	312	182	68

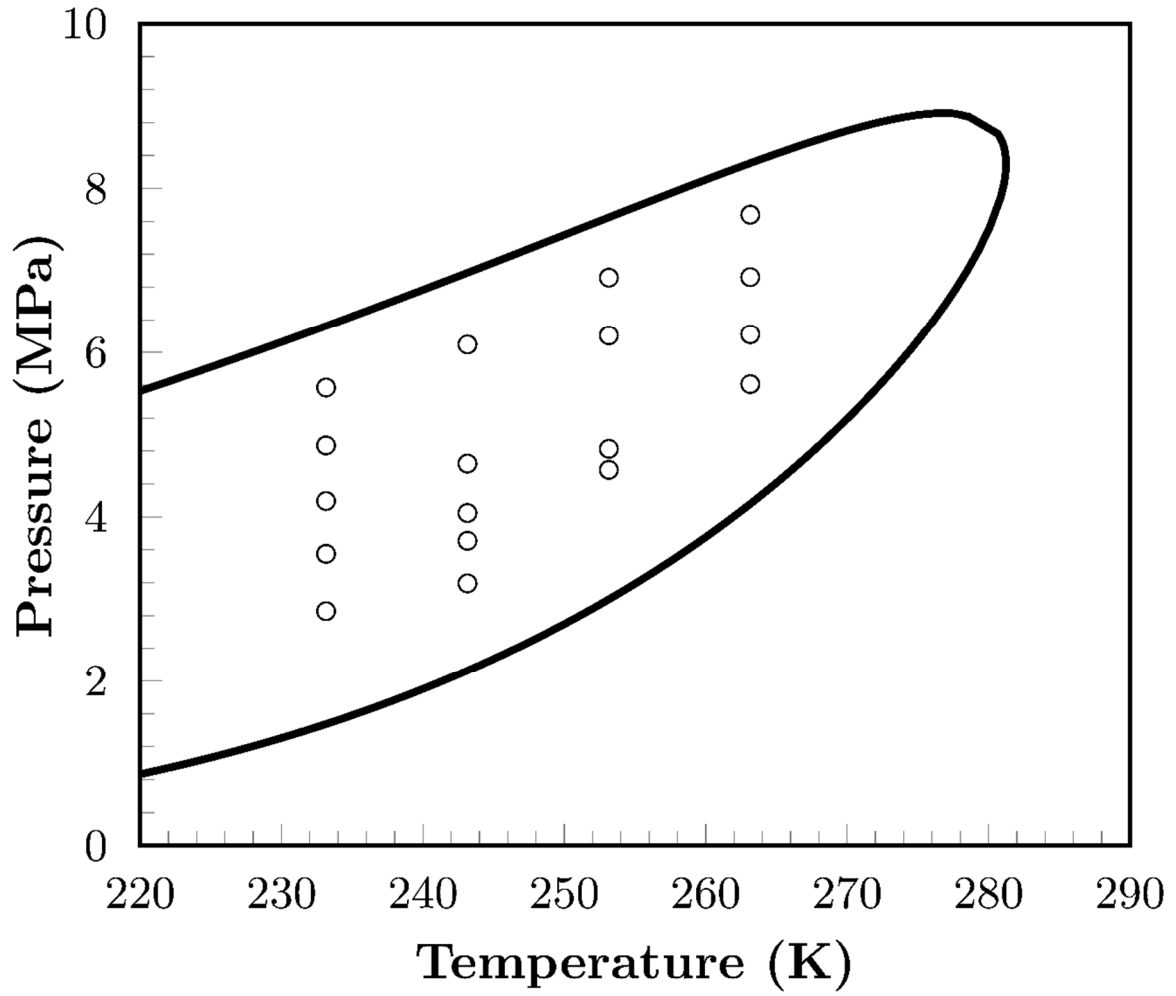
15
16 There was a significant agreement between sCPA and SRK-HV-NRTL predictions and
17 experimental data for MIX 4, see Figure 13. This time, however, the latter showed better
18 results for the whole temperature interval, with ADD = 4.2 %, while the former, 11.8 %.
19 Perhaps the most striking finding was the poor performance of EoS-CG. With the
20 multiparametric model, underestimating predictions with AAD by as much as 38.7 % were
21 observed.



1
2
3 **Figure 13. Experimental and predicted water contents (10^6 mole fraction) for MIX 4 in**
4 **equilibrium with hydrates or water.**

5
6 For MIX 4 an investigation of water content prediction for vapour-liquid equilibrium in
7 the presence of hydrates was undertaken. To do so, 18 measurements were carried out at
8 temperatures between 233.15 and 263.15 K, and moderate pressures (from 2.8 to 7.7 MPa).
9 The experimental conditions are plotted in Figure 14 and phase compositions are listed in
10 Table 13. To obtain these results, a ratio of 5 cc of water for 295 cc of gas was used in each
11 test.

12 As observed in Table 13, experiments conducted in the two-phase region produced a
13 CO_2 -rich liquid phase (with, at least, 77.6% mole/mole), containing between 102 and
14 640 ppmV of water (moles/ 10^6 mole), and a vapour phase containing mainly methane and
15 carbon dioxide and up to 100 ppmV of water vapour.



1

2 **Figure 14. Experimental Conditions for Water Content in MIX 4 in Equilibrium with**
 3 **Hydrates within the 2-phases region. Marks represent coordinates for data in Table 13.**

4

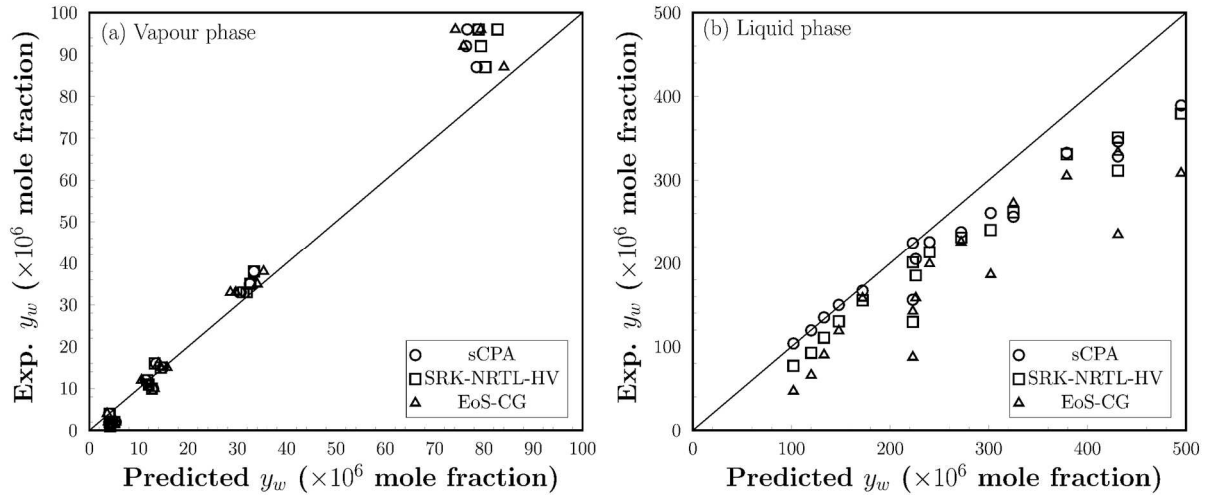
5

Table 13. Water content (Y_w , in ppm, 10^6 mole fraction) and composition of vapour and liquid phases in equilibrium with hydrates for tests with MIX 4. L and V referred to Liquid and Vapour phase, respectively.

