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**Citation**: Rodriguez, C., Vidal, A., Koukouvinis, P. ORCID: 0000-0002-3945-3707, Gavaises, M. ORCID: 0000-0003-0874-8534 and McHugh, M. A. (2018). Simulation of transcritical fluid jets using the PC-SAFT EoS. Journal of Computational Physics, 374, doi: 10.1016/j.jcp.2018.07.030

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Link to published version: http://dx.doi.org/10.1016/j.jcp.2018.07.030

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### Accepted Manuscript

Simulation of transcritical fluid jets using the PC-SAFT EoS

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 PII:
 S0021-9991(18)30491-1

 DOI:
 https://doi.org/10.1016/j.jcp.2018.07.030

 Reference:
 YJCPH 8155

To appear in: Journal of Computational Physics

Received date:1 February 2018Revised date:11 July 2018Accepted date:12 July 2018



Please cite this article in press as: C. Rodriguez et al., Simulation of transcritical fluid jets using the PC-SAFT EoS, J. Comput. Phys. (2018), https://doi.org/10.1016/j.jcp.2018.07.030

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### Highlights

- A numerical framework to simulate transcritical and supercritical flows utilising the compressible form of the Navier–Stokes equations coupled with the Perturbed Chain Statistical Associating Fluid Theory (PC-SAFT) equation of state (EoS) is presented.
- Both conservative and quasi-conservative formulations have been tested.
- Advection test cases and shock tube problems are included to show the overall performance of the developed framework.
- Two-dimensional simulations of nitrogen and dodecane jets are presented to demonstrate the multidimensional capability of the developed model.

- 1 Simulation of transcritical fluid jets using the PC-SAFT EoS 2 3 C. Rodriguez<sup>a,\*</sup>, A. Vidal<sup>a</sup>, P. Koukouvinis<sup>a</sup>, M. Gavaises<sup>a</sup>, M. A. McHugh<sup>b</sup> 4 5 6 <sup>a</sup> School of Mathematics, Computer Science & Engineering, Department of Mechanical Engineering & Aeronautics, City University London, Northampton 7 Square EC1V 0HB, United Kingdom 8 <sup>b</sup> Department of Chemical and Life Science Engineering, 601 West Main Street, 9 Richmond, VA 23284, USA 10 11 \**Corresponding author: Carlos.Rodriguez@city.ac.uk* 12 13 14 Abstract The present paper describes a numerical framework to simulate transcritical and supercritical 15 16 flows utilising the compressible form of the Navier-Stokes equations coupled with the 17 Perturbed Chain Statistical Associating Fluid Theory (PC-SAFT) equation of state (EoS); 18 both conservative and quasi-conservative formulations have been tested. This molecular 19 model is an alternative to cubic EoS which show low accuracy computing the thermodynamic 20 properties of hydrocarbons at temperatures typical for high pressure injection systems. Liquid 21 density, compressibility, speed of sound, vapour pressures and density derivatives are 22 calculated with more precision when compared to cubic EoS. Advection test cases and shock 23 tube problems are included to show the overall performance of the developed framework 24 employing both formulations. Additionally, two-dimensional simulations of nitrogen and 25 dodecane jets are presented to demonstrate the multidimensional capability of the developed 26 model. 27 Keywords: Supercritical, transcritical, PC-SAFT EoS, double-flux model, Riemann problem 28 29 30 Nomenclature 31 32 List of abbreviations 33 Average Absolute Deviation AAD 34 CFD **Computational Fluid Dynamics** 35 CFL Courant–Friedrichs–Lewy 36 **ENO** Essentially Non-Oscillatory 37 Equation of State EoS 38 FC Fully Conservative 39 HLLC Harten-Lax-van Leer-Contact 40 LES Large Eddy Simulation 41 PR Peng-Robinson 42 PC-SAFT Perturbed Chain Statistical Associating Fluid Theory 43 OC Quasi-Conservative 44 RK2 Second-order Runge-Kutta
- 45 SRK Soave-Redlich-Kwong

46	SSP-RI	C3 Third-order strong-stability-preserving Runge–Kutta
47	TVD	Total Variation Diminishing
48	WENO	Weighted Essentially Non-Oscillatory
49		
50	List of .	Symbols
51	$\tilde{a}^{rs}$	Reduced Helmholtz free energy [-]
52	С	Sound speed [m s <sup>-1</sup> ]
53	d	Temperature-dependent segment diameter [Å]
54	g	Radial distribution function [-]
55	Ι	Integrals of the perturbation theory [-]
56	$k_{\scriptscriptstyle B}$	Boltzmann constant [J/K]
57	т	Number of segments per chain [-]
58	$\overline{m}$	Mean segment number in the system [-]
59	р	Pressure [Pa]
60	R	Gas constant [J mol <sup>-1</sup> K <sup>-1</sup> ]
61	Т	Temperature [K]
62	$X_i$	Mole fraction of component i [-]
63	Ż	Compressibility factor [-]
64	L U	Conservative variable vector
65	F	x-convective flux vector
66	G	v-convective flux vector
67	Fv	x-diffusive flux vector
68	G <sub>v</sub>	y-diffusive flux vector
69		
70	Greek	Letters
71	${\cal E}$	Depth of pair potential [J]
72	$\eta$	Packing fraction [-]
73	ρ	Density [kg/m <sup>3</sup> ]
74	$ ho_{\scriptscriptstyle m}$	Total number density of molecules [1/Å <sup>3</sup> ]
75	$\sigma_{\!\scriptscriptstyle d}$	Segment diameter [Å]
76	u	
77	Superse	cripts
78	disp	Contribution due to dispersive attraction
79	hc	Residual contribution of hard-chain system
80	hs	Residual contribution of hard-sphere system
81	id	Ideal gas contribution
82		
83		
84		
85		
86		
87		
88		

#### 89 **1. Introduction**

90 Transcritical and supercritical states occur in modern combustion engines that operate at 91 pressures higher than the critical pressure of the fuels utilised. In Diesel engines for example, 92 the liquid fuel is injected into air at pressure and temperature conditions higher than the 93 critical point of the fuel [1]. The liquid injection temperature is lower than the fuel critical 94 temperature but as the liquid is heated, it may reach supercritical temperature before full 95 vaporisation. This is known as a transcritical injection. Similarly, in liquid rocket engines, 96 cryogenic propellants are injected into chambers under conditions that exceed the critical 97 pressure and temperature of the propellants.

98 A single-species fluid or a mixture reaches a supercritical state when the pressure and 99 temperature surpass its critical properties. In the critical region, repulsive interactions 100 overcome the surface tension resulting in the existence of a single-phase that exhibits properties of both gases and liquids (e.g., gas-like diffusivity and liquid-like density). A 101 102 diffuse interface method is commonly employed in supercritical and transcritical jet 103 simulations to capture the properties of the flow [2]-[4]. Several difficulties should be 104 overcome for simulating the mixing of the jets using a diffused interface [5]. The presence of 105 large density gradients between the liquid-like and the gas-like regions, the need of using a 106 real-fluid EoS, or the spurious pressure oscillations generated in conservative schemes are the 107 main challenges.

108 High order reconstruction methods are usually applied to capture the large density 109 gradients. The authors of [6] performed a two-dimensional large-eddy simulation (LES) of 110 supercritical mixing and combustion employing a fourth-order flux-differencing scheme and a total-variation-diminishing (TVD) scheme in the spatial discretization. In [7] a fourth-order 111 112 central differencing scheme with fourth-order scalar dissipation was applied in order to 113 stabilize the simulation of a cryogenic fluid injection and mixing under supercritical 114 conditions. Moreover, [8] employed an eighth-order finite differencing scheme to simulate 115 homogeneous isotropic turbulence under supercritical pressure conditions, while in [9] a 116 density-based sensor was utilized, which switches between a second-order ENO (Essentially 117 non-oscillatory) and a first-order scheme to suppress oscillations. In the present study a fifth-118 order WENO (Weighted Essentially Non-Oscillatory) scheme [10] is applied due to its high 119 order accuracy and non-oscillatory behaviour.

