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# Density, compressibility and phase equilibrium of high pressure-high temperature reservoir fluids up to 473 K and 140 MPa

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## Abstract

Experimental measurement of volumetric and phase equilibrium properties of two high pressure - high temperature reservoir fluids was performed in this work. One fluid is volatile oil and the other is gas condensate. The density, isothermal compressibility, saturation pressure, and liquid fraction below the saturation pressure were determined in the temperature range up to 473.15 K by using a high pressure vibrating tube densitometer and a PVT cell. The obtained data were modelled by using equations of state including Soave-Redlich-Kwong (SRK), Peng-Robinson (PR), volume translated SRK (SRK-VT), volume translated PR (PR-VT) and Perturbed-Chain Statistical Association Fluid Theory (PC-SAFT). Among these models, the performance in saturation pressure prediction is system-dependent and it is hard to generalize the observation. PC-SAFT models volumetric properties satisfactorily. Regarding density calculation, it performs similarly to SRK-VT and PR-VT for the two fluids investigated here. Concerning isothermal compressibility, PC-SAFT is slightly better.

**Keywords:** density, compressibility, phase equilibrium, high pressure – high temperature, reservoir fluid

## 1. Introduction

The global increase in energy consumption puts a higher demand for energy from different sources including gas and oil. This has led to two clear changes in the oil and gas industry: development of unconventional resources, such as shale production in the US, and exploitation of oil and gas in more challenging environments, such as on-shore or offshore deep reservoirs usually at high pressures and high temperatures. Many of the high-pressure reservoirs related to the second change can be classified as high pressure - high temperature (HPHT) reservoirs, which, in the oil and gas industry, often refer to reservoirs with pressures higher than 69 MPa and temperatures higher than 422.15 K. These limits are somewhat arbitrary and HPHT generally indicates more extreme pressure and temperature conditions where we have less experience with and a much wider temperature and pressure range that the production needs to cover. Development of those reservoirs present numerous challenges in terms of safety, testing, seals, and materials [1]. A central issue is how to realize a cheaper, safer and more efficient development of these reservoirs.

The importance of fundamental thermodynamic properties for relevant HPHT systems can never be overestimated. The availability of thermodynamic properties of the reservoir fluids under these extreme conditions is low and there is only a limited experience in the industry in performing the standard PVT studies at very high pressure and temperature conditions [2]. The volumetric and phase equilibrium properties are necessary for the estimation of oil and gas resources and the description of the dynamic production process which involves single-phase expansion, phase transition and vapour-liquid equilibrium in the two-phase region. Density and compressibility play a crucial role in production forecast of the pressure depletion process [2, 3], as compressibility is a driving force in the primary recovery. Regarding gas condensates, once the reservoir pressure decreases below the saturation pressure, valuable liquid hydrocarbons can drop out and tend to stay in the reservoir [4]. The maximum liquid fraction below the saturation pressure for the gas condensates indicates the richness of the gas condensate [5]. It is desirable to avoid large drop-out since the condensate drop-out near the wellbore can decrease the gas relative permeability dramatically and thus the well productivity, which is known as condensate blocking [4, 6].

In our previous studies [7-9] a series of synthetic binary, ternary and multicomponent mixtures were designed as simple model fluids for real HPHT reservoir fluids. These mixtures include

methane/n-decane, methane/n-butane/n-decane, methane/n-butane/n-dodecane and methane/nbutane/n-hexane/n-decane/n-hexadecane/n-eicosane, and their compositions were designed to represent either volatile oils or gas condensates. Their volumetric properties and phase equilibria were experimentally determined in broad temperature and pressure ranges. We also compared the measured properties with predictions from different equations of state (EoSs), including Soave-Redlich-Kwong (SRK) [10], Peng-Robinson (PR) [11], Soave-Benedict-Webb-Rubin (S-BWR) [12], and Perturbed-Chain Statistical Association Fluid Theory (PC-SAFT) [13]. The comparison involves systems of different composition characteristics and various properties, and it is impossible to single out a model that performs always better than the others in all cases. Nevertheless, for saturation pressures, SRK, PR and PC-SAFT are similar while S-BWR tends to give significantly different phase envelopes for very asymmetric mixtures; for single-phase densities, PC-SAFT and S-BWR are generally better while PC-SAFT's performance seems to be more stable among the synthetic mixtures tested.

In the present work we extend our previous studies to two real reservoir fluids, one HPHT volatile oil and one HPHT gas condensate. We determined experimentally their densities, isothermal compressibilities, saturation pressures, and liquid fractions in the two-phase region in the temperature range from (298.15 to 473.15)K (the density and compressibility determined up to 463.15 K). A high pressure vibrating tube densitometer from Anton Paar was used for density determinations up to 140 MPa, and the study of phase equilibrium was carried out in a PVT cell from Sanchez Technologies. In the modelling part, we used the aforementioned EoSs except for S-BWR because S-BWR tends to predict very different saturation pressures for asymmetric mixtures. For SRK and PR, we also included the calculation with volume translation in order to improve their density calculation results. To apply these EoSs to reservoir fluids, appropriate characterization of their  $C_7^+$  fractions was also performed. We analysed and

compared the model performance for the prediction of volumetric properties and phase behaviour.

#### 2. Materials and methods

#### 2.1. Materials

Both the HPHT volatile oil sample and the HPHT gas condensate sample are bottom hole samples taken from HPHT reservoirs in the Danish North Sea. The samples were received in high pressure cylinders at ambient temperature and pressures of 48 MPa and 93 MPa for the volatile oil and the gas condensate, respectively. The received volatile oil and gas condensate samples were kept at constant pressure and conditioned at 366 K and 363 K, respectively. In order to determine the composition of the reservoir fluids, a separator test was performed for each fluid. The compositional analysis of the obtained gas and oil phases was performed through a gas chromatograph Agilent 7890A gas. The compositional data, together with the gravimetrical and volumetric measurements at ambient pressure allowed to establish the overall composition of the reservoir fluids along with their gas oil ratio (GOR), which is defined as the ratio of the gas volume to the oil volume of a reservoir sample at 0.1013 MPa and 288.75 K. Two replicates of the separator test were performed for each reservoir fluid.

The obtained composition of the reservoir fluids is presented in Table 1 for the HPHT volatile oil and the HPHT gas condensate. The volatile oil has a methane mole fraction of 49.3% whereas the methane mole fraction of the gas condensate is 82.7%, as concerns the mole fraction of  $C_{7^+}$ , it is 27.76% and 4.2% for the HPHT volatile oil and HPHT gas condensate, respectively.

Table 1

Compositional analysis (mole fraction %) and molecular weight of the HPHT reservoir fluids to  $C_{36}^+$ .