	T (K)	P (MPa)	C1	C2	C3	iC4	NC4	iC5	CO ₂	N ₂	nC5	Y _w
L	263.15	5.61	9.5667%	0.9313%	0.4824%	0.1430%	0.1421%	0.0855%	87.7951%	0.7945%	0.0594%	540
L	263.15	6.22	14.9786%	0.9408%	0.4080%	0.1099%	0.1093%	0.0631%	82.0490%	1.2982%	0.0431%	611
L	263.15	6.92	14.4525%	0.9564%	0.4014%	0.1042%	0.1021%	0.0560%	82.5744%	1.3137%	0.0394%	495
L	263.15	7.68	19.3672%	0.9625%	0.3556%	0.0869%	0.0842%	0.0498%	77.5854%	1.4752%	0.0332%	431
L	253.15	4.57	6.8840%	0.9038%	0.4707%	0.1357%	0.1347%	0.0797%	90.9849%	0.3496%	0.0569%	431
L	253.15	4.83	8.5730%	0.8340%	0.4048%	0.1123%	0.1131%	0.0658%	89.2504%	0.5989%	0.0477%	379
L	253.15	6.21	12.3726%	0.8931%	0.3589%	0.0873%	0.0846%	0.0465%	85.2942%	0.8324%	0.0302%	302
L	253.15	6.92	18.5888%	0.9637%	0.3516%	0.0831%	0.0794%	0.0398%	78.4043%	1.4608%	0.0286%	223
L	243.15	3.19	2.3100%	0.7402%	0.4857%	0.1594%	0.1620%	0.1023%	95.7873%	0.1769%	0.0762%	325
L	243.15	3.71	2.9478%	0.7635%	0.4528%	0.1380%	0.1378%	0.0887%	95.2671%	0.1470%	0.0572%	272
L	243.15	4.05	7.0767%	0.8454%	0.3969%	0.1048%	0.1022%	0.0606%	90.9390%	0.4353%	0.0391%	240
L	243.15	4.65	6.7164%	0.8642%	0.4150%	0.1155%	0.1130%	0.0714%	91.1489%	0.5119%	0.0437%	226
L	243.15	6.09	11.4650%	0.8904%	0.3616%	0.0929%	0.0911%	0.0523%	86.4211%	0.5902%	0.0355%	223
L	233.15	2.85	1.5434%	0.8129%	0.4663%	0.1298%	0.1335%	0.0945%	96.6082%	0.1415%	0.0700%	172
L	233.15	3.55	5.0441%	0.9558%	0.4653%	0.1253%	0.1224%	0.0703%	92.9116%	0.2555%	0.0497%	148
L	233.15	4.19	6.4868%	0.9690%	0.4487%	0.1178%	0.1179%	0.0677%	91.3765%	0.3694%	0.0462%	133
L	233.15	4.87	13.3769%	1.0311%	0.4041%	0.0924%	0.0924%	0.0509%	84.2612%	0.6567%	0.0343%	120
L	233.15	5.57	12.1038%	0.9763%	0.3895%	0.0977%	0.0988%	0.0588%	85.6525%	0.5825%	0.0400%	102

V	263.15	5.61	43.875%	0.912%	0.163%	0.020%	0.015%	0.005%	49.701%	5.309%	0.002%	87
V	263.15	6.22	49.418%	0.857%	0.128%	0.016%	0.013%	0.007%	43.658%	5.899%	0.004%	96
V	263.15	6.92	50.217%	0.842%	0.122%	0.013%	0.010%	0.006%	42.410%	6.377%	0.003%	92
V	263.15	7.68	54.373%	0.778%	0.108%	0.012%	0.010%	0.003%	37.780%	6.933%	0.002%	96
V	253.15	4.57	57.824%	0.704%	0.094%	0.011%	0.008%	0.003%	33.397%	7.957%	0.002%	38
V	253.15	4.83	50.077%	0.767%	0.122%	0.015%	0.012%	0.005%	42.475%	6.524%	0.003%	35
V	253.15	6.21	45.183%	0.840%	0.141%	0.017%	0.013%	0.005%	48.212%	5.586%	0.003%	33
V	253.15	6.92	46.163%	0.920%	0.158%	0.019%	0.015%	0.005%	47.627%	5.091%	0.003%	33
V	243.15	3.19	53.049%	0.784%	0.129%	0.018%	0.015%	0.005%	39.087%	6.909%	0.003%	15
V	243.15	3.71	44.626%	0.842%	0.169%	0.026%	0.023%	0.008%	49.033%	5.267%	0.006%	16
V	243.15	4.05	44.884%	0.821%	0.167%	0.026%	0.023%	0.010%	48.376%	5.686%	0.007%	10
V	243.15	4.65	40.028%	0.911%	0.201%	0.032%	0.027%	0.010%	53.811%	4.973%	0.006%	11
V	243.15	6.09	42.233%	0.884%	0.178%	0.026%	0.022%	0.007%	51.688%	4.956%	0.005%	12
V	233.15	2.85	54.481%	0.956%	0.124%	0.012%	0.008%	0.003%	38.253%	6.161%	0.002%	2
V	233.15	3.55	59.642%	0.869%	0.106%	0.010%	0.008%	0.002%	32.379%	6.983%	0.002%	2
V	233.15	4.19	62.415%	0.796%	0.092%	0.009%	0.007%	0.002%	28.992%	7.686%	0.002%	1
V	233.15	4.87	64.098%	0.729%	0.084%	0.008%	0.007%	0.002%	26.603%	8.467%	0.001%	2
V	233.15	5.57	65.117%	0.685%	0.077%	0.008%	0.006%	0.002%	24.914%	9.189%	0.001%	4

1 Figure 15 compares experimental and predicted values for the data displayed in Table
 2 8. Overall, the three different models are predicting the compositions in all phase with the
 3 correct trend of changes. The CPA method showed the best agreement between predictions
 4 and experimental results, particularly in the vapour phase. SRK-HV-NRTL yielded larger
 5 deviations, although, for many conditions, it reproduces sCPA results and tendencies.



6
 7 **Figure 15. Comparative between experimental and predicted water content, data**
 8 **presented in Table 13.**

9
 10 Underprediction was observed for water content in the liquid phase, no matter the
 11 approach used, although, it was more evident for EoS-CG model. Interestingly, sCPA
 12 provided accurate predictions in liquid phase for 233.15 K.

13 **Overall models evaluation:** The present analysis highlights the importance of investigating
 14 capabilities and limitations of equations of state in extrapolating predictions to
 15 multicomponent mixtures. For instance, the accurate multiparametric EoS-CG/GERG
 16 approach showed severe limitations and a massive downgrade in accuracy when used with
 17 methane/carbon dioxide or CO₂-rich multicomponent mixtures with water. It shows how
 18 complex is the task of extending the models based on parameters fitted from binary
 19 systems to complex multicomponent ones. Considering the unexpectedly high average
 20 deviations (50% for the ternary mixture and 27% for MIX 1 to 4), it was also observed that
 21 EoS-CG performed better when dealing with MIX 1 and 3 ($z_{CO_2} > 95\%$), and, when $z_{CO_2} \rightarrow 0$,
 22 in the case of CO₂/CH₄/H₂O (Figure 7). Moreover, the original GERG-2008 (which includes
 23 binary specific departure function only for carbon dioxide - methane) provided better
 24 average absolute deviation (23% against 31%) than EoS-CG (which includes specific

1 departure functions for water – methane and water – carbon dioxide). The most intriguing
2 fact is that this model performs more satisfactorily when there is a major component for
3 which $x_i \rightarrow 1$. Thus, it is possible that, for GERG multiparametric approach, the applied
4 mixing rules are unable to conserve main features from the specific binary functions when
5 extending the approach to multicomponent systems.