120 Cubic EoS models like PR (Peng-Robinson) [11] and SRK (Soave-Redlich-Kwong) EoS 121 [12] are usually used in supercritical and transcritical simulations. In the studies reported in 122 [4], [13]–[15] the SRK EoS was employed in order to close the Navier Stokes equations and 123 compute the fluid properties under supercritical and transcritical conditions. Moreover, the 124 works reported in [3], [9], [16], [17] modeled the non-ideal fluid behavior applying the PR 125 EoS. However, cubic models commonly present low accuracy computing the thermodynamic 126 properties of hydrocarbons at temperatures typical for injection systems [2]. To overcome 127 this, the Statistical Association Fluid Theory Equation of State (SAFT EoS) can be employed. 128 This molecular model is based on the perturbation theory, as extensively studied in [18]–[21] 129 by Wertheim. The authors of [22], [23] developed this EoS by applying Wertheim's theory 130 and extending it to mixtures. Figure 1 shows a schematic representation of the terms 131 considered in the SAFT equation. Each molecule is represented by segments of equal size, 132 assumed to form a repulsive, hard sphere reference fluid. Next, the attractive interactions 133 between segments are added to the model. The segment-segment energy needed to form a 134 chain between the hard-sphere fluid segments is included and, if the segments exhibit 135 associative interactions, such as hydrogen bonding, a term for this interaction is also added.

Among the different variants of the SAFT model, the PC-SAFT is the one implemented here. In this model, hard chains are used as the reference fluid instead of hard spheres. While the SAFT EoS computes segment-segment attractive interactions, the PC-SAFT EoS computes chain-chain interactions, which improves the thermodynamic description of chainlike, fluid mixtures [24].

141 Several papers have been published pointing out the advantages of the SAFT models 142 with respect to the cubic EoS commonly used in CFD simulations. For example, [25] 143 describes how the PC-SAFT model is better than cubic EoS for predicting gas phase 144 compressibility factors and oil phase compressibilities. In [26] the superiority of the PC-145 SAFT performance is demonstrated relative to the Cubic Plus Association (CPA) EoS in 146 correlating second order derivative properties, like speed of sound, dP/dV and dP/dT 147 derivatives, heat capacities and the Joule-Thomson coefficient in the alkanes investigated. 148 Similarly, [27] points out the superiority of the SAFT-BACK EoS over the PR EOS, 149 particularly at high-density conditions, for computing second order derivative properties such 150 as sound velocity and isobaric and isochoric properties. The study of [28] states that cubic 151 EoS predict a linear increase of the Z factor (compressibility factor) with pressure, while the 152 PC-SAFT EoS shows a better pressure dependence. Finally, [29] shows how the sPC-SAFT 153 (simplified PC-SAFT) is more precise than SRK and CPA to compute the speed of sound of 154 normal alkanes and methanol.

155 If a fully conservative (FC) formulation is employed along with a real-fluid EoS, 156 spurious pressure oscillations may appear; the work of [4] has related this problem to 157 computational stability issues, turbulence, and acoustics accuracy loss. The same authors of 158 [4] developed a quasi-conservative (QC) scheme solving a pressure evolution equation 159 instead of the energy conservation equation, while [30] developed a quasi-conservative 160 framework where the artificial dissipation terms in the mass, momentum and energy 161 equations are related and the pressure differential is considered to be zero. In [31] the double 162 flux model was developed to avoid spurious pressure oscillations in simulations of 163 compressible multicomponent flows that employ a perfect gas EoS; [32] extended it to 164 reactive flows; and finally, [3], [17], [33] extended the double flux model to real-fluids and 165 transcritical conditions. However, recently it has been reported that the large energy 166 conservation error in quasi-conservative schemes maybe produce an unphysical quick heat-up 167 of the jet [2].



168 Figure 1. Schematic representation of the attractive and repulsive contributions of the SAFT EoS

169 170

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and the PC-SAFT EoS [24]
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171 The novelty of the approach described here is the coupling of the PC-SAFT EoS with 172 the Navier-Stokes equations, which it is not present in the literature. During the last years 173 conservative and quasi-conservative formulations have been employed in the simulation of

174 supercritical and transcritical jets. For this reason, two codes have been developed employing 175 both schemes: the conservative and the so-called quasi-conservative approach, where the 176 double flux model of [3], [17], [33] is utilized. The aim of this research is not to solve the 177 spurious pressure oscillations characteristic of FC schemes when real-fluid EoS are applied or 178 the energy conservation error of QC formulations but to present how the Navier-Stokes 179 equations can be closed with the PC-SAFT in both scenarios. Advection test cases and shock 180 tube problems are included to show the overall performance of the developed framework 181 using both formulations. Moreover, two-dimensional simulations of nitrogen and dodecane 182 jets are presented to demonstrate the capability of the code to predict fluid mixing.

183 184

#### 2. Numerical Method

The Navier-Stokes equations for a non-reacting multi-component mixture containing Nspecies in a x-y 2D Cartesian system are given by:

187  
188 
$$\frac{\partial \mathbf{U}}{\partial t} + \frac{\partial \mathbf{F}}{\partial x} + \frac{\partial \mathbf{G}}{\partial y} = \frac{\partial \mathbf{F}_{v}}{\partial x} + \frac{\partial \mathbf{G}_{v}}{\partial y}$$
(1)  
189

190 The vectors of eq. 1 are:

191 
$$\mathbf{U} = \begin{bmatrix} \rho \mathbf{Y}_{1} \\ \vdots \\ \rho \mathbf{Y}_{N} \\ \rho u \\ \rho v \\ \rho E \end{bmatrix}, \mathbf{F} = \begin{bmatrix} \rho u \mathbf{Y}_{I} \\ \vdots \\ \rho u \mathbf{Y}_{N} \\ \rho u^{2} + p \\ \rho u v \\ (\rho E + p)u \end{bmatrix}, \mathbf{G} = \begin{bmatrix} \rho v \mathbf{Y}_{I} \\ \vdots \\ \rho v \mathbf{Y}_{N} \\ \rho v u \\ \rho v^{2} + p \\ (\rho E + p)v \end{bmatrix}, \mathbf{F}_{v} = \begin{bmatrix} \mathbf{J}_{x,N} \\ \vdots \\ \mathbf{J}_{x,N} \\ \sigma_{xx} \\ \sigma_{xy} \\ u \sigma_{xx} + v \sigma_{xy} - q_{x} \end{bmatrix}, (2)$$
192 
$$\mathbf{G}_{v} = \begin{bmatrix} \mathbf{J}_{y,1} \\ \vdots \\ \mathbf{J}_{y,N} \\ \sigma_{yx} \\ \sigma_{yy} \\ u \sigma_{yx} + v \sigma_{yy} - q_{y} \end{bmatrix}$$

193

where  $\rho$  is the fluid density, *u* and *v* are the velocity components, p is the pressure, *E* is the total energy,  $J_i$  is the mass diffusion flux of species *i*,  $\sigma$  is the deviatoric stress tensor and *q* is the diffusion heat flux vector.

197 The finite volume method has been applied in this work for obtaining a numerical 198 solution to the above equations. The PC-SAFT EoS is implemented to simulate supercritical 199 and transcritical states. The developed numerical framework considers a condition of 200 thermodynamic equilibrium in each cell. Phase separations or metastable thermodynamic 201 states are beyond the scope of this research and are not considered.

### 202

#### 203 2.1 Formulations

204 Since PC-SAFT EoS is rarely used in CFD simulations, two codes have been 205 developed employing different formulations (conservative and quasi-conservative) to 206 determine which one is more appropriate for the simulation of transcritical and supercritical 207 fluid jets.

#### 208 2.1.1 Conservative formulation

209 Operator splitting [34] is adopted to divide the physical processes into hyperbolic and 210 parabolic sub-steps. The global time step is computed using the CFL (Courant-Friedrichs-211 Lewy) criterion of the hyperbolic operator.

212

#### 213 Hyperbolic sub-step

214 The HLLC (Harten-Lax-van Leer-Contact) solver is used to solve the Riemann 215 problem. The conservative variables are interpolated onto the cell faces using a fifth-order 216 WENO scheme [10] due to its high order accuracy and non-oscillatory behaviour. TVD 217 (Total Variation Diminishing) limiters [34] are applied to avoid oscillations near 218 discontinuities. Time integration is performed using a SSP-RK3 (third-order strong-stability-219 preserving Runge-Kutta) method [35].

220

#### 221 Parabolic sub-step

222 The method developed in [36] is applied to calculate the values of the dynamic 223 viscosity and thermal conductivity of the mixture. The model of [37] is implemented to 224 compute the diffusion coefficient. A RK2 (second-order Runge-Kutta) scheme is employed 225 to perform the time integration of this sub-step. Linear interpolation is performed for 226 computing the conservative variables, enthalpy and temperature on faces from cell centres.

227

#### 228 2.1.2 Quasi-conservative formulation

229 The physical processes are divided into hyperbolic and parabolic sub-steps using 230 operator splitting as well [34]. The CFL criterion of the hyperbolic operator is used to 231 compute the global time step.