Component	HPHT Volatile oil	HPHT Gas condensate
CO <sub>2</sub>	0.14	2.40
$N_2$	1.41	0.10
C1	49.3	82.7

$C_2$	9.78	6.43
C <sub>3</sub>	4.67	2.34
iC <sub>4</sub>	0.62	0.38
$C_4$	1.32	0.65
iC <sub>5</sub>	0.22	0.19
C <sub>5</sub>	2.96	0.26
$C_6$	1.82	0.35
$C_7$	3.05	0.67
$C_8$	3.46	0.93
C <sub>9</sub>	2.69	0.61
$C_{10}$	2.30	0.40
C <sub>11</sub>	1.40	0.20
C <sub>12</sub>	2.30	0.25
C <sub>13</sub>	3.19	0.20
$C_{14}$	1.57	0.16
C <sub>15</sub>	0.96	0.11
C <sub>16</sub>	0.88	0.11
C <sub>17</sub>	0.65	0.07
C <sub>18</sub>	0.56	0.06
C <sub>19</sub>	0.40	0.05
C <sub>20</sub>	0.37	0.04
C <sub>21</sub>	0.40	0.04
C <sub>22</sub>	0.37	0.04
C <sub>23</sub>	0.32	0.03
C <sub>24</sub>	0.30	0.03
C <sub>25</sub>	0.26	0.02
C <sub>26</sub>	0.24	0.02
C <sub>27</sub>	0.21	0.02
$C_{28}$	0.20	0.02
C <sub>29</sub>	0.17	0.01
$C_{30}$	0.16	0.01
C <sub>31</sub>	0.14	0.01
C <sub>32</sub>	0.13	0.01
C <sub>33</sub>	0.11	0.01
C <sub>34</sub>	0.10	0.01
C <sub>35</sub>	0.09	0.01
$C_{36}^{+}$	0.78	0.05
Molecular weight	70.8	25.2

As concerns the gas oil ratio (GOR), it was found to be 224 Sm<sup>3</sup>/Sm<sup>3</sup> for the HPHT volatile oil and 2654 Sm<sup>3</sup>/Sm<sup>3</sup> for the HPHT gas condensate, where Sm<sup>3</sup>/Sm<sup>3</sup> stands for standard cubic meters of gas at 0.1013 MPa and 288.75 K per cubic meter of stock tank oil (STO) at the same conditions. The measured densities of the stock tank oil at 0.1013 MPa and 288.75 K were 0.8012 and 0.8001 g·cm<sup>-3</sup> for the HPHT volatile oil STO and the HPHT gas condensate STO, respectively. More details about the  $C_7^+$  and  $C_{36}^+$  fractions of the reservoir fluids are given in Table 2. For the volatile oil the density of the  $C_7^+$  fraction was obtained from the density measurements of the STO and the C<sub>6</sub> fraction (obtained from true boiling point distillation), whereas the  $C_{36}^+$  density was obtained by using the single carbon number densities previously reported by Katz and Firoozabadi [14] (extrapolated to  $C_{72}$  fraction) together with the STO mass fractions of the volatile oil measured with GC up to C72. As concerns the gas condensate, the densities of the  $C_{7^{+}}$  and the  $C_{36^{+}}$  fractions were obtained by combining the compositional data we have measured from the separator oil up to C72 with the density of the single carbon numbers reported by Katz and Firozobadi [14] (extrapolated to C<sub>72</sub> fraction).

Molecular weight and de	Molecular weight and density of the $C_7^+$ and $C_{36}^+$ fractions.							
Fluid	Fraction	Mole fraction %	Molecular weight	$ ho^*/ ext{g·cm}^{-3}$				
HPHT volatile oil	${ m C_7}^+$	27.76	189.4	0.8076				
	$C_{36}^{+}$	0.78	611.6	0.9367				
HPHT gas condensate	${ m C_7}^+$	4.20	162.4	0.8008				
	${ m C_{36}}^+$	0.05	593.7	0.9339				

\*Density at 288.75 K and 0.1 MPa

## 2.2. Density measurements

Table 2

The experimental determination of the density was carried out by using a high pressure vibrating tube densitometer DMA HPM from Anton Paar. In this apparatus, the oscillation period of a U-tube filled with the sample is measured and displayed with 7 significant figures. The calibration of the device was performed from (298.15 to 463.15)K and pressures up to 140 MPa, following a modified Lagourette et al. method [15, 16] which uses vacuum, Milli-Q water and n-dodecane as reference fluids. After calibration, the density of the studied samples was obtained from the oscillation period measurements. The density determination was performed for the reservoir fluids in the single phase, i.e. the measurements were performed at pressures higher than the saturation pressure  $(p^{sat})$ . The uncertainty of the density through this technique has been rigorously calculated by Segovia et al. [17]. In the present work the expanded (k=2) uncertainty of the density measurements is considered to be  $7 \cdot 10^{-4}$ g·cm<sup>-3</sup> at  $T \le 373.15$ K and 3·10<sup>-3</sup> g·cm<sup>-3</sup> at other temperature conditions.

The pressure was generated in the vibrating tube densitometer through a pressure line comprised of high pressure pipes and valves and a high pressure generator (HiP 37-6-30). The

pressure was measured by using a pressure transducer SIKA type P that can measure pressure up to 150 MPa with a 0.05% FS uncertainty. As concerns temperature control, it was performed by means of a liquid circulator bath Julabo PRESTO A30 and the temperature was measured through a Pt-100 located inside the measurement cell with an uncertainty of  $\pm 0.02$ K. A schematic of the complete setup for the density measurements has been previously published [7].

#### 2.3. Phase equilibrium measurements

The measurement of the phase equilibrium was performed in a PVT cell PVT 240/1500 FV with full visibility from Sanchez Technologies. The measurements can be performed in the temperature range up to 473.15 K and pressures up to 150 MPa. This apparatus consists of a stainless steel hollow cylinder with a motor driven piston in one end and a sapphire window in the other end. The motor driven piston is used to change the pressure inside the PVT cell by increasing or decreasing the cell volume, the maximum volume of the cell is 240 cm<sup>3</sup>. A stirring system composed of four retractable blades is located in the head of the piston and driven by a motor through a magnetic coupling, this stirring system is used to homogenize the studied sample. The full visibility of the cell is achieved through a CCD digital camera Lumenera Lw1335C located in front of the sapphire window together with a set of two lamps connected to four optical fibers. The cell has a rocking mechanism that allows its rotation to a certain angle depending on the type of fluid under study. The whole apparatus is computer controlled through the Falcon Software. A schematic of the experimental setup has been published elsewhere [9].

As concerns the temperature control, it is performed through a set of 8 electrical resistances located in the wall of the cell. Additionally, a liquid circulating thermostat Julabo Presto A40 is connected to a jacket covering the cell, the thermostating liquid circulates with a temperature set point 5 K lower than the sample temperature in order to improve the temperature stability

of the system. The temperature is measured by using a Pt-100 located in the wall of the cell with an uncertainty of  $\pm 0.02$  K. Regarding pressure, a pressure transducer Dynisco PT435A inserted in the wall of the cell is used for the pressure measurements, this pressure transducer is newly calibrated against a reference pressure transducer, as a function of temperature and pressure, prior to the loading of a new sample in the PVT cell. The pressure standard uncertainty is 0.06 MPa.

The experimental determination of the saturation pressure was performed through visual observation. The pressure was slowly decreased at constant temperature from the single phase with a constant flow rate of 1.4·10<sup>-4</sup> or 2.8·10<sup>-4</sup> cm<sup>3</sup>·s<sup>-1</sup> under continuous stirring until the appearance of a new phase was observed, at this point the pressure reading indicated the saturation pressure and it was recorded. Each determination was performed by triplicate for both reservoir fluid samples at selected temperatures covering the range from (298.15 to 473.15)K. The combined standard uncertainty of the determination of the equilibrium pressure is estimated to be 0.10 MPa for bubble points and 0.15 MPa for dew points. This uncertainty was estimated by combining the standard uncertainty of the pressure readings with the repeatability of the visual determinations of saturation pressure.

A constant mass expansion (CME) procedure was applied to both reservoir fluid samples at selected temperatures in the range from (298.15 to 473.15)K. This procedure was performed automatically through the Falcon Software. The measurements started from the single phase sample at constant temperature, a series of pressure steps were made in the single phase region, in each of them the sample was stirred for a certain period (at least 60 s) and after this a set of waiting periods of 600 s were applied until a pressure stability ( $\pm 0.05$  MPa) was achieved. After stabilization, the pressure and volume of the system were recorded and a photo of the sample was taken through the Euclide software version 1.4.2. Once the saturation pressure was reached through the aforementioned pressure steps, a series of volume steps were performed

in the two phase region. Equilibrium pressure and volume were also recorded for each step, as well as a photo of the sample. Afterwards, the liquid fraction percentage in the two phase region was measured from the photos by using the Euclide software, which based on geometrical relations determines the liquid volume for each of the steps. The maximum standard uncertainty of the liquid fractions reported in this work is 2.1%.