6 On the contrary, sCPA has demonstrated a good ability to deal with water containing
7 CO₂-rich systems, within the scope of this study. Although, it exhibited lower accuracy for
8 MIX 2 and MIX 4 and at very low temperatures, its overall predictions can be considered
9 satisfactory.

10 SRK-HV-NRTL showed better results when Huron-Vidal approach was applied only to
11 water – carbon dioxide and water – methane. It suggests that the simpler, the better in
12 incorporating NRTL to SRK. In general, SRK-HV-NRTL model is unable to reproduce the same
13 overall accuracy as sCPA, although, considering the use of only two parameters, it displayed
14 impressive predictive abilities. Particularly for those multicomponent mixtures with initial
15 composition closer to equimolar CO₂/CH₄ (MIX 2 and 4), Huron-Vidal has yielded the best
16 results.

17 In summary, the cubic-plus association version of SRK (previously evaluated for
18 methane-water and carbon dioxide – water binary systems, and that has showed
19 satisfactory water content prediction capabilities [20,32,103–105]) was found to be the the
20 only model, among those considered in this study, capable of satisfactorily predict water
21 content on CO₂-CH₄ and CO₂-rich multicomponent, at temperatures between 233.15 and
22 288.15K and pressures up to 15 MPa, which also include equilibrium conditions with
23 hydrates. It was observed that sCPA was also able to deal with the effect of common
24 impurities, particularly at low concentration, on the water content, as considerable
25 reductions compared with pure carbon dioxide was observed.

26

27 **5. Conclusion**

28

29 An experimental and modelling study of CO₂-rich mixtures with water was performed
30 for temperatures between 233.15 and 288.15 K and pressures up to 15 MPa. This included

1 equilibrium between hydrates or liquid water with liquid or vapour methane/carbon dioxide
2 or CO₂-rich mixtures containing light hydrocarbons (Methane, Ethane, Propane, n- Butane, i-
3 Butane, n-Pentane and I-Pentane) and inert gases (Nitrogen, Hydrogen and Argon). In
4 addition, water content measurements inside the two-phase region in the presence of
5 hydrates were carried out. Complete composition for liquid and vapour phases were
6 reported.

7 Focusing on industrial applications, the capabilities and limitations of three different
8 equation of state approaches (association model theory, Gibbs excess energy model
9 coupling with cubic equations of state and highly accurate multiparametric GERG) were
10 undertaken. Evaluations were focused on water content predictions for CO₂-rich fluid
11 phases. From this work, it can be concluded that:

12

13 • In the application of a SRK-HV-NRTL model fitted from binary systems data, the best
14 results were obtained when Huron-Vidal mixing rules were limited to methane –
15 water and carbon dioxide – water parameters.

16

17 • Small amounts of inert gases in rich-CO₂ streams resulted in a considerable reduction
18 in water content in the fluid phases. It was found that less than 5% permanent gases
19 in initial composition resulted in a 20% water concentration reduction at 288.15 K
20 and 15 MPa. A similar tendency was observed when hydrocarbons are present (92%,
21 81 % and 55% average reduction compared to pure CO₂ in the 0.75 CH₄ + 0.25 CO₂,
22 0.50 CH₄ +0.50 CO₂ and 0.25 CH₄ + 0.75 CO₂ systems, respectively). These results are
23 somewhat counterintuitive and indicate that, although very tempting, approximate
24 CO₂-rich mixtures (even when Z_{CO₂} > 95%) to pure carbon dioxide will lead to
25 considerable water content overestimations (and all the consequent implications
26 such as overdesign, hydrates inhibitors overdosage, etc...).

27

28 • Although regarded as a highly accurate model, EoS-CG/GERG approach has been
29 found very limited when dealing with water plus CO₂/CH₄ or CO₂-rich-mixtures. In

1 fact, no AAD lower than 18% could be found in any studied case. Considering the
2 unexpectedly high average deviations (49% for the ternary mixture and 23.7% for
3 MIX 1 to 4), it was also observed that EoS-CG performed better when dealing with
4 MIX 1 and 3 ($z_{\text{CO}_2} > 95\%$), and, when $z_{\text{CO}_2} \rightarrow 0$, in the case of $\text{CO}_2/\text{CH}_4/\text{H}_2\text{O}$. Thus, it is
5 possible that, for GERG multiparametric approach, the applied mixing rules are
6 unable to conserve the main features from the specific binary functions when
7 extrapolated to multicomponent systems.

- 8
- 9 • For the CO_2 -rich mixtures studied in this work, sCPA exhibited overall better
10 predictions and lower average absolute deviations. For the specific case of MIX 2 and
11 4, SRK-HV-NRTL showed slightly better results. Considering that Huron-Vidal
12 approach used only two parameters, these were considered impressive predictive
13 abilities.
 - 14
 - 15 • In addition, water content measurements were carried out in two phase-region in
16 the presence of hydrates. Composition of liquid and vapour phases were also
17 reported. Despite the overall satisfactory agreement, underpredicted results were
18 observed for water content in the liquid phase, no matter the approach used
19 (although, it was more evident for EoS-CG model).

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23
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28 29 **Abbreviations**

AAD	Average absolute deviation
BIP	Binary Interaction Parameters
CCS	Carbon capture and storage
CR-1	Combining Rule for association parameters
EOR	Enhanced oil recovery
EoS	Equation of State
EoS-CG	Combustion Gas Equation of State
FID	Flame Ionisation Detector
GERG	The European Gas Research Group
HV	Huron-Vidal EoS/ G^{ex} mixing rule
L	Liquid Phase
L_w	Liquid aqueous phase
L_{CO_2}	Liquid carbon dioxide-rich phase
NAMAS	National Measurement Accreditation Service
NRTL	Non-random two-liquid
PC-SAFT	Perturbed Chain Statistical Association Theory
PR	Peng-Robinson
PRSV	Peng–Robinson–Stryjek–Vera
PTR	Platinum Resistance Thermometer
PR-UMR	Peng-Robinson Universal Mixing Rules
SAFT	Statistical Association Fluid Theory
SAFT-VR	Variable Range Statistical Association Fluid Theory
SC	Supercritical
sCPA	Simplified Cubic-Plus Association
SRK	Soave-Redlich-Kwong
TCD	Thermal Conductive Detector
TDLAS	Tunable diode laser absorption spectroscopy
V	Vapour Phase
VLE	Vapor-Liquid Equilibrium
VLLE	Vapor-Liquid-Liquid Equilibrium
VTPR	Volume Translated Peng-Robinson
WAG	Water alternated gas
WS	Wong-Sandler EoS/ G^{ex} mixing rule

1

2 Nomenclature

3

a	Mixture attractive parameter for cubic EoS
a_i	Pure component attraction parameter for Cubic EoS
b	Mixture Co-volume
b_i	Pure component co-volume
c_1, c_2, c_3	coefficients for Mathias-Copeman alpha function
F_{ij}	Binary coefficient for departure functions for GERG/EoS-CG
G^{ex}	Gibbs excess energy
P	Pressure
v	Molar volume
R	Universal gas constant
T	Temperature
g	Radial distribution function (RDF) for sCPA
x_i	Mole fraction of component i
X^{ai}	Fraction of non-bonded components at site A_i
z_i	Initial mole fraction of component i