232

#### 233 Hyperbolic sub-step

234 The double flux model of [3], [17], [33] has been implemented. The HLLC solver is 235 used to solve the Riemann problem. In the one-dimensional cases presented, the primitive 236 variables are interpolated onto the cell faces using a fifth-order WENO scheme [10]. In the 237 two-dimensional cases, a sensor that compares the value of the density in the faces and the 238 centre of the cells is employed to determine in which regions a more dissipative scheme must 239 be applied [3]. If the sensor is activated, TVD limiters [34] are employed. The solution is 240 then blended with a first-order scheme (90% WENO). Time integration is performed using a 241 SSP-RK3 method [35].

242

244

245

243 The following steps were followed to implement the double flux model [3], [17], [33]:

1) In each cell are stored the values of  $\gamma^*$  (eq.3) and  $e_0^*$  (eq.4).

$$\gamma^* = \frac{\rho c^2}{p} \tag{3}$$

(4)

246

 $e_0^* = e - \frac{pv}{\gamma^* - 1}$ 247 where p is the pressure, c is the sound speed, e is the internal energy and v is the 248 specific volume.

250 2) Runge-Kutta scheme

• Step 1: The fluxes at the faces are computed using the primitive variables. The total energy in the left (L) and right (R) states are computed using eq.5.

$$(\rho E)_{L,R}^{n} = \frac{p_{L,R}^{n}}{\gamma_{j}^{*,n} - 1} + \rho_{L,R}^{n} e_{0,j}^{*,n} + \frac{1}{2} \rho_{L,R}^{n} \mathbf{u}_{L,R}^{n} \cdot \mathbf{u}_{L,R}^{n}$$
(5)

- Step 2: Update conservative variables using the RK scheme
- Step 3: Update primitive variables (using the double flux model to compute the pressure).
- 257

253

3) Update total energy: The total energy is updated from primitive variables based on the EoS
(eq.6). Only at this point the PC-SAFT EoS is used to compute the internal energy, sound
speed, temperature and enthalpy.

(6)

261 
$$\rho E = \rho e + \frac{1}{2} \rho \mathbf{u} \cdot \mathbf{u}$$

262

### 263 Parabolic sub-step

The diffusion fluxes are calculated conservatively in the same way that is explained in the conservative formulation.

266

#### 267 2.2 PC-SAFT EoS subroutine

A different subroutine has been developed for each formulation because of the differentinputs of the EoS subroutine.

270

### 271 *Conservative formulation*

272 The thermodynamic variables computed in the CFD code by the PC-SAFT EoS are 273 the temperature, pressure, sound speed and enthalpy. The algorithm inputs are the density, 274 internal energy, molar fractions and three pure component parameters per component 275 (number of segments per chain, energy parameter of each component and segment diameter), 276 see Table 1. The density and the internal energy are obtained from the conservative variables 277 of the CFD code. The molar fractions are computed using the mass fractions employed in the 278 continuity equations and the molar weights of the components. The pure component 279 parameters are specified in the initialization of the simulation. A detailed description of the 280 PC-SAFT EoS can be found in the Appendix A.

281 The Newton-Raphson method is employed to compute the temperature that is needed 282 to calculate the value of all other thermodynamic variables. The temperature dependent 283 function used in the iterative method is the internal energy. Initially, a temperature value is 284 assumed (for example the value of the temperature from the previous time RK sub-step or 285 from the previous time step) to initialize the iteration process. In most cells, this value is close 286 to the solution. Then the compressibility factor is calculated as the sum of the ideal gas 287 contribution (considered to be 1), the dispersion contribution and the residual hard-chain 288 contribution (Appendix A):

289

$$290 Z = 1 + Zhc + Zdisp (7)$$

291

292 The pressure is then calculated using eq.8 once the compressibility factor is known [38]:

(8)

(9)

293 
$$p = Zk_B T \rho_m (10^{10})^3$$

294 where k is the Boltzmann constant and  $\rho_m$  is the total number density of molecules.

Finally, the internal energy is estimated as the sum of the ideal internal energy and the residual internal energy. The ideal internal energy is computed using the ideal enthalpy. The residual internal energy is calculated using eq.9 [39]:

298

299 
$$\frac{e^{res}}{RT} = -T \left(\frac{\partial \tilde{a}^{res}}{\partial T}\right)_{\rho, x_i}$$

300 where  $\tilde{a}^{res}$  is the reduced Helmholtz free energy.

301

302 If the difference between the internal energy computed with the PC-SAFT model and the 303 value obtained from the conservative variables is bigger than 0.001J/kg, the Newton-Raphson 304 method is applied to calculate a new value of the temperature and the aforementioned steps 305 are repeated, see Appendix D.

306

#### 307 Quasi-conservative formulation

The thermodynamic variables computed in the CFD code by the PC-SAFT EoS are the temperature, internal energy, sound speed and enthalpy. The algorithm inputs are the density, pressure, molar fractions and three pure component parameters per component. The density and mass fractions (used to compute the molar fractions) are obtained from the conservative variables. The pressure is obtained employing the double flux model. The temperature is iterated until the difference between the pressure computed with the PC-SAFT model and the value obtained from the double flux model is lower than 0.001Pa, see Appendix D.

315

#### 316 **2.3 Peng-Robinson EoS and PC-SAFT EoS comparison**

The most attractive feature of the PC-SAFT EoS is the better prediction of derivative properties such as compressibility and speed of sound. [27] shows the inaccuracy of cubic models to predict second derivative properties such as isobaric heat capacity and sound velocity in hydrocarbons at high density ranges. In the case of the sonic fluid velocity, the AAD% (Average Absolute Deviation) by PR EoS for methane, ethane, and propane are 28.6%, 14.7%, and 61.2%, respectively.

323 Figure 2 presents a comparison of the thermodynamic properties of n-dodecane at 324 6MPa computed using the PC-SAFT EoS and the Peng-Robinson EoS. NIST Refprop [40] 325 has been used as reference due to its extensive validation with experimental data. While the 326 results of both EoS are quite similar at density values lower than 550 Kg/m<sup>3</sup> there is a 327 significant difference at higher densities, especially in the sound speed. Cubic models 328 commonly present low accuracy computing the thermodynamic properties of hydrocarbons at temperatures typical for injection systems [2]. However, the PC-SAFT EoS shows an 329 330 accuracy similar to NIST without the need of an extensive model calibration as only three 331 parameters are needed to model a specific component. Another advantage is the possibility of 332 computing the thermodynamic properties of mixtures; NIST has limited mixture 333 combinations.



Figure 2: Comparison of thermodynamic properties of n-dodecane at 6MPa computed using the PC-SAFT EoS and the Peng-Robinson EoS: (a) density, (b) sound speed, (c) internal energy

#### **336 3. Results**

Initially, advection test cases and shock tube problems are solved to validate the hyperbolic part of the numerical framework employing the conservative and quasiconservative formulations, while the parabolic part is omitted. Following, two-dimensional simulations of transcritical and supercritical nitrogen and dodecane jets are presented, including the parabolic part, to prove the multi-dimensional capability of the code.

342

### 343 **3.1. One-dimensional cases**

### 344 **3.1.1 Advection test cases**

### 345 *Conservative formulation*

346 Figure 3 shows the results of the supercritical Advection Test Case 1, see Table 2. 347 Nitrogen is used as working fluid (The critical properties of nitrogen are  $p_{c,N2} = 3.4$  MPa and 348  $T_{c,N2} = 126.2$  K). The computational domain is x  $\in [0, 1]$  m; the initial conditions in 0.25 m < x < 0.75 m are  $\rho = 250$  kg/m<sup>3</sup>, p=5 MPa, and T=139.4 K; in the rest of the domain are  $\rho = 45.5$ 349 350 kg/m<sup>3</sup>, p=5 MPa, and T=367.4 K. The advection velocity applied is 50m/s; periodic boundary 351 conditions are utilized; a uniform grid spacing of 0.01m is employed; the simulated time is 352 t=0.02s; and the CFL is set to be 0.5. Four spatial discretization schemes are compared: fifth-353 order WENO, second-order (based on the Minmod limiter), first order and a blend of the 354 fifth-order WENO and the first-order schemes (95% WENO and rest 1st order).