## 2.4. Modelling

The experimental data measured in this work have been modelled by using two cubic equations of state (EoSs), Soave-Redlich-Kwong (SRK), and Peng-Robinson (PR) and a non-cubic EoS, Perturbed Chain Statistical Associating Fluid Theory (PC-SAFT) [13, 18]. For the last EoS, its simplified version described by von Solms et al. [13] was used in the calculations.

In the volumetric calculations, the Peneloux [19] volume correction (*c*) can be applied to the cubic EoSs in order to compensate for their poor density prediction. The volumes calculated by the cubic EoSs ( $V_{\text{SRK}}$  and  $V_{\text{PR}}$ ) are related to the volumes calculated by the volume translation EoSs (SRK-VT and PR-VT) through the volume correction parameter of SRK ( $c_{\text{SRK}}$ ) and PR ( $c_{PR}$ ), as follows:

$$V_{SRK-VT} = V_{SRK} - c_{SRK} \tag{1}$$

$$V_{PR-VT} = V_{PR} - c_{PR} \tag{2}$$

where  $V_{\text{SRK-VT}}$  and  $V_{\text{PR-VT}}$  are the molar volumes calculated with SRK and PR with volume translation, respectively,  $V_{\text{SRK}}$  and  $V_{\text{PR}}$  are the molar volumes calculated with SRK and PR, respectively, and  $c_{\text{SRK}}$  and  $c_{PR}$  are the Peneloux volume correction for SRK and PR, respectively.

The Peneloux volume correction is calculated according to the following equation for both SRK ( $c_{SRK}$ ) and PR ( $c_{PR}$ ):

$$c = \sum_{i} x_i c_i \tag{3}$$

where  $x_i$  is the mole fraction of the single carbon number C<sub>i</sub> and  $c_i$  is the Peneloux volume correction of the single carbon number C<sub>i</sub>.

For single carbon number components with carbon number smaller than 7, their Peneloux volume corrections are calculated by equations 4 and 5 for SRK and PR, respectively [20]:

$$c_{SRK} = \frac{0.40768RT_c \left(0.29441 - Z_{RA}\right)}{p_c} / cm^3 mol^{-1}$$
(4)

$$c_{PR} = \frac{0.50033RT_c \left( 0.25969 - Z_{RA} \right)}{p_c} / cm^3 mol^{-1}$$
(5)

where *R* is the universal gas constant,  $T_c$  is the critical temperature,  $p_c$  is the critical pressure and  $Z_{RA}$  (Rackett compressibility factor) is given by the following equation [21]:

$$Z_{R4} = 0.29056 - 0.08775\omega \tag{6}$$

where  $\omega$  is the acentric factor.

For components in  $C_7^+$ , their shift parameters were obtained through characterization by matching the density of  $C_7^+$ . In this sense, the density calculation for SRK and PR with volume translation has involved certain adjustment to experimental data although in an indirect manner. For the characterization of the  $C_7^+$  fraction of the reservoir fluid samples, we used the methods described by Yan et al. [22] and Varzandeh et al. [3] which are based on the characterization method of Pedersen et al. [23, 24]. The difference from the method of Pedersen et al. lies in how the model parameters are estimated. The first step of characterization is the same for all EoS models: an exponential distribution was used for the calculation of mole fraction of the single carbon number (SCN) components in the plus fraction. In the second step, the parameter estimation for SRK and PR was performed by using Twu's correlation [25] for  $p_c$  and  $T_c$  and Lee-Kesler correlation [26] for  $\omega$ . For PC-SAFT, the correlations of Varzandeh et al. [3] were used to calculate the model parameters. In the last step the SCN components were lumped into a few pseudo-components with approximately equal mass.

The single compound model parameters as well as those for the lumped components are presented in tables A.1 and A.2 from the supplementary material. The binary interaction parameters,  $k_{ij}$ , used in this work and the mixing rules are provided in tables A.3 and A.4 of the supplementary material.

An in-house calculation code [3, 22] was used for all the calculations. In principle, group contribution based characterization methods such as the method of Skjold-Jørgensen [27] could also be used for characterization. But they require additional information for chemical groups which are usually not available from the chemical analytical data. The classical  $C_7^+$  characterization based on boiling points and fraction densities is therefore more appropriate here. Regarding the selection of EoS models, we include two cubics and one non-cubic with strong theoretical basis. But we have not included the association model like the cubic plus association (CPA) model [28] here. This is because the CPA model reduces to SRK for systems without associating components and the calculation is essentially SRK with a different set of parameters [28]. The application of CPA with reservoir fluids and associating compounds can be found in Yan et al. [29]

## 3. Results and discussion

In the present work, the comparison between the experimental data and the model predictions is performed through the absolute average deviation (AAD), the absolute average relative deviation (AARD) and the standard deviation ( $\sigma$ ), which are defined as follows:

$$AAD = \frac{1}{N} \sum_{i=1}^{N} \left| Y_i^{cal} - Y_i^{exp} \right|$$
(7)

AARD / % = 
$$\frac{100}{N} \sum_{i=1}^{N} \left| \frac{Y_i^{cal} - Y_i^{exp}}{Y_i^{exp}} \right|$$
 (8)

$$\sigma = \sqrt{\frac{\sum_{i=1}^{N} \left(Y_i^{\exp} - Y_i^{cal}\right)^2}{N - p}}$$
(9)

where  $Y_i^{exp}$  is the value of the experimental property determined in this work,  $Y_i^{cal}$  is the value predicted through the model for the same property, N is the number of experimental data points and p is the number of fitting parameters. In the comparison of density-related properties, the original SRK and PR, their volume translated versions, and PC-SAFT are included. However, volume-translated SRK and PR are excluded in the calculation of saturation pressures since volume translation does not affect phase equilibrium.

## 3.1. Density

The experimental density data of the two HPHT reservoir fluids in the studied temperature and pressure range is gathered in Table 3. Under the studied conditions, the density of the HPHT volatile oil varies in the range (0.54 to 0.74) g·cm<sup>-3</sup> whereas the density of the HPHT gas condensate varies in the range (0.30 to 0.50) g·cm<sup>-3</sup>. The expected trends of density with temperature and pressure were found, i.e. density increases with pressure along isotherms, whereas it decreases with temperature along isobars.

T/V	p/MPa					
1/K	40.00	60.00	80.00	100.00	120.00	140.00
			HPHT vol	latile oil		
298.15	0.6781	0.6945	0.7080	0.7197	0.7301	0.7393
323.15	0.6579	0.6768	0.6918	0.7047	0.7161	0.7261
348.15	0.6374	0.6590	0.6758	0.6899	0.7021	0.7129
358.15	0.6279	0.6505	0.6680	0.6826	0.6950	0.7062
373.15	0.6153	0.6395	0.6581	0.6734	0.6865	0.6980
423.15	0.5755	0.6060	0.6284	0.6463	0.6614	0.6745
432.15	0.5685	0.6002	0.6233	0.6417	0.6572	0.6707
463.15	0.5433	0.5799	0.6056	0.6258	0.6426	0.6570

Table 3 Experimental density data<sup>a</sup>,  $\rho$ , of the HPHT reservoir fluids in g·cm<sup>-3</sup>.