4

5 Greek Symbols

6

α	Dimensionless Helmholtz energy
----------	--------------------------------

$\alpha(T)$	Alpha function for SRK and PR EoS
α_{ij}	Non-randomness parameter between component i and j in NRTL Gibbs excess model
β^{AiBj}	Association volume parameter between sites Ai and Bj used in sCPA
Δ^{AiBj}	Strength of interaction between sites Ai and Bj used in sCPA
$\beta_{v,ij}$ $\beta_{T,ij}$	Adjustable parameters for GERG and EoS-CG equations
δ	Reduced mixture density used in GERG and EoS-CG models
ε^{ij}	Association energy between sites Ai and Bj used in sCPA model
ρ	Molar density
$\gamma_{v,ij}$ $\gamma_{T,ij}$	Adjustable parameters for GERG and EoS-CG equations
τ	Reduced mixture temperature used in GERG and EoS-CG models
τ_{ij} , τ_{ji}	Parameters for NRTL model

1

2

3 **References**

4

- 5 [1] J.J. Carrol, Natural Gas Hydrates - A Guide for Engineers, Gulf Professional Publishing,
6 2009. <https://doi.org/https://doi.org/10.1016/B978-0-7506-8490-3.X0001-8>.
- 7 [2] A.M.T. De Andrade, C.E.M. Vaz, J. Ribeiro, L.G.R. Lopreato, R.F.S. do Nascimento,
8 Offshore Production Units for Pre-Salt Projects, Offshore Technol. Conf. (2015).
- 9 [3] A. Chapoy, R. Burgass, A. Terrigeol, C. Coquelet, A.T.C.C. Antonin Chapoy Rod Burgass,
10 A. Chapoy, R. Burgass, A. Terrigeol, C. Coquelet, Water Content of CO₂-rich
11 Mixtures: Measurements and Modeling using the Cubic-Plus-Association Equation of
12 State, J. Nat. Gas Eng. 1 (2016) 85–97. <https://doi.org/10.7569/jnge.2015.692505>.
- 13 [4] A. Dhima, J.C. De Hemptinne, J. Jose, Solubility of hydrocarbons and CO₂ mixtures in
14 water under high pressure, Ind. Eng. Chem. Res. 38 (1999) 3144–3161.
15 <https://doi.org/10.1021/ie980768g>.
- 16 [5] Y. Bi, T. Yang, K. Guo, Determination of the upper-quadruple-phase equilibrium region
17 for carbon dioxide and methane mixed gas hydrates, J. Pet. Sci. Eng. (2013).
18 <https://doi.org/10.1016/j.petrol.2012.11.019>.
- 19 [6] P. Kastanidis, G.E. Romanos, A.K. Stubos, I.G. Economou, I.N. Tsimpanogiannis, Two-
20 and three-phase equilibrium experimental measurements for the ternary CH₄ +
21 CO₂ + H₂O mixture, Fluid Phase Equilib. 451 (2017) 96–105.
22 <https://doi.org/10.1016/j.fluid.2017.08.002>.
- 23 [7] J.H. Erbar, A.K. Jagota, S. Muthswamy, M. Mosheghian, Prediction synthetic gas and
24 natural gas thermodynamic properties using a modified Soave-Redlich-Kwong
25 equation of state, Tulsa, 1980.
- 26 [8] P. Kastanidis, V.K. Michalis, G.E. Romanos, A.K. Stubos, I.G. Economou, I.N.
27 Tsimpanogiannis, Solubility of Methane and Carbon Dioxide in the Aqueous Phase of
28 the Ternary (Methane + Carbon Dioxide + Water) Mixture: Experimental
29 Measurements and Molecular Dynamics Simulations, J. Chem. Eng. Data. 63 (2018)
30 1027–1035. <https://doi.org/10.1021/acs.jced.7b00777>.
- 31 [9] L.N. Legoix, L. Ruffine, J.P. Donval, M. Haeckel, Phase equilibria of the CH₄-CO₂ binary
32 and the CH₄-CO₂-H₂O ternary mixtures in the presence of a CO₂-rich liquid phase,
33 Energies. (2017). <https://doi.org/10.3390/en10122034>.
- 34 [10] K. Ohgaki, K. Takano, H. Sangawa, T. Matsubara, S. Nakano, Methane exploitation by
35 carbon dioxide from gas hydrates - Phase equilibria for CO₂-CH₄ mixed hydrate
36 system, J. Chem. Eng. Japan. (1996). <https://doi.org/10.1252/jcej.29.478>.

- 1 [11] J. Qin, R.J. Rosenbauer, Z. Duan, Experimental Measurements of Vapor–Liquid
2 Equilibria of the H₂O + CO₂ + CH₄ Ternary System, *J. Chem. Eng. Data.* 53 (2008)
3 1246–1249. <https://doi.org/10.1021/je700473e>.
- 4 [12] K.Y. Song, R. Kobayashi, The Water Content of a CO₂-Rich Gas Mixture Containing
5 5.31 mol % Methane along the Three-Phase and Supercritical Conditions, *J. Chem.*
6 *Eng. Data.* 35 (1990) 320–322. <https://doi.org/10.1021/je00061a026>.
- 7 [13] C. Jarne, S.T. Blanco, M.A. Gallardo, E. Rauzy, S. Otín, I. Velasco, Dew points of ternary
8 methane (or ethane) + carbon dioxide + water mixtures: Measurement and
9 correlation, *Energy and Fuels.* 18 (2004) 396–404.
10 <https://doi.org/10.1021/ef030146u>.
- 11 [14] W.A. Fouad, M. Yarrison, K.Y. Song, K.R. Cox, W.G. Chapman, High pressure
12 measurements and molecular modeling of the water content of acid gas containing
13 mixtures, *AIChE J.* 61 (n.d.) 3038–3052. <https://doi.org/10.1002/aic.14885>.
- 14 [15] S.Z.S. Al Ghafri, E. Forte, G.C. Maitland, J.J. Rodriguez-Henríquez, J.P.M. Trusler,
15 Experimental and Modeling Study of the Phase Behavior of (Methane + CO₂ + Water)
16 Mixtures, *J. Phys. Chem. B.* 118 (2014) 14461–14478.
17 <https://doi.org/10.1021/jp509678g>.
- 18 [16] V. Belandria, A. Eslamimanesh, A.H. Mohammadi, P. Théveneau, H. Legendre, D.
19 Richon, Compositional analysis and hydrate dissociation conditions measurements for
20 carbon dioxide + methane + water system, *Ind. Eng. Chem. Res.* 50 (2011) 5783–5794.
21 <https://doi.org/10.1021/ie101959t>.
- 22 [17] Y.T. Seo, H. Lee, J.H. Yoon, Hydrate phase equilibria of the carbon dioxide, methane,
23 and water system, *J. Chem. Eng. Data.* 46 (2001) 381–384.
24 <https://doi.org/10.1021/je000237a>.
- 25 [18] H. Bruusgaard, J.G. Beltrán, P. Servio, Solubility measurements for the CH₄ + CO₂ +
26 H₂O system under hydrate-liquid-vapor equilibrium, *Fluid Phase Equilib.* 296 (2010)
27 106–109. <https://doi.org/10.1016/j.fluid.2010.02.042>.
- 28 [19] Y.T. Seo, H. Lee, Multiple-phase hydrate equilibria of the ternary carbon dioxide,
29 methane, and water mixtures, *J. Phys. Chem. B.* 105 (2001) 10084–10090.
30 <https://doi.org/10.1021/jp011095+>.
- 31 [20] A. Chapoy, H. Haghighi, R. Burgass, B. Tohidi, Gas hydrates in low water content
32 gases: Experimental measurements and modelling using the CPA equation of state,
33 *Fluid Phase Equilib.* 296 (2010) 9–14.
34 <https://doi.org/https://doi.org/10.1016/j.fluid.2009.11.026>.
- 35 [21] A. Chapoy, H. Haghighi, R. Brugrass, B. Tohidi, On the phase behaviour of the (carbon
36 dioxide + water) system at low temperatures: Experimental and modeling, *J. Chem.*
37 *Eng.* 47 (2012) 6–12.
- 38 [22] L. V. Jasperson, J.W. Kang, C.S. Lee, D. MacKlin, P.M. Mathias, R.J. McDougal, W.G.
39 Rho, D. Vonniederhausern, Experimental Determination of the Equilibrium Water
40 Content of CO₂ at High Pressure and Low Temperature, *J. Chem. Eng.*
41 *Data.* 60 (2015) 2674–2683. <https://doi.org/10.1021/acs.jced.5b00320>.
- 42 [23] G.K. Folas, E.W. Froyna, J. Lovland, G.M. Kontogeorgis, E. Solbraa, Data and prediction
43 of water content of high pressure nitrogen, methane and natural gas, *Fluid Phase*
44 *Equilib.* 252 (2007) 162–174.
45 <https://doi.org/https://doi.org/10.1016/j.fluid.2006.12.018>.
- 46 [24] A. Chapoy, C. Coquelet, D. Richon, Solubility measurement and modeling of water in
47 the gas phase of the methane/water binary system at temperatures from 283.08 to