The oscillations are more severe when high-order reconstruction schemes are applied. By blending a high-order scheme and a low-order model, dissipation can be used to smooth the numerical solution. If the advection test case is initialized using a smooth profile no spurious pressure oscillation appear in the solution as the sharp jumps in the thermodynamic properties between cells are avoided, see Figure 4. The smooth initial interface was generated as described in [13] using eq.10.

$$361 q = q_L(1 - f_{sm}) + q_R f_{sm} (10)$$

$$362 \qquad f_{sm} = \frac{(1 + erf[\Delta R / \varepsilon])}{2} \tag{11}$$

363 Where L and R refers to the left and right states respectively and  $\Delta R$  is the distance from the 364 initial interface.  $\mathcal{E} = C_{\varepsilon} \Delta x$ , where  $\Delta x$  is the grid spacing and  $C_{\varepsilon}$  is a free parameter to 365 determine the interface smoothness set to be 8.

366





Figure 3. Advection Test Case 1 (N<sub>2</sub>), FC formulation, CFL = 0.5, u = 50 m/s, 100 cells, t=0.02 s. Comparison of the (a) density, (b) temperature, (c) pressure and (d) x-velocity between the analytical and the numerical solution.





#### 374 Quasi-conservative formulation

Figure 5 presents the results of the transcritical Advection Test Case 1 solved using the QC formulation. The advection velocity applied is 50m/s; periodic boundary conditions are applied; a uniform grid spacing of 0.01m is used; the simulated time is t=0.02s; and the CFL is set to be 1. Unlike the fully conservative scheme, spurious pressure oscillations are not present in the solution.

380



Figure 5. Advection Test Case 1 (N<sub>2</sub>), FC and QC formulations, CFL(FC) = 0.5, CFL(QC)=1.0, u = 50 m/s, 100 cells, t=0.02 s. Comparison of the (a) density, (b) temperature, (c) pressure and (d) x-velocity between the analytical and the numerical solution.

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386 Figure 6 presents the results of the transcritical Advection Test Case 2 where nitrogen is used as working fluid, see Table 2. The computational domain is  $x \in [0, 1]$  m; the initial 387 conditions in 0.25 m < x < 0.75 m are p=804.0 kg/m<sup>3</sup>, p=4 MPa, and T=84.41 K; in the rest of 388 389 the domain the initial conditions are  $p=45.5 \text{ kg/m}^3$ , p=4 MPa, and T=299.0 K. The advection 390 velocity utilized is 100 m/s; periodic boundary conditions are used; the computational domain 391 is x  $\in [0, 1]$  m; 150 cells are employed; the simulated time is t=0.01 s; a fifth-order WENO 392 discretization scheme is used; and the CFL is set to be 1.0. It can be observed how large 393 density gradients are solved without spurious pressure oscillations applying the double flux 394 model.

395 Figure 7 shows the results of the transcritical advection of n-dodecane at supercritical pressure and subcritical temperature (pc,n-dodecane =1.817 MPa, Tc,n-dodecane =658.1 K) in 396 397 supercritical nitrogen, Advection Test Case 3 (Table 2). The computational domain is x  $\epsilon$ 398 [0,1] m; the initial conditions in 0.25 m < x < 0.75 m are  $\rho_{n-dodecane} = 700.0 \text{ kg/m}^3$ ,  $p_{n-dodecane} =$ 399 6MPa, and  $T_{n-dodecane}$  = 360.1 K; in the rest of the domain  $\rho_{N2}$  = 20.0 kg/m<sup>3</sup>,  $p_{N2}$  = 6 MPa, and  $T_{N2}$ 400 =965.7 K. The advection velocity utilized is 100 m/s; periodic boundary conditions are used; 401 150 cells are employed; the simulated time is t=0.01 s; a fifth-order WENO discretization 402 scheme is used; and the CFL is set to be 1.0. Unlike conservative codes, velocity and pressure 403 equilibriums are preserved in multicomponent cases if the double flux model is applied.

Table 1. PC-SAFT pure component parameters [38]							
		m	$\sigma$ [Å]	$\mathcal{E}/k[K]$			
NITROGEN		1.2053	3.3130	90.96			
DODECANE		5.3060	3.8959	249.21			
		<b>Table 2. 1</b>	D Test Cases				
		ADVECTION	N TEST CASES				
CASE 1		Pressure [MPa]	Density [kg/m <sup>3</sup> ]	Temperature [K]			
0.25  m < x < 0.25  m	).75 m	N <sub>2</sub> , 5	N <sub>2</sub> , 250	N <sub>2</sub> , 139.4			
$0.25 \text{ m} > x \text{ or } x^{2}$	> 0.75 m	N <sub>2</sub> , 5	N <sub>2</sub> , 45.5	N <sub>2</sub> , 367.4			
CASE 2	2		C				
0.25  m < x < 0.25  m	).75 m	N2, 4	N <sub>2</sub> , 804	N <sub>2</sub> , 84.4			
0.25  m > x  or  x > 100  or  x > 1	> 0.75 m	N2, 4	N <sub>2</sub> , 45.5	N <sub>2</sub> , 299.0			
CASE 3	;						
0.25 m < x < 0.75 m		n-dodecane, 6.0	n-dodecane, 700.0	n-dodecane, 360.1			
0.25  m > x  or  x > 0.75  m		N <sub>2</sub> , 6.0	N <sub>2</sub> , 20.0	N <sub>2</sub> , 965.7			
		SHOCK TU	BE PROBLEM				
PROBLE	М	Pressure [MPa]	Density [kg/m <sup>3</sup> ]	Temperature [K]			
x < 0.5 m		n-dodecane, 13.0	n-dodecane, 700.0	n-dodecane, 372.8			
x > 0.5 n	n	n-dodecane, 6.0	n-dodecane, 150.0	n-dodecane, 944.4			
	ľ						
Table 3. 2D Test Cases							
CASE A	Press	sure [MPa]	Density [kg/m <sup>3</sup> ]	Temperature [K]			
JET	1	N <sub>2</sub> , 4.0	N <sub>2</sub> , 804.0	N <sub>2</sub> , 84.4			
CHAMBER		N <sub>2</sub> , 4.0	N <sub>2</sub> , 45.5	N <sub>2</sub> , 299.5			
CASE B							
JET	]	N <sub>2</sub> , 4.0	N <sub>2</sub> , 440.0	N <sub>2</sub> , 127.0			
CHAMBER	1	N <sub>2</sub> , 4.0	N <sub>2</sub> , 44.5	N <sub>2</sub> , 305.0			
CASE C							
JET	n-dod	lecane, 11.1	n-dodecane, 450.0	n-dodecane, 687.2			
CHAMBER	Ν	J <sub>2</sub> , 11.1	N <sub>2</sub> , 37.0	N <sub>2</sub> , 972.9			



#### 423 Energy conservation error in the quasi-conservative formulation

The evolution of the energy conservation error of the Advection Test Case 2 is presented in
Figure 8 . The error has been evaluated employing eq.12 [3].

427 
$$\varepsilon = \left| \frac{\int_{\Omega} \left[ (\rho E)(t) - (\rho E)(0) \right] dx}{\int_{\Omega} (\rho E)(0) dx} \right|$$
(12)

428 where  $\varepsilon$  is the relative error of the total energy respect to initial conditions and  $\Omega$  is the 429 computational domain.

430

431 The energy conservation error is higher using the PC-SAFT EoS than Peng-Robinson EoS. 432 This is related to the fact that the profiles of  $\gamma^*$  and  $e_0^*$  are smoother employing the cubic 433 model. There are shaper jumps in the internal energy and speed of sound employing the PC-434 SAFT EoS, see Figure 10. The error in the conservation of the energy depends on the jumps 435 in the variables  $1/(\gamma^* - 1)$  and  $e^*$  [3]. A convergence of the error to 0 exists increasing the 436 refinement.

> 10 Energy conservation error 10 = 400 PR 10 N = 400 PC-SAFT 0 = 200 PR = 200 PC-SAFT 100 PR 10 0 = 100 PC-SAFT N = 50 PR N = 50 PC-SAFT 10 0.008 0.01 Õ 0.002 0.004 0.006 t[s]

437

Figure 8. Relative energy conservation error computed using eq.10 of the QC formulation for the
 Advection Test Case 2 (Transcritical nitrogen) using the Peng-Robinson EoS (PR) and the PC SAFT EoS. N is the number of cells employed.

441



442

Figure 9. Relative energy conservation error computed using eq.10 of QC formulation for the
 Advection Test Case 3 using the PC-SAFT EoS. N is the number of cells employed.



445 446

Figure 10. Advection Test Case 2 (N<sub>2</sub>), QC formulation, CFL = 1.0, u = 150 m/s, 100 cells, t=0.01s. Comparison of  $\gamma^*$  and  $e_0^*$  computed using the Peng Robinson EoS (PR EoS) and the PC-SAFT in the Advection Test Case 2.