			HPHT gas c	ondensate		
298.15	_	_	0.4476	0.4681	0.4848	0.4987
323.15	_	_	0.4288	0.4510	0.4689	0.4839
348.15	—	0.3781	0.4107	0.4346	0.4536	0.4694
363.15	_	0.3649	0.3986	0.4236	0.4434	0.4599
373.15	—	0.3577	0.3930	0.4189	0.4392	0.4560
393.15	_	0.3435	0.3809	0.4082	0.4294	0.4466
423.75	_	0.3226	0.3621	0.3911	0.4135	0.4316
463.15	_	0.2962	0.3384	0.3693	0.3925	0.4112

<sup>a</sup>Expanded density uncertainty  $U(\rho)$  (k=2):  $0.7 \cdot 10^{-3}$  g·cm<sup>-3</sup> at T < 373.15 K;  $3 \cdot 10^{-3}$  g·cm<sup>-3</sup> at other temperature conditions; standard temperature uncertainty u(T): 0.02 K; standard pressure uncertainty u(p): 0.08 MPa.

The experimental density data for both reservoir fluids have been correlated as a function of temperature and pressure by using the following modified Tammann-Tait equation:

$$\rho(T,p) = \frac{\rho(T,p_{ref})}{1 - C \cdot \ln\left(\frac{B(T) + p}{B(T) + p_{ref}}\right)}$$
(10)

where  $\rho(T, p_{ref})$  is the density in g·cm<sup>-3</sup> as a function of temperature (K) at a reference pressure (MPa), given by the following equation:

$$\rho(T, p_{ref}) = \sum_{i=0}^{3} A_i T^i$$
(11)

*C* is a temperature and pressure independent parameter and B(T) is a temperature dependent parameter given by:

$$B(T) = \sum_{j=0}^{2} B_j T^j$$
(12)

The fitting parameters of Equation 10 are given in Table 4 together with  $\sigma$  and  $\sigma^*$ , which are the standard deviations for Equations 11 and 10, respectively. The absolute average relative deviation of the fit is also provided.

Table 4

Modified Tammann-Tait (Equation 10) parameters for the HPHT volatile oil and the HPHT gas condensate.  $\sigma$  is the standard deviation for Equation (11) and  $\sigma^*$  the standard deviation for equation (10).

HPHT volatile oil HPHT gas condensate

<i>p<sub>ref</sub></i> / MPa	40	60
$A_0/\mathrm{g}\cdot\mathrm{cm}^{-3}$	0.9434	0.8736
$10^{4} \cdot A_{I} / \text{g} \cdot \text{cm}^{-3} \cdot \text{K}^{-1}$	-9.35	-22.95
$10^{7} \cdot A_{2} / \text{g} \cdot \text{cm}^{-3} \cdot \text{K}^{-2}$	1.53	32.12
$10^9 \cdot A_3 / g \cdot cm^{-3} \cdot K^{-3}$		-2.046
$10^3 \cdot \sigma / \mathrm{g \cdot cm^{-3}}$	0.6	0.7
С	0.1024	0.145
$B_0$ /MPa	280.7	117.34
$B_1$ /MPa·K <sup>-1</sup>	-1.1244	-0.6568
$10^{3} \cdot B_{2} / MPa \cdot K^{-2}$	1.0	0.66
$10^3 \cdot \sigma^* / \text{g} \cdot \text{cm}^{-3}$	0.7	0.9
AARD / %	0.08	0.16

The experimental density values are plotted in Fig. 1 at selected temperatures together with modelling results. A quantitative comparison of the model performance for density prediction is shown in Fig. 2, where it can be observed that the best density prediction for the volatile oil is obtained through the cubic EoSs with volume translation and PC-SAFT, these three EoSs provided similar deviations in the density predictions. Regarding the gas condensate, the best density prediction was obtained through PR-VT closely followed by PC-SAFT and SRK-VT. The worst density predictions were given by the cubic EoSs with volume translation, SRK and PR. For the two HPHT reservoir fluids studied here, simple cubic EoSs with volume translation can achieve similar performance for density prediction to the advanced thermodynamic model PC-SAFT which has a more theoretical background in statistical mechanics.



Fig. 1. Experimental density data,  $\rho$ , at 298.15 K ( $\blacklozenge$ ), 373.15 K ( $\blacklozenge$ ) and 463.15 K ( $\blacksquare$ ) for the HPHT volatile oil (a, b, c) and for the HPHT gas condensate (d, e, f). Lines represent the model predictions SRK (•••), SRK-VT (- - -), PR (—), PR-VT (- • -) and PC-SAFT (——).



Fig. 2. Absolute average relative deviations of the density predictions for the HPHT volatile oil and the HPHT gas condensate through SRK (solid orange), SRK-VT (striped orange), PR (solid red), PR-VT (striped red) and PC-SAFT (solid purple).

## 3.2. Isothermal compressibility

The isothermal compressibility values ( $\kappa_T$ ) of the two reservoir fluids were obtained by differentiation from the modified Tammann-Tait fitting (Equation 10) according to the following equation:

$$\kappa_T(T,p) = \frac{1}{\rho} \left( \frac{\partial \rho}{\partial p} \right)_T$$
(13)

The isothermal compressibility values of the two fluids studied in this work are presented in Table 5, whereas the compressibility is plotted against pressure together with model predictions at selected temperatures in Fig. 3. A quantification of the model performance for prediction of isothermal compressibility is presented in Fig. 4, where it is observed that even though PC-SAFT provides a slightly better prediction, the predictions of this property for the HPHT volatile oil are very similar for the different models, with average deviations around 10%. Concerning the HPHT gas condensate, the compressibility predictions are better than the ones obtained for the volatile oil, with AARD lower than 9%. The best prediction for this fluid was obtained with PC-SAFT with an AARD of 4.4 %.

$T/\mathbf{K}$	p/MPa			
	60.00	80.00	100.00	120.00
	HPHT v	olatile oil		
298.15	1.07	0.91	0.79	0.70
323.15	1.22	1.02	0.87	0.76
348.15	1.40	1.14	0.96	0.83
358.15	1.48	1.19	1.00	0.86
373.15	1.61	1.27	1.06	0.91
423.15	2.11	1.57	1.26	1.06
432.15	2.21	1.63	1.30	1.09
463.15	2.55	1.81	1.42	1.18
	HPHT g	as conden	isate	
298.15		2.58	2.03	1.68
323.15		2.91	2.23	1.83
348.15		3.27	2.45	1.98
363.15		3.50	2.58	2.07
373.15		3.66	2.67	2.13
393.15		3.98	2.85	2.25
423.75	_	4.46	3.11	2.43
463.15		4.93	3.36	2.60

Table 5 Isothermal compressibility values<sup>a</sup>,  $10^3 \kappa_T$  (MPa<sup>-1</sup>), of the HPHT reservoir fluids.

<sup>a</sup>Relative expanded isothermal compressibility uncertainty  $U_r(\kappa_T)$  (k=2): 0.013



Fig. 3. Experimental compressibility data,  $\kappa_T$ , at 298.15 K ( $\diamond$ ), 373.15 K ( $\bigcirc$ ) and 463.15 K ( $\bigcirc$ ) for the HPHT volatile oil (a, b, c) and for the HPHT gas condensate (d, e, f). Lines represent the model predictions SRK (•••), SRK-VT (---), PR ( $\longrightarrow$ ), PR-VT (- •-) and PC-SAFT ( $\longrightarrow$ 



Fig. 4. Absolute average relative deviations of the isothermal compressibility predictions for the HPHT volatile oil and the HPHT gas condensate through SRK (solid orange), SRK-VT (striped orange), PR (solid red), PR-VT (striped red) and PC-SAFT (solid purple).

Recently, Burgess et al. [30] published experimental data and analyzed the predictive capability of PC-SAFT and SRK-VT for density and isothermal compressibility of two crude oils and their synthetic mixtures with methane at HPHT conditions (pressures up to 276 MPa and temperatures up to 523 K). They used only three pseudo-components to represent the crude oil together with a set of high pressure parameters, and their prediction of density values was within 1-4%, whereas the prediction of isothermal compressibility was within 1-10%, which is in agreement with the predictions of density and isothermal compressibility obtained in the present work for real reservoir fluids. However, when they used Gross and Sadowski [18] parameters for PC-SAFT, instead of the high pressure parameters, the compressibility prediction was within 40%, in contrast to the present work where we get a good prediction of compressibility values (maximum deviation 14.2%) with these last parameters. These authors [30] reported also poor predictions of the isothermal compressibility by using SRK with volume translation at all p,T conditions with deviations reaching in some points more than 50%, in contrast to the present work where the maximum average deviations of cubic EoSs with volume translation for isothermal compressibility are around 10%, with maximum deviations of 27%.