- 1 318.12K and pressures up to 34.5MPa, *Fluid Phase Equilib.* 214 (2003) 101–117.
2 [https://doi.org/https://doi.org/10.1016/S0378-3812\(03\)00322-4](https://doi.org/https://doi.org/10.1016/S0378-3812(03)00322-4).
- 3 [25] K.Y. Song, R. Kobayashi, Water Content of CO₂ in Equilibrium with Liquid Water
4 and/or Hydrates, *SPE Form. Eval.* 2 (1987) 500–508.
- 5 [26] H.G. Donnelly, D.L. Katz, Phase Equilibria in the Carbon Dioxide–Methane System, *Ind.*
6 *Eng. Chem.* (1954). <https://doi.org/10.1021/ie50531a036>.
- 7 [27] R. Wiebe, V.L. Gaddy, Vapor Phase Composition of Carbon Dioxide-Water Mixtures at
8 Various Temperatures and at Pressures to 700 Atmospheres, *J. Am. Chem. Soc.* 63
9 (1941) 475–477. <https://doi.org/10.1021/ja01847a030>.
- 10 [28] A.D. King, C.R. Coan, Solubility of water in compressed carbon dioxide, nitrous oxide,
11 and ethane. Evidence for hydration of carbon dioxide and nitrous oxide in the gas
12 phase, *J. Am. Chem. Soc.* 93 (1971) 1857–1862.
13 <https://doi.org/10.1021/ja00737a004>.
- 14 [29] P.C. Gillespie, G.M. Wilson, GPA Research Report: Vapor-Liquid and Liquid-Liquid
15 Equilibria: Water - Methane, Water - Carbon Dioxide, Water - Hydrogen Sulfide,
16 Water - n-Pentane, Water - Methane - n-Pentane, 1982.
- 17 [30] K. Ohgaki, M. Nishikawa, T. Furuichi, T. Katayama, Entrainer Effect of H₂O and
18 Ethanol on alpha-Tocopherol Extraction by Compressed CO₂, *Kagaku Kogaku*
19 *Ronbunshu.* (1988).
- 20 [31] T. Nakayama, H. Sagara, K. Arai, S. Saito, High pressure liquid-liquid equilibria for the
21 system of water, ethanol and 1,1-difluoroethane at 323.2 K, *Fluid Phase Equilib.* 38
22 (1987) 109–127. [https://doi.org/https://doi.org/10.1016/0378-3812\(87\)90007-0](https://doi.org/https://doi.org/10.1016/0378-3812(87)90007-0).
- 23 [32] R. Burgass, A. Chapoy, P. Duchet-Suchaux, B. Tohidi, Experimental water content
24 measurements of carbon dioxide in equilibrium with hydrates at (223.15 to 263.15)K
25 and (1.0 to 10.0)MPa, *J. Chem. Thermodyn.* 69 (2014) 1–5.
26 <https://doi.org/https://doi.org/10.1016/j.jct.2013.09.033>.
- 27 [33] J.A. Briones, J.C. Mullins, M.C. Thies, B.-U. Kim, Ternary phase equilibria for acetic
28 acid-water mixtures with supercritical carbon dioxide, *Fluid Phase Equilib.* 36 (1987)
29 235–246. [https://doi.org/https://doi.org/10.1016/0378-3812\(87\)85026-4](https://doi.org/https://doi.org/10.1016/0378-3812(87)85026-4).
- 30 [34] I.P. Sidorov, Y.S. Kazarnovskii, A.M. Goldman, The Solubility of Water in compressed
31 Gases, *Tr. Gos. Nauchno Issled. Proektn. Inst. Azotn. Promst. Prod. Org. Sin.* 1 (1953)
32 48–67.
- 33 [35] R.D. Smith, H.R. Udseth, B.W. Wright, Micro-scale methods for characterization of
34 supercritical fluid extraction and fractionation processes, *Process Technol. Proc.* 3
35 (1985) 191–223.
- 36 [36] G. Mueller, phd. thesis, Univ. Kaiserslautern, 1983.
- 37 [37] J.A. Nighswander, N. Kalogerakis, A.K. Mehrotra, Solubilities of Carbon Dioxide in
38 Water and 1 Wt % NaCl Solution At Pressures up to 10 MPa and Temperatures from
39 80 to 200 °C, *J. Chem. Eng. Data.* (1989). <https://doi.org/10.1021/je00057a027>.
- 40 [38] A. Bamberger, G. Sieder, G. Maurer, High-pressure (vapor+liquid) equilibrium in
41 binary mixtures of (carbon dioxide+water or acetic acid) at temperatures from 313 to
42 353 K, *J. Supercrit. Fluids.* 17 (2000) 97–110.
43 [https://doi.org/https://doi.org/10.1016/S0896-8446\(99\)00054-6](https://doi.org/https://doi.org/10.1016/S0896-8446(99)00054-6).
- 44 [39] D.-Q.Q. Zheng, W.-D.D. Ma, R. Wei, T.-M.M. Guo, Solubility study of methane, carbon
45 dioxide and nitrogen in ethylene glycol at elevated temperatures and pressures, *Fluid*
46 *Phase Equilib.* 155 (1999) 277–286. [https://doi.org/https://doi.org/10.1016/S0378-](https://doi.org/https://doi.org/10.1016/S0378-3812(98)00469-5)
47 [3812\(98\)00469-5](https://doi.org/https://doi.org/10.1016/S0378-3812(98)00469-5).