447 448

Figure 9 presents the evolution of the energy conservation error of the Advection Test Case 3. Because of the different thermodynamic properties of the components, a higher energy conservation error than in the single-species cases appears. Although, a convergence to 0 is observed in one-dimensional cases increasing the refinement like in the single-species cases.

454

### 455 **3.1.2 Shock tube problems**

The Euler equations are solved in this validation so a direct comparison with the exact solver can be done. The exact solution has been computed using the methodology described in [41].

458

### 459 Quasi-conservative formulation

460 The domain is  $x \in [0, 1]$  m. The working fluid employed is dodecane. A fifth-order 461 WENO scheme is employed to interpolate the primitive variables onto the cell faces. 800 462 equally spaced cells were used. Wave transmissive boundary conditions are implemented in 463 the left and right sides. The double flux model is applied. The pressure exceeds the critical 464 value in all the domain while there is a transition in the temperature from subcritical to 465 supercritical from left to right. The initial conditions in the left state are  $\rho_L = 700 \text{ kg/m}^3$ ,  $p_L = 13$ MPa,  $u_L=0$  m/s; and in the right state are  $\rho_R=150$  kg/m<sup>3</sup>,  $p_R=6$  MPa,  $u_R=0$  m/s. The simulated 466 467 time is t=0.2 ms.

Figure 11 displays the results obtained for density, temperature, pressure and velocity. Despite being a quasi-conservative scheme, the double flux model [3], [17], [33] can solve strong shock waves in transcritical cases with a high degree of accuracy without generating spurious pressure oscillations.



Figure 11. Shock Tube Problem 1 (Dodecane), QC formulation, CFL = 1.0, 800 cells, t=0.2 ms. Comparisons of (a) density, (b) temperature, (c) velocity and (d) pressure profiles: exact solution and numerical solution.





#### 481 Conservative formulation

482 The same shock tube problem described before is solved. A fifth-order WENO 483 scheme is employed to interpolate the conservative variables onto the cell faces. Large 484 spurious pressure oscillations appear in the solution because of the sharp jumps in the 485 thermodynamic properties between cells, see Figure 12.

486

#### 487 Comparison with the Peng-Robinson EoS (Quasi-conservative formulation)

488 Figure 13 shows the density, temperature, pressure, velocity, sound speed and 489 internal energy of the same shock tube problem solved in a larger domain x  $\in [0, 2]$  m using 490 the PC-SAFT and the Peng-Robinson EoS. The simulated time is t=0.3 ms. The quasi-491 conservative formulation has been employed. 800 equally spaced cells were used. A 492 significant difference can be observed in the results between the two EoS. Due to the high 493 deviation in the sound speed computed by the Peng-Robinson EoS in the high-density region, 494 the expansion wave travels much faster using the cubic model. Moreover, the calculated 495 temperatures are much lower using the Peng-Robinson EoS in the high-density region. 496





#### 501 3.2 Two-dimensional cases

Planar two-dimensional simulations of transcritical and supercritical jets are presented in this section. The initial conditions are summarized in Table 3. The parabolic sub-step is included into these simulations, without sub-grid scale modelling for turbulence or heat/species diffusion.

506

#### 507 Transcritical nitrogen injection (Quasi-Conservative formulation, Case A)

508 A structured mesh is applied with a uniform cell distribution. The cell size is 0.043 mm 509  $\times$  0.043 mm. The domain used is 30mm  $\times$  15mm. Transmissive boundary conditions are 510 applied at the top, bottom and right boundaries while a wall condition is employed at the left 511 boundary. A flat velocity profile is imposed at the inlet. The case is initialized using a 512 pressure in the chamber of 4 MPa, the density of the nitrogen in the chamber is  $45.5 \text{ kg/m}^3$ 513 and the temperature is 299.5 K. The temperature of the jet is 84.4 K and the density is 804.0 514 kg/m. A summary of the initial conditions can be found in Table 3. The velocity of the jet is 515 100 m/s and the diameter of the exit nozzle is 1.0 mm.

516



517

518 519

Figure 14. 2D Test Case A, CFL = 1.0, 245000 cells, QC formulation. Density results of the simulation of the planar cryogenic nitrogen jet at various times.

520 When the jet enters the elevated temperature environment of the chamber, the 521 velocity gradients at the jet surface generate a vortex rollup that finally breakup into ligament-522 shaped structures, see Figure 14. The Kelvin Helmholtz instability can be observed in the 523 shear layer, which is similar to a gas/gas turbulent mixing case. No droplets are formed at 524 these conditions. The jet is quickly heated to a gas-like supercritical state after the injection 525 takes place. It must be highlighted that the mesh resolution is not enough to resolve all the 526 scales (the aim of these simulations is to test the developed numerical framework). Moreover, 527 2D simulation cannot resolve turbulence. Figure 17 shows the density, temperature, pressure 528 and sound speed results at  $4 \times 10^{-4}$  s.

Figure 15 shows a scatter plot of pressure as a function of density for the planar cryogenic nitrogen jet. The simulated case remains in the hyperbolic region of the governing equations with a real-valued speed of sound (Appendix B). The mixing trajectory passes close to the critical point with a few individual points inside the saturation curve, which means that phase separation does not occur [42]. The larger fluctuations caused by the confined domain or the two-dimensionality of the case could be the reason why a small number of cells are in the unstable region [3].

Although one of reasons of the prevailing usage of cubic EoS is their efficiency, practical simulations can be performed using the PC-SAFT EoS. The quasi-conservative formulation is computationally less expensive than the conservative scheme because the PC-SAFT EoS has to be used only once in the hyperbolic operator in each time step. The computational time is 65-70% higher using the PC-SAFT EoS than utilizing the PR EoS. Figure 16 shows the time taken by the code to solve the transcritical nitrogen injection case depending on the number of cells used (only one core is used to perform the simulation).

The PC-SAFT EoS is implemented using loops that depend on the number of components solved, which means that it takes more time to compute the properties of mixtures. However, knowing the mass fractions it is possible to determine how many components are present in a cell a priori. The PC-SAFT is then only solved for that specific number of components. Most cells along the simulation in the combustion chamber contain only nitrogen. For this reason, a significant increment on time has not been observed performing two-component simulations.



Figure 15. Scatter plot of pressure as a function of density for the transcritical nitrogen jet (Case
 A). The vapor dome, non-convex region and the region with complex speed of sound (SOS) are
 included.





- (Case A) at  $t = 4 \times 10^{-4}$  employing a variable number of cells.
- 556 557



Figure 17. 2D Test Case A, CFL = 1.0, 245000 cells, QC formulation. Results of the simulation of the planar cryogenic nitrogen jet at t = 4 x 10<sup>-4</sup> s using the quasi-conservative formulation: (a) density, (b) temperature, (c) pressure, (d) sound speed.

#### 562 Supercritical nitrogen injection (Conservative formulation, Case B)

The case is initialized using a pressure in the chamber of 4 MPa, the density of the nitrogen in the chamber is 45.5 kg/m3 and the density of the jet is 440.0 kg/m<sup>3</sup> (Table 3). The velocity of the jet is 50 m/s. The spatial reconstruction is carried out using a blending of the fifth-order WENO and the first-order schemes (95% fifth-order WENO). The CFL number is set at 0.4. Transmissive boundary conditions are applied at the top, bottom and right boundaries while a wall condition is employed at the left boundary. A flat velocity profile is imposed at the inlet.

570 If sharp interface methods (i.e. front tracking method) are not applied, the interfaces 571 are not sharp one-point jumps but smooth as they are resolved [43]. This is the reason why the 572 wiggles that appear in this 2D simulation are not as severe as in the 1D cases presented in 573 Section 3.1 initialized using a sharp interface, see Figure 18. The study of [43] shows how 574 smooth interfaces can reduce the spurious pressure oscillations.

The minimum pressure encountered along the simulation is higher than the nitrogen critical pressure so there are no cells in the vapor-liquid equilibrium region. The heat-up of the jet follows the same density-temperature trajectory employing a FC or a QC formulation in single-species cases, see Figure 19. In the works of [2], [44] a different behaviour in multicomponent cases can be observed, where QC formulations follow an isobaric-isochoric mixing model for binary mixtures while conservative schemes follow an isobaric-adiabatic mixing model.



583Figure 18. 2D Test Case B, CFL = 0.4, 180000 cells, FC formulation. Results of the simulation of<br/>the supercritical nitrogen jet at t = 7.84 x 10<sup>-4</sup> s: (a) density, (b) pressure.







Figure 19. 2D Test Case B solved using the FC and QC formulations. Scattered data of density and temperature. The nitrogen vapor dome is included.