#### 3.3. Phase equilibrium

The measured saturation pressures ( $p^{sat}$ ) for the HPHT reservoir fluids are presented in Table 6. In the studied temperature range a maximum saturation pressure of 26.34 MPa was measured for the HPHT volatile oil at 432.12 K, whereas for the HPHT gas condensate a maximum saturation pressure of 69.55 MPa was measured at 298.15 K. These experimental data are plotted together with the different model predictions for this property in Fig. 5. The plot of the saturation pressure of the HPHT gas condensate against temperature is similar to the one previously reported by Ungerer et al. [31] for a high pressure natural gas where high dew pressures up to 70 MPa were reported and attributed to the simultaneous abundance of methane

and significant amount of heavy hydrocarbons in the sample. In this last figure, it can also be observed that the models that reproduce better the experimental data are SRK and PR for the HPHT volatile oil and PC-SAFT for the HPHT gas condensate. A quantitative analysis of the model deviations is presented in Fig. 6 where, as already observed in Fig. 5, it is shown how the simple cubic EoSs perform better than PC-SAFT in the prediction of the saturation pressure of the HPHT volatile oil, the opposite trend was found for the HPHT gas condensate where PC-SAFT provides better prediction. Among the three models studied in this work it is not possible to find one that performs equally well for the two HPHT reservoir fluids analysed in the present work without tuning of binary interaction factors,  $k_{ij}$ .

Table 6 Saturation pressure<sup>a</sup>,  $p^{\text{sat}}$ , of the HPHT reservoir fluids studied in this work.

T/K	p <sup>sat</sup> / MPa	T/K	$p^{\mathrm{sat}}$ / MPa
HPHT v	olatile oil	HPHT g	as condensate
298.06	20.92	298.15	69.55
323.13	22.65	323.15	64.41
348.00	24.20	348.15	59.57
358.10	24.64	363.16	57.32
373.08	25.24	372.93	56.17
398.02	25.94	393.18	53.92
423.09	26.29	413.20	51.88
432.12	26.34	423.79	50.87
448.00	26.32	433.15	49.81
473.04	26.03	447.97	48.27
		473.15	45.60

<sup>a</sup>Standard temperature uncertainty u(T): 0.02 K; Combined standard pressure uncertainty u(p):0.10 MPa for the HPHT volatile oil and 0.15 MPa for the HPHT gas condensate.



Fig. 5. Experimental saturation pressure (O) of (a) HPHT volatile oil and (b) HPHT gas condensate. Lines represent the model predictions SRK (•••), PR (—) and PC-SAFT (– –).



Fig. 6. Absolute average relative deviations obtained in the prediction of the experimental saturation pressure for the HPHT volatile oil and for the HPHT gas condensate through SRK (orange), PR (red) and PC-SAFT (purple).

An example of the images of the PVT cell in the two-phase region during the CME process is presented in Fig. 7 and in Fig. 8. These pictures were used to determine the liquid fraction percentage  $(100 \cdot V_{liq}/V_{tot})$ , i.e. the ratio between the liquid volume and the total volume at given *p*, *T* conditions below the saturation pressure. The yellow line in Fig. 7 indicates the level of the vapour-liquid interface in contact with the sapphire window of the PVT cell, as the experiments were performed in a cell positioning angle of 45°, the vapour-liquid interface can be also observed in contact with the head of the cell piston. The results for the constant mass expansion (CME) procedure are gathered in Tables 7 and 8 for the HPHT volatile oil and the HPHT gas condensate, respectively. Measurements of the liquid fraction percentage and the relative volume ( $V_{tot}/V_{sat}$ ),which is the ratio between the total volume and the volume at saturation pressure at given *T* are presented in these tables.



Fig. 7. Pictures showing the liquid fraction in the CME experiment of the HPHT volatile oil at 373.15 K. The yellow line represents the liquid level. PVT cell positioning angle: 45°.



Fig. 8. Pictures showing the liquid fraction in the CME experiment of the HPHT gas condensate at 393.15 K. PVT cell positioning angle: 315°.

Table 7

Liquid fraction percentage  $(100 V_{\text{liq}}/V_{\text{tot}})$  and relative volume  $(V_{\text{tot}}/V_{\text{sat}})$  for the HPHT volatile oil.<sup>a</sup>

p	%	p	%	p	%	p	%
MPa	$V_{\rm liq}/V_{\rm tot}$ $V_{\rm tot}/V_{\rm sat}$	MPa	$V_{\rm liq}/V_{\rm tot}$ $V_{\rm tot}/V_{\rm sat}$	MPa	$V_{\rm liq}/V_{\rm tot}$ $V_{\rm tot}/V_{\rm sat}$	MPa	$V_{\rm liq}/V_{\rm tot}$ $V_{\rm tot}/V_{\rm sat}$
	<i>T</i> =298.15 K	1 7	7=323.15 K		T=358.15 K	7	Г=373.15 К
51.79	0.939	53.71	0.930	51.63	0.923	52.59	0.917
51.12	0.940	52.47	0.932	51.22	0.924	51.82	0.919
49.34	0.944	50.80	0.935	49.43	0.929	50.22	0.922
47.53	0.947	49.17	0.938	47.51	0.933	48.25	0.926
45.41	0.950	46.96	0.942	45.39	0.938	46.34	0.931
43.38	0.953	45.16	0.946	43.47	0.943	44.32	0.937
41.51	0.956	42.93	0.950	41.37	0.948	42.46	0.941
39.60	0.960	41.02	0.954	39.50	0.953	40.46	0.947
37.41	0.964	39.26	0.957	37.42	0.958	38.45	0.952
35.50	0.968	37.11	0.962	35.42	0.964	36.43	0.958
33.47	0.972	35.14	0.967	33.53	0.969	34.38	0.965
31.67	0.975	33.29	0.971	31.51	0.976	32.43	0.971
29.62	0.980	31.19	0.976	29.44	0.982	30.36	0.978
27.45	0.984	29.22	0.981	27.38	0.989	28.46	0.985
25.49	0.989	27.11	0.986	25.42	0.997	28.21	0.986
23.34	0.994	25.17	0.992	25.27	0.997	26.39	0.994
21.32	0.999	23.23	0.998	25.25	0.997	26.26	0.994