- 1 [40] C.F. Prutton, R.L. Savage, The Solubility of Carbon Dioxide in Calcium Chloride-Water
2 Solutions at 75, 100, 120° and High Pressures, *J. Am. Chem. Soc.* (1945).
3 <https://doi.org/10.1021/ja01225a047>.
- 4 [41] J. Kiepe, S. Horstmann, K. Fischer, J. Gmehling, Experimental determination and
5 prediction of gas solubility data for CO₂ + H₂O mixtures containing NaCl or KCl at
6 temperatures between 313 and 393 K and pressures up to 10 MPa, *Ind. Eng. Chem.*
7 *Res.* (2002). <https://doi.org/10.1021/ie020154i>.
- 8 [42] D.Q. Zheng, T.M. Guo, H. Knapp, Experimental and modeling studies on the solubility
9 of CO₂, CHClF₂, CHF₃, C₂H₂F₄ and C₂H₄F₂ in water and aqueous NaCl solutions
10 under low pressures, *Fluid Phase Equilib.* (1997). [https://doi.org/10.1016/s0378-](https://doi.org/10.1016/s0378-3812(96)03177-9)
11 [3812\(96\)03177-9](https://doi.org/10.1016/s0378-3812(96)03177-9).
- 12 [43] M.S.-W. Wei, T.S. Brown, A.J. Kidnay, E.D. Sloan, Vapor + Liquid Equilibria for the
13 Ternary System Methane + Ethane + Carbon Dioxide at 230 K and Its Constituent
14 Binaries at Temperatures from 207 to 270 K, *J. Chem. Eng. Data.* 40 (1995) 726–731.
15 <https://doi.org/10.1021/je00020a002>.
- 16 [44] S. Bando, F. Takemura, M. Nishio, E. Hihara, M. Akai, Solubility of CO₂ in aqueous
17 solutions of NaCl at (30 to 60)°C and (10 to 20) MPa, *J. Chem. Eng. Data.* (2003).
18 <https://doi.org/10.1021/je0255832>.
- 19 [45] L. Ruffine, J.P.M. Trusler, Phase behaviour of mixed-gas hydrate systems containing
20 carbon dioxide, *J. Chem. Thermodyn.* (2010).
21 <https://doi.org/10.1016/j.jct.2009.11.019>.
- 22 [46] J.M. Han, H.Y. Shin, B.M. Min, K.H. Han, A. Cho, Measurement and correlation of high
23 pressure phase behavior of carbon dioxide + water system, *J. Ind. Eng. Chem.* (2009).
24 <https://doi.org/10.1016/j.jiec.2008.09.012>.
- 25 [47] P.J. Carvalho, L.M.C. Pereira, N.P.F. Gonçalves, A.J. Queimada, J.A.P. Coutinho, Carbon
26 dioxide solubility in aqueous solutions of NaCl: Measurements and modeling with
27 electrolyte equations of state, *Fluid Phase Equilib.* (2015).
28 <https://doi.org/10.1016/j.fluid.2014.12.043>.
- 29 [48] A. Zawisza, B.B. Malesinska, Solubility of carbon dioxide in liquid water and of water
30 in gaseous carbon dioxide in the range 0.2-5 MPa and at temperatures up to 473 K, *J.*
31 *Chem. Eng. Data.* 26 (1981) 388–391. <https://doi.org/10.1021/je00026a012>.
- 32 [49] S. Muromachi, A. Shijima, H. Miyamoto, R. Ohmura, Experimental measurements of
33 carbon dioxide solubility in aqueous tetra-n-butylammonium bromide solutions, *J.*
34 *Chem. Thermodyn.* (2015). <https://doi.org/10.1016/j.jct.2015.01.008>.
- 35 [50] S.X. Hou, G.C. Maitland, J.P.M. Trusler, Measurement and modeling of the phase
36 behavior of the (carbon dioxide + water) mixture at temperatures from 298.15 K to
37 448.15 K, *J. Supercrit. Fluids.* 73 (2013) 87–96.
38 <https://doi.org/10.1016/j.supflu.2012.11.011>.
- 39 [51] R. D'Souza, J.R. Patrick, A.S. Teja, High Pressure Phase Equilibria in the Carbon Dioxide
40 - n-Hexadecane and Carbon Dioxide - Water Systems, *Can. J. Chem. Eng.* (1988).
- 41 [52] S.D. Zaalishvili, The solubility of carbon dioxide mixed with hydrogen and nitrogen in
42 water under pressure, *Zh. Fiz. Khim.* 14 (1940) 413–417.
- 43 [53] F. Lucile, P. Cézac, F. Contamine, J.P. Serin, D. Houssin, P. Arpentinier, Solubility of
44 carbon dioxide in water and aqueous solution containing sodium hydroxide at
45 temperatures from (293.15 to 393.15) K and pressure up to 5 MPa: Experimental
46 measurements, *J. Chem. Eng. Data.* (2012). <https://doi.org/10.1021/je200991x>.
- 47 [54] L.A. Webster, A.J. Kidnay, Vapor-Liquid Equilibria for the Methane-Propane-Carbon

- 1 Dioxide Systems at 230 K and 270 K, *J. Chem. Eng. Data.* 46 (2001) 759–764.
2 <https://doi.org/10.1021/je000307d>.
- 3 [55] M.B. King, A. Mubarak, J.D. Kim, T.R. Bott, The mutual solubilities of water with
4 supercritical and liquid carbon dioxides, *J. Supercrit. Fluids.* 5 (1992) 296–302.
5 [https://doi.org/https://doi.org/10.1016/0896-8446\(92\)90021-B](https://doi.org/https://doi.org/10.1016/0896-8446(92)90021-B).
- 6 [56] M. Frost, E. Karakatsani, N. Von Solms, D. Richon, G.M. Kontogeorgis, Vapor-liquid
7 equilibrium of methane with water and methanol. Measurements and modeling, *J.*
8 *Chem. Eng. Data.* 59 (2014) 961–967. <https://doi.org/10.1021/je400684k>.
- 9 [57] M. Rigby, J.M. Prausnitz, Solubility of water in compressed nitrogen, argon, and
10 methane, *J. Phys. Chem.* 72 (1968) 330–334. <https://doi.org/10.1021/j100847a064>.
- 11 [58] A.H. Mohammadi, A. Chapoy, D. Richon, B. Tohidi, Experimental Measurement and
12 Thermodynamic Modeling of Water Content in Methane and Ethane Systems, *Ind.*
13 *Eng. Chem. Res.* 43 (2004) 7148–7162. <https://doi.org/10.1021/ie049843f>.
- 14 [59] Y.D. Zel'venskii, Solubility of CO₂ in water under pressure, *Zh. Khim. Prom.* 14 (1937)
15 1250–1257.
- 16 [60] S. Mao, Z. Duan, D. Zhang, L. Shi, Y. Chen, J. Li, Thermodynamic modeling of binary
17 CH₄–H₂O fluid inclusions, *Geochim. Cosmochim. Acta.* 75 (2011) 5892–5902.
18 <https://doi.org/https://doi.org/10.1016/j.gca.2011.07.021>.
- 19 [61] J. Davalos, W.R. Anderson, R.E. Phelps, A.J. Kidnay, Liquid-vapor equilibria at
20 250.00 deg.K for systems containing methane, ethane, and carbon dioxide, *J. Chem.*
21 *Eng. Data.* 21 (1976) 81–84. <https://doi.org/10.1021/je60068a030>.
- 22 [62] T.A. Al-Sahhaf, A.J. Kidnay, E.D. Sloan, Liquid + vapor equilibriums in the nitrogen +
23 carbon dioxide + methane system, *Ind. Eng. Chem. Fundam.* 22 (1983) 372–380.
24 <https://doi.org/10.1021/i100012a004>.
- 25 [63] Q. Nasir, K.M. Sabil, K.K. Lau, Measurement of isothermal (vapor+liquid) equilibria,
26 (VLE) for binary (CH₄+CO₂) from T=(240.35 to 293.15) K and CO₂ rich synthetic
27 natural gas systems from T=(248.15 to 279.15) K, *J. Nat. Gas Sci. Eng.* 27 (2015) 158–
28 167. <https://doi.org/https://doi.org/10.1016/j.jngse.2015.08.045>.
- 29 [64] F.A. Somait, A.J. Kidnay, Liquid-vapor equilibriums at 270.00 K for systems containing
30 nitrogen, methane, and carbon dioxide, *J. Chem. Eng. Data.* 23 (1978) 301–305.
31 <https://doi.org/10.1021/je60079a019>.
- 32 [65] N. Xu, J. Dong, Y. Wang, J. Shi, High pressure vapor liquid equilibria at 293 K for
33 systems containing nitrogen, methane and carbon dioxide, *Fluid Phase Equilib.* 81
34 (1992) 175–186. [https://doi.org/https://doi.org/10.1016/0378-3812\(92\)85150-7](https://doi.org/https://doi.org/10.1016/0378-3812(92)85150-7).
- 35 [66] M.J. Huron, J. Vidal, New mixing rules in simple equations of state for representing
36 vapour-liquid equilibria of strongly non-ideal mixtures, *Fluid Phase Equilib.* (1979).
37 [https://doi.org/10.1016/0378-3812\(79\)80001-1](https://doi.org/10.1016/0378-3812(79)80001-1).
- 38 [67] K.S. Pedersen, J. Milter, C.P. Rasmussen, Mutual solubility of water and a reservoir
39 fluid at high temperatures and pressures, *Fluid Phase Equilib.* 189 (2001) 85–97.
40 [https://doi.org/10.1016/s0378-3812\(01\)00562-3](https://doi.org/10.1016/s0378-3812(01)00562-3).
- 41 [68] A. Austegard, E. Solbraa, G. de Koeijer, J. Molnkiv, Thermodynamic Models for
42 Calculating Mutual Solubilities in H₂O-CO₂-CH₄ mixtures, *Chem. Eng. Res. Des.* 84
43 (2006).
- 44 [69] G.M. Kontogeorgis, E.C. Voutsas, I. V. Yakoumis, D.P. Tassios, An Equation of State for
45 Associating Fluids, *Ind. Eng. Chem. Res.* 35 (1996) 4310–4318.
46 <https://doi.org/10.1021/ie9600203>.
- 47 [70] G.M. Kontogeorgis, I. V Yakoumis, H. Meijer, E. Hendriks, T. Moorwood,