### 588

#### 589 Supercritical dodecane injection (Conservative formulation, Case C)

590 Dodecane/nitrogen mixtures are Type IV as stated by [45], which means that the 591 critical temperature of the mixture is an intermediate value of the critical temperature of both 592 components and the mixture critical pressure is higher than the critical pressure of either 593 component, see Figure 23. A simulation of a dodecane jet where VLE (Vapor-Liquid 594 Equilibrium) conditions are avoided injecting the dodecane at a temperature higher than its 595 critical value has been included to prove the multi-species capability of the code. To check 596 that any cell is in a thermodynamic metastable state, the vapor-liquid saturation curves were 597 computed (Appendix C).

598 A structured mesh is applied with a uniform cell distribution. The cell size is  $8.3 \mu m \times$ 599  $8.3\mu m$ . The domain used is  $5mm \times 2.5mm$ . Transmissive boundary conditions are applied at 600 the top, bottom and right boundaries while a wall condition is employed at the left boundary. 601 A flat velocity profile is imposed at the inlet. The case is initialized using a pressure in the 602 chamber of 11.1 MPa, the density and the temperature of the nitrogen in the chamber are 37.0 603 kg/m<sup>3</sup> and 973 K (high-load Diesel operation conditions [46]) respectively. The density and 604 temperature of the jet are 450.0 kg/m<sup>3</sup> and 687 K (Table 3). The velocity of the jet is 200 m/s and the diameter of the exit nozzle is 0.1 mm. 605

As in the transcritical nitrogen case ligament-shaped structures appear and the Kelvin Helmholtz instability can be observed in the shear layer, see Figure 20. The jet is quickly heated-up from a liquid-like supercritical state to a gas-like supercritical state. Some spurious oscillations appear in the pressure field because of the high non-linearity of the EoS. The quasi-conservative formulation was not employed because of the incorrect prediction of the jet heat-up that appear in multi-component cases [2], [44].

A comparison of averaged scattered data of composition and temperature and an isobaric-adiabatic mixing process can be seen in Figure 21. As [44] stated, fully conservative schemes describe an isobaric-adiabatic mixing process. The isobaric-adiabatic line in the Figure 21 was computed using eq.13-14 and the initial conditions of this case.

$$\dot{m}_3 = \dot{m}_1 + \dot{m}_2 \tag{13}$$

$$\dot{m}_3 h_3 = \dot{m}_1 h_1 + \dot{m}_2 h_2 \tag{14}$$



Figure 20. 2D Test Case C, CFL = 0.5, 180000 cells, FC formulation. Results of the simulation of the supercritical dodecane jet at t = 2.5 x 10<sup>-5</sup> s: (a) density, (b) temperature, (c) pressure, (d) sound speed.





#### 625 **4.** Conclusions

626 The Perturbated Chain Statistical Associating Fluid Theory (PC-SAFT) is utilized to close the 627 Navier-Stokes equations using both a conservative and a quasi-conservative formulation, 628 where the double flux model of [3], [17], [33] is applied. The PC-SAFT EoS presents a 629 precision similar to NIST without the need of an extensive calibration as only three 630 parameters are needed to model a specific component. It is presented as an alternative to the 631 commonly used cubic EoS that present a low accuracy for computing the thermodynamic 632 properties of hydrocarbons at temperatures typical for high pressure injection systems. 633 Advection test cases and shock tube problems have been used to validate the hyperbolic 634 operator of the developed numerical framework. The conservative formulation generates 635 spurious pressure oscillations, like it has been reported with other diffuse interface density-636 based codes employing a real-fluid EoS. Due to fact that the interfaces are not sharp one-point 637 jumps but smooth, as they are resolved in 2D simulations, the wiggles generated do not 638 compromise the stability of the simulation. The quasi-conservative scheme can model 639 transcritical single- and multicomponent cases without spurious pressure oscillations. Errors 640 in the energy conservation that appear employing this formulation may produce an unphysical 641 quick heat-up of the injected jet in multicomponent cases. Two-dimensional simulations of 642 nitrogen and dodecane jets have been presented to demonstrate the multidimensional and 643 multicomponent capability of the numerical framework.

#### 644 Acknowledgments

645 This project has received funding from the European Union Horizon-2020 Research646 and Innovation Programme with grant Agreement No 675528.

- 647
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- 649
- 650
- 651

#### 652 Appendix A PC-SAFT EoS

The PC-SAFT EoS is expressed as the sum of all the residual Helmholtz free energy
contributions. These contributions correspond to the distinct types of molecular interactions.
The residual Helmholtz free energy is computed using eq.15 [38].

$$657 \qquad \tilde{a}^{res} = \tilde{a}^{hc} + \tilde{a}^{disp} \tag{15}$$

658

656

659 The hard-chain term,  $\tilde{a}^{hc}$ , for a mixture of nc components, is given in eq. 16

660 
$$\tilde{a}^{hc} = \bar{m}\tilde{a}^{hs} - \sum_{i}^{nc} x_i (m_i - 1) \ln g_{ii}^{hs} (\sigma_{dii})$$
 (16)

where  $\overline{m}$  is the number of segments for a multicomponent mixture (eq. 17),  $x_i$  is the mole fraction of every component i in the fluid,  $\tilde{a}^{hs}$  is the hard sphere contribution (eq. 18),  $g_{ii}^{hs}$  is the radial distribution function of the hard-sphere fluid (eq.23) and  $m_i$  is the number of segments per chain of every component.

665

666 The number of segments for a multicomponent mixture is:

$$667 \qquad \overline{m} = \sum_{i}^{nc} x_i m_i \tag{17}$$

668 The hard sphere contribution is:

669 
$$\tilde{a}^{hs} = \frac{1}{\varsigma_0} \left[ \frac{3\varsigma_1\varsigma_2}{(1-\varsigma_3)} + \frac{\varsigma_2^3}{\varsigma_3(1-\varsigma_3)^2} + \left(\frac{\varsigma_2^3}{\varsigma_3^2} - \varsigma_0\right) \ln(1-\varsigma_3) \right]$$
(18)

670  $\boldsymbol{\zeta}_n$  is defined as:

671 
$$\zeta_n = \frac{\pi}{6} \rho_m \sum_i x_i m_i d_i^n$$
  $n \in \{0, 1, 2, 3\}$  (19)

672 where  $\rho_m$  is the molecular density and  $d_i$  is the temperature-dependent segment diameter of 673 component i (eq.21).

674 
$$\rho_m = \frac{6}{\pi} \eta \left( \sum_i x_i m_i d_i^3 \right)^{-1} \text{ being } \eta = \varsigma_3$$
(20)

675 
$$d_{i} = \sigma_{di} \left[ 1 - 0.12 \exp\left(-3\frac{\varepsilon_{i}}{kT}\right) \right]$$
(21)

where k is the Boltzmann constant, T is the temperature and  $\varepsilon_i$  is the depth of pair potential of the component.

The mixture parameters  $\sigma_{ij}$  and  $\varepsilon_{ij}$  which are defined for every pair of unlike segments are modeled using a Berthelot-Lorentz combining rule.

681 
$$\sigma_{ij} = \frac{1}{2}(\sigma_i + \sigma_j)$$
(22)

$$\epsilon_{ij} = \sqrt{\varepsilon_i \varepsilon_j} (1 - k_{ij})$$
(23)

683 where  $k_{ij}$  is the binary interaction parameter.