21.05 21.02 20.70 20.39 19.63 18.33 16.45 13.61	97.76 94.76 90.14 81.06	0.999 0.999 1.001 1.005 1.015 1.033 1.071 1.166	23.06 22.94 22.57 22.28 21.62 20.39 18.53 15.47	99.33 97.69 94.69 89.98 80.48	0.998 0.998 1.000 1.004 1.013 1.031 1.068 1.160	24.78 24.45 23.87 22.77 20.99 17.85 14.15 10.37	99.49 97.75 94.39 88.99 79.14 66.84 51.99	0.999 1.003 1.011 1.029 1.064 1.152 1.327 1.677	26.13 26.13 25.74 25.13 24.59 23.57 21.83 18 70	99.83 97.97 94.43 89.09 78 58	$\begin{array}{c} 0.995\\ 0.995\\ 0.997\\ 1.000\\ 1.009\\ 1.026\\ 1.060\\ 1.146\end{array}$
10.65	68.76	1.356	12.13	68.30	1.344	7.58	39.63	2.202	14.91	66.40	1.317
7.77	53.56	1.735	8.85	52.77	1.712	,			10.96	51.73	1.659
5.69	40.28	2.303	6.46	39.74	2.263				8.03	38.57	2.172
,	T=423.15	Κ	1 7	-432.15	Κ	1 7	-473.15	Κ			
52.33		0.888	51.32		0.892	50.80		0.862			
51.05		0.891	50.46		0.894	50.03		0.864			
49.32		0.897	48.82		0.899	48.27		0.870			
47.29		0.902	46.80		0.905	46.30		0.877			
45.33		0.909	44.84		0.911	44.33		0.885			
43.42		0.914	42.88		0.918	42.35		0.893			
41.46		0.921	40.91		0.925	40.38		0.902			
39.39		0.929	38.93		0.933	38.34		0.912			
37.43		0.937	36.86		0.941	36.39		0.922			
35.42		0.944	34.83		0.950	34.39		0.934			
33.47		0.953	32.94		0.960	32.39		0.947			
31.43		0.963	30.84		0.971	30.41		0.961			
29.40		0.974	28.92		0.982	28.39		0.978			
29.23		0.975	28.77		0.983	28.25		0.978			
27.38		0.986	26.88		0.996	26.37		0.997			
27.27		0.98/	26.69		0.997	26.28		0.998			
27.00		0.988	20.51		0.998	20.10	00.71	0.999			
27.00		0.988	20.43		0.999	25.90	99./1	1.002			
20.77		0.990	20.28	00 00	1.001	25.04	90.40	1.009			
20.52	07 77	0.995	20.15	90.00	1.004	23.02	90.95	1.025			
25.03	91.11	1.001	23.74	90.07	1.011	25.90	03.49 72 11	1.032			
23.04	93.12 87.50	1.01/	24.93	92.55	1.027	18 19	72.14 50 50	1.122			
20.75	07.30 76.76	1.040	20.87	75 87	1.050	14.06	15 05	1.204			
16.08	63.86	1.120	17 20	63.02	1.155	14.00	+3.73	1.34/			
12 75	49 10	1.595	12.99	48.25	1.599						

<sup>a</sup>Standard temperature uncertainty u(T): 0.02 K; Standard pressure uncertainty u(p): 0.06 MPa; Maximum standard liquid fraction uncertainty u(Liquid fraction percentage): 2.1%; Maximum standard relative volume uncertainty u(Relative volume): 0.035.

Table 8

Liquid fraction percentage  $(100 \cdot V_{\text{liq}}/V_{\text{tot}})$  and relative volume  $(V_{\text{tot}}/V_{\text{sat}})$  for the HPHT gas condensate.<sup>a</sup>

р	%	р	%	р	%	р	%
MPa	$V_{ m liq}/V_{ m tot} V_{ m tot}/V_{ m sat}$	/MPa	$V_{\rm liq}/V_{\rm tot} V_{\rm tot}/V_{\rm sat}$	MPa	Vliq/Vtot Vtot/Vsat	MPa	$V_{\rm liq}/V_{\rm tot} V_{\rm tot}/V_{\rm sat}$
/	<i>T</i> =298.15 K	1	<sup>-</sup> =323.15 K	<i>T</i> =348.15 K		1	Г=363.15 K
73.43	0.987	71.36	0.974	65.34	0.971	65.31	0.958
73.08	0.989	70.97	0.976	65.21	0.972	64.93	0.959
72.12	0.991	70.08	0.979	64.29	0.976	63.99	0.964
71.17	0.994	69.10	0.982	63.29	0.981	62.96	0.969
70.19	0.997	68.11	0.985	62.34	0.985	61.98	0.973
70.02	0.998	67.13	0.988	61.33	0.990	60.99	0.979
69.85	0.998	66.10	0.992	60.31	0.995	59.99	0.984
69.72	0.998	65.12	0.996	60.22	0.996	59.00	0.990
69.57	0.999	64.93	0.997	60.10	0.996	57.97	0.995
68.48	1.002	64.83	0.997	60.02	0.997	57.89	0.996
67.30	1.006	64.80	0.997	59.90	0.997	57.73	0.997

64.47		1.014	64.69		0.998	59.80		0.998	57.63		0.997	
61.87		1.023	63.85		1.001	59.70		0.998	57.61		0.997	
59.58		1.032	63.02		1.005	59.68		0.998	57.05		1.001	
57.33	0.06	1.041	60.99		1.013	59.04		1.002	56.56		1.004	
55.12	0.17	1 0 5 0	58.96		1 022	58 46		1 005	55 27		1 012	
53 11	0.17	1.059	57 19		1.030	56.98		1.003	54.06		1 020	
51 20	0.72	1.057	55 16	0.03	1.030	55 53		1.013	52 87		1.020	
11.29	1.08	1.007	53 70	0.03	1.037	5/ 11		1.022	51 76		1.026	
20.66	2 /1	1.100	52.19	0.15	1.057	52.97	0.01	1.030	50.72	0.02	1.030	
25.50	3.41 4 75	1.143	50 77	0.54	1.057	51.67	0.01	1.030	10.72	0.03	1.044	
22.29	4.75	1.104	30.77	0.55	1.000	50.44	0.07	1.04/	49.00	0.19	1.035	
32.37	5.89	1.224	45.04	1.03	1.102	50.44	0.24	1.055	48.09	0.31	1.001	
28.71	/.12	1.285	41.40	2.74	1.139	49.28	0.41	1.064	45.18	1.03	1.094	
25.95	8.25	1.346	38.01	3.67	1.176	45.28	1.25	1.098	42.21	1.61	1.12/	
20.44	9./1	1.551	35.19	4.66	1.213	41.94	2.11	1.133	39.67	2.43	1.101	
15.48	9.28	1.967	31.82	5.75	1.270	39.15	2.81	1.16/	37.48	2.95	1.194	
			29.17	6.37	1.327	36.74	3.38	1.203	34.67	3.64	1.245	
			23.44	7.92	1.520	33.73	4.42	1.256	32.35	4.26	1.296	
			17.78	7.66	1.909	31.32	5.04	1.309	26.84	5.42	1.468	
			13.60	6.54	2.494	25.71	6.20	1.488	20.68	5.65	1.813	
			10.56	5.02	3.277	19.68	6.48	1.849	15.84	4.90	2.334	
			8.31	3.79	4.256	15.05	5.51	2.394	12.25	3.92	3.030	
						11.64	4.30	3.121	9.59	3.02	3.900	
						9.13	3.39	4.031				_
T=	=393.15	Κ	T=	=423.75	Κ	T	=448.15	Κ	T	=473.15	Κ	
60.27		0.957	59.24		0.933	54.22		0.941	54.18		0.905	
59.63		0.960	56.51		0.952	53.84		0.944	51.22		0.934	
58.64		0.966	55.51		0.960	52.85		0.953	50.24		0.944	
57.64		0.973	54.51		0.968	51.87		0.962	49.25		0.955	
56.63		0.980	53.50		0.976	50.88		0.972	48.23		0.966	
55.66		0.987	52.51		0.985	49.88		0.982	47.25		0.978	
54.64		0.994	51.49		0.994	48.88		0.993	46.24		0.991	
54.56		0.995	51.44		0.995	48.80		0.994	46.08		0.993	
54.47		0.995	51.33		0.996	48.70		0.995	46.03		0.994	
54.35		0.996	51.23		0.997	48.60		0.996	45.97		0.995	
54.25		0.997	51.11		0.998	48.50		0.997	45.87		0.996	
54.14		0.998	51.02		0.999	48.45		0.998	45.83		0.997	
54.03		0.999	50.98		0.999	48.41		0.999	45.73		0.998	
53.95		1.000	50.93		1.000	48.35		0.999	45.68		0.999	
53.90		1.000	50.90		1.000	48.31		1.000	45.63		0.999	
53.87		1.000	50.83		1.001	48.25		1.000	45.58		1.000	
53.82		1.001	50.78		1.001	48.25		1.000	45.53		1.001	
53.75		1.001	50.73		1.002	48.03		1.003	45.34		1.003	
53.65		1.002	50.69		1.002	47.80		1.005	45.17		1.005	
53.61		1.003	50.69		1.002	47.28		1.012	44.74		1.011	
53.23		1.006	50.43		1.005	46.73		1.018	44.33		1.017	
52.85		1 009	50.15		1 008	46.22		1 025	43.92		1 023	
51.00		1.016	49 47		1.015	45 72		1.020	43 51		1 029	
51.01		1.010	48 76		1.012	45.72		1.031	43 12		1.035	
50.20		1.021	48 16		1.022	44 75	0.00	1.050	42 75		1.033	
<u>49</u> 38		1.031	47 53		1.025	44 29	0.00	1.011	42.75	0.00	1.047	
48 57	0.01	1.037	46.94		1.033	42 52	0.00	1.031	40.93	0.00	1.071	
17.87	0.01	1.040	16 32		1.042	10 01	0.00	1 103	30.60	0.00	1.001	
47.02 47.00	0.00	1.054	45 70	0.02	1.049	30.74	0.22	1 1 20	38 38	0.09	1 1 1 1 0	
<u>11.07</u> <u>11</u> 12	0.15	1.002	43.65	0.02	1.050	38 12	0.59	1.129	37.72	0.20	1.119	
12 US	1.04	1 1 2 2	A1 76	0.20	1 1 1 2	36 21	0.57	1.133	35.66	0.50	1.143	
42.08	1.04	1.123	41.70	0.31	1.113	21 20	0.03	1.194	24 22	0.40	1.1/9	
40.02	1.30	1.1.04	40.03	0.79	1.141	20.20	1.0/	1.234	20.20	0.08	1.213	
25.1/ 25.70	1.0/	1.100	26 11	1.05	1.109	20.29	1.39	1.303	50.29 21 01	1.03	1.555	
33./ð	2.40	1.232	24.50	1.41	1.212	24.43	2.08	1.030	24.81	1.40	1.3//	
33./1	2.80	1.2/9	34.38	1./0	1.200	19.18	2.09	2.028	19./0	1.32	1.941	
28.56	5.81	1.436	29.81	2.57	1.398	15.02	1.89	2.360	15.96	1.38	2.367	
22.36	4.19	1.752	23.71	2.86	1.685	11.85	1.54	3.225	13.14	1.25	2.854	