- 1 Multicomponent phase equilibrium calculations for water–methanol–alkane
2 mixtures, *Fluid Phase Equilib.* 158–160 (1999) 201–209.
3 [https://doi.org/https://doi.org/10.1016/S0378-3812\(99\)00060-6](https://doi.org/https://doi.org/10.1016/S0378-3812(99)00060-6).
- 4 [71] A. Valtz, A. Chapoy, C. Coquelet, P. Paricaud, D. Richon, Vapour–liquid equilibria in
5 the carbon dioxide–water system, measurement and modelling from 278.2 to 318.2K,
6 *Fluid Phase Equilib.* 226 (2004) 333–344.
7 <https://doi.org/https://doi.org/10.1016/j.fluid.2004.10.013>.
- 8 [72] Z.L. Yang, H.Y. Yu, Z.W. Chen, S.Q. Cheng, J.Z. Su, A compositional model for CO₂
9 flooding including CO₂ equilibria between water and oil using the Peng–Robinson
10 equation of state with the Wong–Sandler mixing rule, *Pet. Sci.* (2019).
11 <https://doi.org/10.1007/s12182-018-0294-2>.
- 12 [73] J. Gernert, R. Span, EOS-CG: A Helmholtz energy mixture model for humid gases and
13 CCS mixtures, *J. Chem. Thermodyn.* 93 (2016) 274–293.
14 <https://doi.org/10.1016/j.jct.2015.05.015>.
- 15 [74] S. Herrig, New Helmholtz-Energy Equations of State for Pure Fluids and CCS-Relevant
16 Mixtures, (2018) 260.
- 17 [75] A. Aasen, M. Hammer, G. Skaugen, J.P. Jakobsen, Ø. Wilhelmsen, Thermodynamic
18 models to accurately describe the PVTxy-behavior of water / carbon dioxide mixtures,
19 *Fluid Phase Equilib.* 442 (2017) 125–139.
- 20 [76] C.H. Twu, D. Bluck, J.R. Cunningham, J.E. Coon, A cubic equation of state with a new
21 alpha function and a new mixing rule, *Fluid Phase Equilib.* 69 (1991) 33–50.
22 [https://doi.org/10.1016/0378-3812\(91\)90024-2](https://doi.org/10.1016/0378-3812(91)90024-2).
- 23 [77] M.-J. Huron, J. Vidal, New mixing rules in simple equations of state for representing
24 vapour-liquid equilibria of strongly non-ideal mixtures, *Fluid Phase Equilib.* 3 (1979)
25 255–271. [https://doi.org/https://doi.org/10.1016/0378-3812\(79\)80001-1](https://doi.org/https://doi.org/10.1016/0378-3812(79)80001-1).
- 26 [78] D.S.H. Wong, S.I. Sandler, A theoretically correct mixing rule for cubic equations of
27 state, *AIChE J.* 38 (1992) 671–680. <https://doi.org/10.1002/aic.690380505>.
- 28 [79] E. Petropoulou, G.D. Pappa, E. Voutsas, Fluid Phase Equilibria Modelling of phase
29 equilibrium of natural gas mixtures containing associating compounds, *Fluid Phase*
30 *Equilib.* 433 (2017) 135–148. <https://doi.org/10.1016/j.fluid.2016.10.028>.
- 31 [80] N. Von Solms, M.L. Michelsen, G.M. Kontogeorgis, Computational and Physical
32 Performance of a Modified PC-SAFT Equation of State for Highly Asymmetric and
33 Associating Mixtures, *Ind. Eng. Chem. Res.* 42 (2003) 1098–1105.
34 <https://doi.org/10.1021/ie020753p>.
- 35 [81] O. Kunz, W. Wagner, The GERG-2008 Wide-Range Equation of State for Natural Gases
36 and Other Mixtures: An Expansion of GERG-2004, *J. Chem. Eng. Data.* 57 (2012) 3032–
37 3091. <https://doi.org/10.1021/je300655b>.
- 38 [82] E. Voutsas, K. Magoulas, D. Tassios, Universal mixing rule for cubic equations of state
39 applicable to symmetric and asymmetric systems: Results with the Peng-Robinson
40 equation of state, *Ind. Eng. Chem. Res.* 43 (2004) 6238–6246.
41 <https://doi.org/10.1021/ie049580p>.
- 42 [83] E. Collinet, J. Gmehling, Prediction of phase equilibria with strong electrolytes with
43 the help of the volume translated Peng-Robinson group contribution equation of
44 state (VTPR), *Fluid Phase Equilib.* (2006). <https://doi.org/10.1016/j.fluid.2006.05.033>.
- 45 [84] Z.-L. Yang, H.-Y. Yu, Z.-W. Chen, S.-Q. Cheng, J.-Z. Su, A compositional model for CO₂
46 flooding including CO₂ equilibria between water and oil using Peng-Robinson
47 equation of state with Wong-Sandler mixing rule, *Pet. Sci.* (2019) 1–16.