684

685 The radial distribution function of the hard-sphere fluid is:

686 
$$g_{ij}^{hs} = \frac{1}{(1-\zeta_3)} + \left(\frac{d_i d_j}{d_i + d_j}\right) \frac{3\zeta_2}{(1-\zeta_3)^2} + \left(\frac{d_i d_j}{d_i + d_j}\right)^2 \frac{3\zeta_2^2}{(1-\zeta_3)^3}$$
 (24)

687 The dispersion term is defined as:

688 
$$\tilde{a}^{disp} = -2\pi p_m I_1(\eta, \bar{m}) \overline{m^2 \varepsilon \sigma_d^3} - \pi p_m \bar{m} C_1 I_2(\eta, \bar{m}) \overline{m^2 \varepsilon^2 \sigma_d^3}$$
(25)

689 where  $\eta = \zeta_3$  is the reduced density,  $I_1$  and  $I_2$  are integrals approximated by simple power 690 series in density

691 
$$I_1(\eta, \overline{m}) = \sum_{i=0}^6 a_i(\overline{m})\eta^i$$
(26)

692 
$$I_2(\eta, \overline{m}) = \sum_{i=0}^6 b_i(\overline{m})\eta^i$$
(27)

694 
$$a_i(\overline{m}) = a_{0i} + \frac{m-1}{\overline{m}}a_{1i} + \frac{m-1}{\overline{m}}\frac{m-2}{\overline{m}}a_{2i}$$
 (28)

695 
$$b_i(\overline{m}) = b_{0i} + \frac{\overline{m-1}}{\overline{m}} b_{1i} + \frac{\overline{m-1}}{\overline{m}} \frac{\overline{m-2}}{\overline{m}} b_{2i}$$
 (29)

696 Where  $a_{0i}, a_{1i}, a_{2i}, b_{0i}, b_{1i}, b_{2i}$  are constants [38].

697  $C_1$  is defined as:

698

$$C_{1} = \left(1 + Z^{hc} + \rho \frac{\partial Z^{hc}}{\partial \rho}\right)^{-1} = \left(1 + \overline{m} \frac{8\eta - 8\eta^{2}}{(1 - \eta)^{4}} + (1 - \overline{m}) \frac{20\eta - 27\eta^{2} + 12\eta^{3} - 2\eta^{4}}{[(1 - \eta)(2 - \eta)]^{2}}\right)^{-1}$$
(30)

699 The terms  $\overline{m^2 \mathcal{E} \sigma_d^3}$  and  $\overline{m^2 \mathcal{E}^2 \sigma_d^3}$  are defined as:

700 
$$\overline{m^2 \varepsilon \sigma_d^3} = \sum_{i}^{nc} \sum_{j}^{nc} x_i x_j m_i m_j \left(\frac{\varepsilon_{ij}}{kT}\right) \sigma_{ij}^3$$
(31)

701 
$$\overline{m^2 \varepsilon^2 \sigma_d^3} = \sum_i^{nc} \sum_j^{nc} x_i x_j m_i m_j \left(\frac{\varepsilon_{ij}}{kT}\right)^2 \sigma_{ij}^3$$
(32)

#### 702 Compressibility factor

Then the compressibility factor is calculated as the sum of the ideal gas contribution (considered to be 1), the dispersion contribution and the residual hard-chain contribution [38]:

705 
$$Z = 1 + Z^{hc} + Z^{disp}$$
 (33)

706 
$$Z^{hs} = \frac{\zeta_3}{(1-\zeta_3)} + \frac{3\zeta_1\zeta_2}{\zeta_0(1-\zeta_3)^2} + \frac{3\zeta_2^3 - \zeta_3\zeta_2^3}{\zeta_0(1-\zeta_3)^3}$$
(34)

707 
$$Z^{hc} = \overline{m}Z^{hs} - \sum_{i} x_{i}(m_{i}-1)(g_{ii}^{hs})^{-1}\rho_{m}\frac{\partial g_{ii}^{hs}}{\partial \rho_{m}}$$
 (35)

708 
$$Z^{disp} = -2\pi\rho_m \frac{\partial(\eta I_1)}{\partial \eta} \overline{m^2 \varepsilon \sigma_d^3} - \pi\rho_m \overline{m} \left[ C_1 \frac{\partial(\eta I_2)}{\partial \eta} + C_2 \eta I_2 \right] \overline{m^2 \varepsilon^2 \sigma_d^3}$$
(36)

709 where:

710 
$$C_{2} = \frac{\partial C_{1}}{\partial \eta} = -C_{1}^{2} \left( \frac{\overline{m} - 4\eta^{2} + 20\eta + 8}{(1 - \eta)^{5}} + (1 - \overline{m}) \frac{2\eta^{3} + 12\eta^{2} - 48\eta + 40}{[(1 - \eta)(2 - \eta)]^{3}} \right)$$
(37)

$$\rho \frac{\partial g_{ij}^{hs}}{\partial \rho} = \frac{\zeta_3}{(1-\zeta_3)^2} + \left(\frac{d_i d_j}{d_i + d_j}\right) \left(\frac{3\zeta_2}{(1-\zeta_3)^2} + \frac{6\zeta_2 \zeta_3}{(1-\zeta_3)^3}\right) + \frac{1}{(1-\zeta_3)^3} + \frac{1}{(1-\zeta_3)^3} + \frac{1}{(1-\zeta_3)^4} + \frac{1}{(1-\zeta_3)^4$$

712 
$$\frac{\partial(\eta I_1)}{\partial \eta} = \sum_{j=0}^{6} a_j(\overline{m})(j+1)\eta^i$$
(39)

713 
$$\frac{\partial(\eta I_2)}{\partial \eta} = \sum_{j=0}^{6} b_j(\overline{m})(j+1)\eta^i$$
(40)

714

### 715 *Derivative of the Helmholtz free energy respect to temperature.*

716 The temperature derivative of  $\tilde{a}^{res}$  is the sum of two contributions.

718 
$$\left(\frac{\partial \tilde{a}^{res}}{\partial T}\right)_{\rho,x_i} = \left(\frac{\partial \tilde{a}^{hc}}{\partial T}\right)_{\rho,x_i} + \left(\frac{\partial \tilde{a}^{disp}}{\partial T}\right)_{\rho,x_i}$$
 (41)

719 The temperature derivative of the Helmholtz free energy hard-chain reference contribution is:
720

721 
$$\left(\frac{\partial \tilde{a}^{hc}}{\partial T}\right)_{\rho,x_i} = \overline{m} \left(\frac{\partial \tilde{a}^{hs}}{\partial T}\right)_{\rho,x_i} - \sum_i x_i (m_i - 1) (g_{ii}^{hs})^{-1} \left(\frac{\partial g_{ii}^{hs}}{\partial T}\right)_{\rho,x_i}$$
(42)

722

The temperature derivative of the Helmholtz free energy residual contribution of the hard-sphere system is:

725

$$726 \qquad \left(\frac{\partial \tilde{a}^{hs}}{\partial T}\right)_{\rho,x_{i}} = \frac{1}{\varsigma_{0}} \left[ \frac{\frac{3\left(\varsigma_{1,T}\varsigma_{2}+\varsigma_{1}\varsigma_{2,T}\right)}{\left(1-\varsigma_{3}\right)^{2}} + \frac{3\varsigma_{1}\varsigma_{2}\varsigma_{3,T}}{\left(1-\varsigma_{3}\right)^{2}} + \frac{3\varsigma_{2}^{2}\varsigma_{2,T}}{\varsigma_{3}\left(1-\varsigma_{3}\right)^{2}} + \frac{\varsigma_{2}^{3}\varsigma_{3,T}\left(3\varsigma_{3}-1\right)}{\varsigma_{3}^{2}\left(1-\varsigma_{3}\right)^{3}} + \left[ \frac{3\varsigma_{2}^{2}\varsigma_{2,T}}{\varsigma_{3}^{2}\left(1-\varsigma_{3}\right)^{3}} + \frac{3\varsigma_{2}^{2}\varsigma_{2,T}}{\varsigma_{3}^{2}\left(1-\varsigma_{3}\right)^{2}} + \frac{\varsigma_{2}^{3}}{\varsigma_{3}^{2}\left(1-\varsigma_{3}\right)^{3}} + \frac{1}{\varsigma_{2}^{2}\left(1-\varsigma_{3}\right)^{3}} + \frac{1}{\varsigma_{2}^{2}\left(1-\varsigma_{3}\right)^{3}} + \frac{1}{\varsigma_{3}^{2}\left(1-\varsigma_{3}\right)^{3}} + \frac{1}{\varsigma_{3}^{2}\left(1-\varsigma_{3}\right)^{$$

728 with abbreviations for two temperature derivatives:

729 
$$\zeta_{n,T} = \frac{\partial \zeta_n}{\partial T} = \frac{\pi}{6} \rho \sum_i x_i m_i n d_{i,T} (d_i)^{n-1} \qquad n \in \{0, 1, 2, 3\}$$
(44)  
730 
$$d_i = \frac{\partial d_i}{\partial t} = \sigma \left(3 \frac{\varepsilon_i}{2}\right) \left[-0.12 \exp\left(-3 \frac{\varepsilon_i}{2}\right)\right]$$

730 
$$d_{i,T} = \frac{\partial d_i}{\partial T} = \sigma_i \left( 3 \frac{\varepsilon_i}{kT^2} \right) \left[ -0.12 \exp\left( -3 \frac{\varepsilon_i}{kT} \right) \right]$$