17.23	3.85	2.228	18.43	2.82	2.118
13.35	3.12	2.865	14.35	2.41	2.696
10.47	2.47	3.661	11.28	1.90	3.420

<sup>a</sup>Standard temperature uncertainty u(T): 0.02 K; Standard pressure uncertainty u(p): 0.06 MPa; Maximum standard liquid fraction uncertainty u(Liquid fraction percentage): 1.6%; Maximum standard relative volume uncertainty u(Relative volume): 0.063.

The experimental values of the relative volume and the liquid fraction are plotted as a function of pressure in Fig. 9 for the HPHT volatile oil and the HPHT gas condensate at selected temperatures. The model predictions are also depicted in this figure. As regards the relative volume at the selected temperatures, it is not possible to observe a clear transition from the single-phase system to the two-phase system in the relative volume curves vs. pressure due to the small contrast between gas and liquid compressibilities associated to the phase transition. Maximum relative expansions of 2.3 and 4.3 were measured for the HPHT volatile oil and the HPHT gas condensate, respectively. A good prediction of this property was found through the models studied in this work.

As regards the liquid fraction of the HPHT volatile oil in Fig. 9 (b), the experimental data show the typical liquid fraction behaviour under expansion corresponding to a bubble point, i.e. a liquid fraction of 100% is observed at the saturation pressure, and the value of this property decreases with a further decrease in the pressure below the bubble point. The best liquid fraction predictions for the HPHT volatile oil are obtained with the cubic EoSs, whereas PC-SAFT can reproduce in general the slope of liquid fraction decrease with pressure and the deviations in the prediction are mainly caused by the poor prediction of the saturation pressure of the HPHT volatile oil through this EoS. Finally, concerning the liquid fraction of the HPHT gas condensate, it can be observed in Fig. 9 (d) that the experimental data correspond to the typical behaviour of a retrograde gas starting with a liquid fraction of 0% at the saturation pressure which increases with the pressure decrease and passes through a maximum, after which the liquid fraction decreases with a further pressure decrease. An interesting feature from the experimental liquid fraction values is that negligible amount of condensate happened under

a considerable pressure decrease below the dew point, i.e. a big pressure decrease was needed in order to obtain a measurable amount of condensate, this phenomena led to a tail in the liquid fraction vs. pressure curves. The different EoSs present difficulties capturing this tail-like behaviour, as previously stated by Whitson et al. [5]. This tail becomes longer when the temperature decreases, for instance at 298.15 K, the measured dew point was 69.55 MPa, but the pressure needed to be decreased down to 57.33 MPa in order to get a measurable amount of condensate. This tail-like behaviour of the liquid drop-out was previously reported by Robinson [32], who reported a pressure decrease of 15.8 MPa below the dew point to get a 0.5% of liquid fraction for a gas condensate at 283.15 K. However, Ahmed [33] reported that the small build-up of condensate has previously been attributed to sample contamination by hydraulic fluids from various sources during drilling, production and sampling, even though he states that it could also be a feature of real reservoir fluids [33]. Our compositional analysis of the gas condensate (Table 1) does not show any obvious sample contamination. The experimental liquid fraction increases when temperature decreases due to a higher drop out of the heavier compounds at lower temperatures. A maximum liquid fraction of 9.7 % was found at 298.15 K, this value is over predicted by the models used in this work, it can be observed in Fig. 9 (d) that the model prediction for the liquid fraction of the gas condensate becomes poorer when the temperature decreases. Barnum et al. [34] gathered data for gas condensates from 17 fields, concluding that the condensation of the liquid hydrocarbons can affect the gas productivity to great extent in low productivity reservoirs. Also, Afidick et al. [35] studied near wellbore condensate accumulation in a gas condensate from the Arun field (Indonesia), this gas condensate was a lean-gas with a maximum liquid drop-out of 1.1% and they found that despite of being a lean-gas, some of the well productivities were reduced by 50%. These studies [34, 35] indicate that the amount of retrograde condensate of the HPHT gas condensate studied in the present work could affect productivity of this reservoir fluid.



Fig. 9. Relative volume  $(V_{tot}/V_{sat})$  and liquid fraction percentage  $(100 \cdot V_{liq}/V_{tot})$  of the HPHT volatile oil (a, b) and the HPHT gas condensate (c, d) at 298.15 K ( $\diamond$ ), 373.15 K ( $\bullet$ ), 393.15 K ( $\bullet$ ) and 473.15 K ( $\bullet$ ). Lines represent the model predictions SRK (••••), SRK-VT (- - -), PR (—), PR-VT (- • -) and PC-SAFT (— —).

A quantitative representation of the models performance for both the prediction of the relative volume and the liquid fraction percentage is given in Figs. 10 and 11, respectively. In Fig. 10 it can be observed that the relative volume predictions are generally good. For the HPHT volatile oil predictions are especially good through PR, whereas for the HPHT gas condensate the best predictions were obtained with PC-SAFT, closely followed by SRK. For the cubic EoSs the volume translation had negligible effect in the calculation of this property. In Fig. 11 it can be observed how the prediction of the liquid fraction percentage of the HPHT volatile oil is better through the cubic EoSs, as concerns the gas condensate the best prediction is obtained with PC-SAFT.