- 1 [85] A. Chapoy, R. Burgass, A. Terrigeol, C. Coquelet, Water Content of CO₂-rich
2 Mixtures: Measurements and Modeling using the Cubic-Plus-Association Equation of
3 State, *J. Nat. Gas Eng.* 1 (2016) 85–97. <https://doi.org/10.7569/jnge.2015.692505>.
- 4 [86] I. Tsvintzelis, S. Ali, G.M. Kontogeorgis, Modeling phase equilibria for acid gas
5 mixtures using the cubic-plus-association equation of state. 3. Applications relevant
6 to liquid or supercritical CO₂ transport, *J. Chem. Eng. Data.* 59 (2014) 2955–2972.
7 <https://doi.org/10.1021/je500090q>.
- 8 [87] P. Reshadi, K.H. Nasrifar, M. Moshfeghian, Evaluating the phase equilibria of liquid
9 water+natural gas mixtures using cubic equations of state with asymmetric mixing
10 rules, *Fluid Phase Equilib.* 302 (2011) 179–189.
11 <https://doi.org/10.1016/j.fluid.2010.08.007>.
- 12 [88] K.Y.S.K.R.C.W.G.C. Wael A. Fouad Matt Yarrison, High Pressure Measurements and
13 Molecular Modeling of the Water Content of Acid Gas Containing Mixtures, *AIChE J.*
14 (2015).
- 15 [89] K. Nasrifar, F. Alavi, J. Javanmardi, Prediction of water content of natural gases using
16 the PC-SAFT equation of state, *Fluid Phase Equilib.* 453 (2017) 40–45.
17 <https://doi.org/https://doi.org/10.1016/j.fluid.2017.08.023>.
- 18 [90] V. Papaioannou, F. Calado, T. La, S. Dufal, M. Sadeqzadeh, G. Jackson, C.S. Adjiman, A.
19 Galindo, Application of the SAFT- g Mie group contribution equation of state to fluids
20 of relevance to the oil and gas industry, 416 (2016) 104–119.
21 <https://doi.org/10.1016/j.fluid.2015.12.041>.
- 22 [91] H. Zhao, Modeling vapor-liquid phase equilibria of methane-water and methane-
23 carbon dioxide-water systems at 274K to 573K and 0.1 to 150 MPa using PRSV
24 equation of state and Wong-Sandler mixing rule, *Fluid Phase Equilib.* 447 (2017) 12–
25 26. <https://doi.org/10.1016/j.fluid.2017.05.015>.
- 26 [92] H. Zhao, S.N. Lvov, Phase behavior of the CO₂–H₂O system at temperatures of 273–
27 623 K and pressures of 0.1–200 MPa using Peng-Robinson-Stryjek-Vera equation of
28 state with a modified Wong-Sandler mixing rule: An extension to the CO₂–CH₄–H₂O
29 system, *Fluid Phase Equilib.* 417 (2016) 96–108.
30 <https://doi.org/https://doi.org/10.1016/j.fluid.2016.02.027>.
- 31 [93] J.M. Míguez, M.C. Dos Ramos, M.M. Piñeiro, F.J. Blas, An examination of the ternary
32 methane + carbon dioxide + water phase diagram using the SAFT-VR approach, *J.*
33 *Phys. Chem. B.* 115 (2011) 9604–9617. <https://doi.org/10.1021/jp2017488>.
- 34 [94] A. Chapoy, R. Burgass, B. Tohidi, J.M. Austell, C. Eickhoff, Effect of Common Impurities
35 on the Phase Behavior of Carbon-Dioxide-Rich Systems: Minimizing the Risk of
36 Hydrate Formation and Two-Phase Flow, *SPE J.* 16 (2011) 921–930.
37 <https://doi.org/10.2118/123778-PA>.
- 38 [95] A. Chapoy, H. Haghighi, R. Burgass, B. Tohidi, On the phase behaviour of the (carbon
39 dioxide + water) systems at low temperatures: Experimental and modelling, *J. Chem.*
40 *Thermodyn.* 47 (2012) 6–12. <https://doi.org/10.1016/j.jct.2011.10.026>.
- 41 [96] P.M. Mathias, T.W. Copeman, Extension of the Peng-Robinson equation of state to
42 complex mixtures: Evaluation of the various forms of the local composition concept,
43 *Fluid Phase Equilib.* 13 (1983) 91–108.
- 44 [97] J.-N. Jaubert, F. Mutelet, VLE predictions with the Peng–Robinson equation of state
45 and temperature dependent kij calculated through a group contribution method,
46 *Fluid Phase Equilib.* 224 (2004) 285–304.
47 <https://doi.org/https://doi.org/10.1016/j.fluid.2004.06.059>.

- 1 [98] J.N. Jaubert, R. Privat, Relationship between the binary interaction parameters (k_{ij}) of
2 the Peng-Robinson and those of the Soave-Redlich-Kwong equations of state:
3 Application to the definition of the PR2SRK model, *Fluid Phase Equilib.* (2010).
4 <https://doi.org/10.1016/j.fluid.2010.03.037>.
- 5 [99] J. Gernert, R. Span, EOS–CG: A Helmholtz energy mixture model for humid gases and
6 CCS mixtures, *J. Chem. Thermodyn.* 93 (2016) 274–293.
7 <https://doi.org/https://doi.org/10.1016/j.jct.2015.05.015>.
- 8 [100] Z. Duan, S. Mao, A thermodynamic model for calculating methane solubility, density
9 and gas phase composition of methane-bearing aqueous fluids from 273 to 523K and
10 from 1 to 2000bar, *Geochim. Cosmochim. Acta.* 70 (2006) 3369–3386.
11 <https://doi.org/https://doi.org/10.1016/j.gca.2006.03.018>.
- 12 [101] R. Burgass, A. Chapoy, B. Tohidi, Experimental and Modelling Low Temperature Water
13 Content in Multicomponent Gas Mixtures, in: *Proceeding 7th Int. Conf. Gas Hydrates*
14 *(ICGH 2011)*, Edinburgh, 2011.
- 15 [102] A. Chapoy, R. Brugrass, B. Tohidi, J.M. Austell, C. Eickhoff, Effect of Common
16 Impurities on the Phase Behaviour of Carbon-Dioxide-Rich Systems: Minimizing the
17 Risk of Hydrate Formation and Two-Phase Flow, *SPE J.* SPE-123778 (2011).
- 18 [103] H. Haghghi, A. Chapoy, R. Burgass, B. Tohidi, on the Phase Behaviour of the Carbon
19 Dioxide - Water Systems At Low Temperatures, *Icgh 2011.* 44 (2011).
- 20 [104] P. Ahmadi, A. Chapoy, CO₂ solubility in formation water under sequestration
21 conditions, *Fluid Phase Equilib.* 463 (2018) 80–90.
22 <https://doi.org/10.1016/j.fluid.2018.02.002>.
- 23 [105] A. Chapoy, R. Burgass, A. Terrigeol, C. Coquelet, Water Content of CO₂ rich Mixtures:
24 Measurements and Modeling using the Cubic-Plus-Association Equation of State, *J.*
25 *Nat. Gas Eng.* 1 (2016) 85–97.
26 [https://pdfs.semanticscholar.org/e45d/e7c3f6bc55d830d0058e4ade64d141b120f0.p
27 df?_ga=2.159672189.1728404663.1563924027-567500309.1563924027](https://pdfs.semanticscholar.org/e45d/e7c3f6bc55d830d0058e4ade64d141b120f0.pdf?_ga=2.159672189.1728404663.1563924027-567500309.1563924027).

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30