732 The temperature derivative of the radial pair distribution function is:

$$\frac{\partial g_{ii}^{hs}}{\partial T} = \frac{\zeta_{3,T}}{(1-\zeta_3)^2} + \left(\frac{1}{2}d_{i,T}\right) \frac{\zeta_2}{(1-\zeta_3)^2} + \left(\frac{1}{2}d_i\right) \left(\frac{3\zeta_{2,T}}{(1-\zeta_3)^2} + \frac{6\zeta_2\zeta_{3,T}}{(1-\zeta_3)^3}\right) + \left(\frac{1}{2}d_id_{i,T}\right) \frac{2\zeta_2^2}{(1-\zeta_3)^3} + \left(\frac{1}{2}d_i\right)^2 \left(\frac{4\zeta_2\zeta_{2,T}}{(1-\zeta_3)^3} + \frac{6\zeta_2^2\zeta_{3,T}}{(1-\zeta_3)^4}\right)$$
(45)

734

The temperature derivative of the Helmholtz free energy contribution due to dispersiveattraction is:

737

738
$$\begin{pmatrix} \frac{\partial \tilde{a}^{disp}}{\partial T} \\ \frac{\partial C_1}{\partial T} \\ \frac{\partial I_1}{\partial T} - \frac{I_1}{\partial T} \\ \frac{\partial C_2}{\partial T} \\ \frac{\partial I_2}{\partial T} + C_1 \frac{\partial I_2}{\partial T} - 2C_1 \frac{I_2}{T} \\ \frac{\partial T}{\partial T} \\ \frac{\partial C_2}{\partial T} \\ \frac{\partial C_1}{\partial T} \\ \frac{\partial C_2}{\partial T} \\ \frac{\partial I_2}{\partial T}$$

739

740 with

742 
$$\frac{\partial I_1}{\partial T} = \sum_{i=0}^6 a_i(\overline{m}) i \zeta_{3,T} \eta^{i-1}$$
(47)

743

744 
$$\frac{\partial I_2}{\partial T} = \sum_{i=0}^{6} b_i(\overline{m}) i \varsigma_{3,T} \eta^{i-1}$$

745

746 
$$\frac{\partial C_1}{\partial T} = \varsigma_{3,T} C_2$$

747

### 748 Estimation of enthalpy and sound speed.

The enthalpy is used to compute the thermal diffusion vector in the parabolic sub-step. It is computed as the sum of the ideal contribution (obtained by integrating the ideal heat capacity at constant pressure with respect to the temperature) and the residual enthalpy [38]:

752 
$$\frac{\hat{h}^{res}}{RT} = -T \left( \frac{\partial \tilde{a}^{res}}{\partial T} \right)_{\rho, x_i} + (Z - 1)$$
(50)

753

Sound speed is computed using the equation applied by [47]:

755 
$$c = \sqrt{\frac{C_p}{C_v} \left(\frac{\partial P}{\partial \rho_m}\right)_T}$$
(51)

where  $C_p$  and  $C_v$  are the heat capacities at constant pressure and volume respectively [39].

758 The derivatives needed to compute the sound speed are:

759 
$$\left(\frac{\partial P}{\partial \rho_m}\right)_{T,x_i} = \left(\frac{\partial P}{\partial \eta}\right)_{T,x_i} \left(\frac{\partial \eta}{\partial \rho_m}\right)_{T,x_i}$$
 (52)

760 
$$\left(\frac{\partial \eta}{\partial \rho_m}\right)_{T,x_i} = \frac{\pi}{6} \left(\sum_i x_i m_i d_i^3\right)$$
 (53)

761 
$$\left(\frac{\partial P}{\partial \eta}\right)_{T,x_i} = k_B T \left(10^{10}\right)^3 \left[\rho_m \left(\frac{\partial Z}{\partial \eta}\right)_{T,x_i} + Z \left(\frac{\partial \rho_m}{\partial \eta}\right)_{T,x_i}\right]$$
(54)

762

763 
$$\left(\frac{\partial \rho_m}{\partial \eta}\right)_{T,x_i} = \frac{6}{\pi} \left(\sum_i x_i m_i d_i^3\right)^{-1}$$
(55)

764 
$$\left(\frac{\partial Z}{\partial \eta}\right)_{T,x_i}$$
 can be found in [48]. (56)

(48)

(49)

### 765 Appendix B Hyperbolicity of Euler system with PC-SAFT EoS

The hyperbolicity of the Euler system relies on a real speed of sound [3]. Using the PC-SAFT, the speed of sound is always real outside of the vapor-liquid equilibrium state. Inside the vapor-liquid equilibrium region, the spinodal curves (determined by  $(\partial p / \partial v)_T = 0$ ) enclose the unstable / non-convex region where a complex speed of sound could be found, see Figure 22.

### 771 Appendix C Pressure-composition phase diagram for the N2+C12H26

#### 772 system

The calculation of the number of phases present in a mixture in a certain condition is a recognized problem in the utilization of any EoS. In some cases, the number of phases is assumed a priori and then the composition in every phase is calculated by imposing equilibrium conditions. However, this technique often leads to divergence in the iterative methods used to achieve these. In our case, this is solved by an isothermal flash calculation after a stability analysis using the Tangent Plane Criterion Method proposed by [49] and applied to the PC-SAFT EoS by [50], see Figure 23.





Figure 22. The vapor dome, non-convex region and the region with complex speed of sound of dodecane computed using the PC-SAFT EoS.



#### Appendix D PC-SAFT EoS subroutines 816

#### Algorithm 1: Fully conservative formulation 817 \_\_\_\_

igoritiim	I: runy	conservative formulation	

		From conservative variables: $\rho, e, Y_i$
		Inputs Temperature from the previous time step
		Specific values of each component: $\sigma_{,\varepsilon} / K_{,k_{ii}}, m$
		1) Compute the mole fraction of each component
		DO
		if [(abs (e(CSV)-e(PC-SAFT)) > 0.001] <i>then</i>
		2) Compute segment diameter of each component (eq.21)
		3) Compute mean segment number (eq.17)
		4) Compute the coefficients a, and b, (eq.28-29)
		5) Compute abbreviations (eq.31-32)
		5) Compute $\zeta_n$ (eq.19)
		6) Compute radial distribution function of the hard sphere fluid (eq.24)
		7) Compute contribution of the hard sphere fluid to the compressibility factor (eq.34)
		8) Compute contribution of the hard chain to the compressibility factor (eq.35)
		9) Compute dispersion contribution to the compressibility factor (eq.36)
		10) Compute total compressibility (eq.33)
	Algorithm -	11) Compute pressure (eq.8)
		12) Compute partial derivative of the Helmholtz free energy respect to temperature
		13) Compute residual internal energy
		14) Compute residual enthalpy
		15) Compute sonic fluid velocity
		16) Compute ideal enthalpy
		17) Compute ideal internal energy
		18) Compute total enthalpy
		19) Compute total internal energy
		20) Compute the new temperature using the Newton-Raphson method. The temperature
		dependent function used is the internal energy
819		ELSE
017		RETURN
		END IF
		END DO
820		
821		
822		
823		
824		
825		
826		

### 827 Algorithm 2: Quasi-conservative formulation

828			
		[	From conservative variables: $\rho$ , $Y_i$
		Turneta	From double flux model: p
		Inputs	Temperature from the previous time step
			Specific values of each component: $\sigma, \varepsilon / K, k_{ij}, m$
		1) Com	npute the mole fraction of each component
		DO	
		if [(ab	s (p(double flux model)-p(PC-SAFT)) > 0.001 <i>then</i>
		2) 0	Compute segment diameter of each component (eq.21)
		3) 0	Compute mean segment number (eq.17)
		4) 0	Compute the coefficients a <sub>i</sub> and b <sub>i</sub> (eq.28-29)
		5) 0	Compute abbreviations (eq.31-32)
		5) 0	Compute $\varsigma_n$ (eq.19)
		6) 0	Compute radial distribution function of the hard sphere fluid (eq.24)
		7) 0	Compute contribution of the hard sphere fluid to the compressibility factor (eq.34)
		8) (	Compute contribution of the hard chain to the compressibility factor (eq.35)
		9) (	Compute dispersion contribution to the compressibility factor (eq.36)
		10)	Compute total compressibility (eq.33)
	Algorithm	{ 11)	Compute pressure (eq.8)
		12)	Compute partial derivative of the Helmholtz free energy respect to temperature
		13)	Compute residual internal energy
		14)	Compute residual enthalpy
		15)	Compute sonic fluid velocity
		16)	Compute ideal enthalpy
		17)	Compute ideal internal energy
		18)	Compute total enthalpy
		19)	Compute total internal energy
		20)	Compute the new temperature using the Newton-Raphson method. The temperature
		deper	ident function used is the pressure
829		ELSE	
		RET	URN
		END I	F
		END I	00
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03U 021			
001			
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833			
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