Fig. 10. Absolute average deviation for the prediction of the relative volume ( $V_{tot}/V_{sat}$ ) of the HPHT reservoir fluids through SRK (solid orange), SRK-VT (striped orange), PR (solid red), PR-VT (striped red) and PC-SAFT (solid purple).



Fig. 11. Absolute average deviation for the prediction of the liquid fraction percentage  $(100 \cdot V_{\text{liq}}/V_{\text{tot}})$  of the HPHT reservoir fluids through SRK (solid orange), SRK-VT (striped orange), PR (solid red), PR-VT (striped red) and PC-SAFT (solid purple).

Finally, the overall deviations for various properties of the two HPHT reservoir fluids calculated by the five models are summarized in Table 9. The smallest deviations and those "sufficiently" close to the smallest deviations are marked in bold letters. The definition of "sufficiently" close is a bit arbitrary. For density, compressibility and saturation pressure, we select values differing from the smallest value by less than 0.5. For relative volume and liquid

fraction, we marked essentially just the smallest deviations. Note that the deviations for density, compressibility and saturation pressures are relative and those for relative volume and liquid fraction are absolute. The table provides an overview of the performance of the five models. It is again difficult to single out a model that performs always the best for the studied properties although PC-SAFT tends to give smaller deviations as indicated by the number of bold letters. Among these properties, the performance of saturation pressure prediction is case-dependent and it is difficult to make a general conclusion which model is better. This is somewhat in agreement with our previous observations [3, 7, 9, 22]. For reservoir fluids, the prediction is also affected by the characterization procedure and uncertainties in interaction parameters and other characterized model parameters. It is therefore more difficult to expect that a model is consistently much better than others in the saturation pressure calculation. It should be noted that the deviations in liquid fraction are closely related to the deviations in saturation pressure. Although the vapor- and liquid-phase densities are also used in the calculation of liquid fractions, the correct location of the saturation pressure is apparently the most influencing factor. Our comparison here is for predictive calculation. For models tuned against saturation pressure, the effect of density modeling may become more visible. The calculation of relative volumes is generally insensitive to the models partly because the two-phase total volume is controlled mainly by the vapor phase. The single-phase density and compressibility are related to the volumetric performance of EoS models. Volume translation is obviously needed to improve the density calculation for SRK and PR. But the effect of volume translation on compressibility is not as straightforward. PR-VT, SRK-VT and PC-SAFT performs almost equally well for the volumetric properties with PC-SAFT seemingly slightly better in compressibility. It should be noted that PR-VT and SRK-VT have used the experimental density of  $C_7^+$  through characterization.

It would be useful if the performance of an EoS can be linked to the methane content. The methane mole fraction for the volatile oil is around 49% and that for the gas condensate is around 83%. The density prediction improves a lot for SRK, deteriorates clearly for PR, and deteriorates slightly for PC-SAFT. The changes for SRK and PR can be explained by the fact that SRK describes methane gas density better than PR. But the performance change in compressibility is more difficult to explain, and so are the changes for the volume translated versions of SRK and PR. Regarding the saturation pressure, the prediction worsens for SRK and PR and improves for PC-SAFT when the methane fraction increases. However, we note that the above observations cannot be generalized even for the densities from SRK and PR. In our previous studies for well-defined mixtures containing methane [7-9], the trends with the methane content are not always in agreement with the observations in this study. For instance, although the density prediction by SRK improves with the methane content for two sevencomponent synthetic mixtures [8], the prediction by PR also improves with the methane content.

One consideration to involve PC-SAFT in the comparison is to see whether this advanced EoS shows some particular advantages over classical cubic models in describing some challenging reservoir fluids systems. PC-SAFT is apparently more complex than SRK and PR. But its robustness is comparable to SRK and PR in practical phase equilibrium calculations. This is especially the case if the Wertheim association term is not used, as in this study for hydrocarbon systems. The potential advantage for PC-SAFT without the association term comes from its more rigorous repulsive and attractive terms. Our previous comparative study [22] has shown that PC-SAFT is clearly better than cubics in describing volumetric properties for pure components but similar to cubics in binary and multicomponent VLE calculations. After  $C_7^+$  characterization [3, 22], the advantage of using PC-SAFT in PVT calculation becomes less clear. This can be attributed to many reasons, such as the uncertainty associated with the  $C_7^+$ 

characterization, the sensitivity of the model parameters, the quality of the PVT data, and the type of fluids used in comparison. Regarding the sensitivity, our previous study [22] shows that PC-SAFT is more sensitive to its model parameters. This can make PC-SAFT more susceptible to uncertainties in reservoir fluids. In practice, tuning of a PVT model is usually performed to anchor the model to some experimental data, which can somehow reduce the influence from the uncertainties. For further application of PC-SAFT in reservoir fluids modelling, one major challenge is to identify situations where PC-SAFT can clearly outperform the classical models. Answering the challenge calls for both theoretical analyses indicating where PC-SAFT can be more advantageous and more experimental data for challenging systems like HPHT fluids to test the hypotheses.

Table 9

Summary of the overall deviations for different properties of the two HPHT reservoir fluids.

	SRK	SRK-VT	PR	PR-VT	PC-SAFT
	Volatile Oil				
Single-phase density	14	1.1	4.5	1.3	1.3
Single-phase compressibility	10.8	11.3	11.5	10.3	9.7
Saturation pressure	1.6		2.2		6.1
Relative volume	0.16	0.15	0.008	0.005	0.02
Liquid fraction	1.3	2.4	3.3	2.5	5.3
	Gas condensate				
Single-phase density	2.7	2.4	6.5	0.6	2.0
Single-phase compressibility	5.6	8.6	7.6	5.7	4.4
Saturation pressure	14		23		3.5
Relative volume	0.06	0.06	0.12	0.13	0.05
Liquid fraction	2.7	2.1	1.9	1.5	1.2

## 4. Conclusions

As a continuation of our previous studies on HPHT fluids, we extend our measurement to real reservoir fluids including one HPHT volatile oil and one HPHT gas condensate. We systematically measured the single-phase density, single-phase compressibility, saturation pressure, and liquid fraction in the two-phase region of these two fluids. The measurement was carried out in a wide temperature and pressure range, with maximum temperature and pressure

reaching 473.15 K and 140 MPa, respectively. Since it is difficult to mimic completely the phase behavior of real reservoir fluids with well-defined synthetic mixtures, the measured data is valuable for developing and evaluating relevant models for HPHT applications.

We also compared experimental results with model predictions by common cubic EoSs (SRK and PR), their volume translated versions (SRK-VT and PR-VT), and the more advanced PC-SAFT model. The saturation pressure calculation is case-dependent: the simple SRK and PR models perform better than PC-SAFT for the volatile oil whereas PC-SAFT performs better for the gas condensate. It is difficult to generalize this to other fluids. The HPHT gas condensate shows a minute amount of liquid drop-out in a considerable pressure range below the dew point. None of the current models can capture this long tail-like liquid drop-out curve. For volumetric properties like density and compressibility, PC-SAFT is clearly better than SRK and PR especially in density but PC-SAFT, PR-VT and SRK-VT perform similarly in density calculation for the two fluids investigated here. PC-SAFT seems to be slightly better in compressibility prediction but the difference is not large. Overall, the findings from the model comparison in this study are in certain agreement with the findings from our previous studies [7-9]. One can generally expect that SRK, PR and PC-SAFT perform similarly in equilibrium calculation whereas PC-SAFT may be slightly better in volumetric properties. However, the difference between models seems to be blurred for ill-defined reservoir fluids. The performance of models can well vary with the fluid type. The two fluids investigated here have relatively light  $C_7^+$  and their overall compositions fall into the range of reservoir fluids with medium gas oil ratios. It is recommended to study experimentally and through modeling other types of HPHT reservoir fluids, e.g., heavier oil systems or even lighter gas systems, in order to give a more complete picture of the performance of various models in modeling HPHT reservoir fluids.

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