Developing intermolecular-potential models for use with the SAFT-VR Mie Equation of State

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Abstract A major advance in the statistical associating fluid theory for potentials of variable range (SAFT-VR) has recently been made with the incorporation of the Mie (generalized Lennard-Jones) interaction between the segments comprising the molecules in the fluid. [Lafitte et al. J. Chem. Phys. 2013:139:154504] The Mie potential offers greater versatility in allowing one to describe the softness/hardness of the repulsive interactions and the range of the attractions, which govern fine details of the fluid-phase equilibria and thermodynamic derivative properties of the system. In our current work, the SAFT-VR Mie equation of state is employed to develop models for a number of prototypical fluids, including some of direct relevance to the oil and gas industry: methane, carbon dioxide and other light gases, alkanes, alkyl benzenes, and perfluorinated compounds. A complication with the use of more-generic force fields like the Mie potential is the additional number of parameters that have to be considered to specify the interactions between the molecules, leading to a degree of degeneracy in the parameter space. A formal methodology to isolate intermolecularpotential models and assess the adequacy of the description of the thermodynamic properties in terms of the complex parameter space is developed. Fluid-phase equilibrium properties (the vapour pressure and saturated-liquid density) are chosen as the target properties in the refinement of the force fields; the predictive capability for other properties such as the enthalpy of vaporization, single-phase density, speed of sound, isobaric heat capacity, and Joule-Thomson coefficient, is appraised. It is found that an overall improvement of the representations of the thermophysical properties of the fluids is obtained by using the more-generic Mie form of interaction; in all but the simplest of fluids, one finds that the Lennard-Jones interaction is not the most appropriate.

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INTRODUCTION

Modelling the thermodynamics and fluid-phase equilibria of a system at the level of its constituent molecules is of central importance to chemical engineering and the applied physical sciences in general¹. As pointed out by Redlich and Kwong² more than sixty years ago, it is desirable and convenient to have an algebraic method based on a suitable equation of state (EOS) for this purpose. The primary interest of Redlich and Kwong lay in the provision of an empirical relation for the description of (supercritical) gases (liquids are barely mentioned in their manuscript), and the nature of their modification of the pioneering cubic EOS of van der Waals^{3,4} probably reflects this. With the passage of time the demands and interests of the engineering community have changed and evolved; Soave's modification⁵ of the Redlich-Kwong EOS allowed for better performance in relation to describing vapour-liquid equilibrium, while Peng and Robinson's EOS⁶ provided further improvement in relation to the hitherto poor descriptions of liquid densities with the Redlich-Kwong and Soave-Redlich-Kwong forms. Although a good description of fluid-phase equilibria in (p, V, T) space (where p is the pressure, V the volume and T the temperature) probably remains the most important goal, there has subsequently been an ever increasing demand for methodologies that allow for greater predictive capabilities. This is true not only in terms of describing the complex phase behaviour exhibited by the multicomponent mixtures that are frequently of interest in modern engineering applications, but also in terms of providing the reliable and simultaneous description of other key derived thermodynamic properties, such as heat capacities, speeds of sound and Joule-Thomson coefficients; moreover, one may require a description of fluid phase behaviour in terms of other thermodynamic variables – for example, in the analysis of refrigeration or power cycles, a representation in either (T, S) or (p, H)space is more appropriate (where S is the entropy and H the enthalpy).

Although many further developments have been made with respect to the modelling of fluids and fluid mixtures with cubic EOSs and the current-day description is frequently excellent (see, for example, the book of Kontogeorgis and Folas⁷ for a more-detailed discussion), one is ultimately constrained by the simple nature of the underlying molecular model, which remains, at heart, van der Waals' attracting hard-sphere molecular model. The need to be able to predict or, at least, adequately describe the thermodynamic properties of fluids of greater molecular functionality in progressively more-complicated processes has therefore

driven the development of increasingly sophisticated EOS, based on more physically realistic molecular models. Among these are the molecular-based family of EOSs developed within the framework of the statistical associating fluid theory (SAFT)^{8,9}, the origins of which are firmly grounded in a formal statistical mechanical treatment of continuum fluids^{10–15}. To capture the essentials of asymmetries in molecular shape, molecules are modelled as chains of tangentially bonded spherical monomeric segments, which interact with each other through a prescribed potential. Moreover, the directional nature of intermolecular interactions between associating molecules can be accounted for by incorporating bonding sites on the molecular cores, mimicking the hydrogen bonding in fluids such as water or methanol.

Within the basic SAFT EOS methodology, one constructs the free energy of the fluid in the form of a perturbation theory starting with that of a reference monomeric fluid and adding contributions relating to the changes in free energy that would be brought about by, for example, irreversibly joining the monomers into chains, and by allowing molecules to associate with each other. The nature and expression of the different terms corresponding to these perturbative contributions are specific to each particular version of SAFT; indeed, it is these differences that characterize the various incarnations of the EOS. Among the more popular are the original SAFT^{8,9}, Huang-Radosz (HR) SAFT¹⁶ (sometimes referred to as CK-SAFT), simplified SAFT¹⁷, SAFT-LJ (Lennard-Jones)¹⁸⁻²¹, SAFT-VR (Variable Range)^{22,23}, Soft SAFT^{24,25}, PC-SAFT (Perturbed Chain)^{26,27}, and simplified PC-SAFT²⁸. Group-contribution (GC) re-formulations of SAFT have been also developed by a number of groups^{29–38}; these EOSs are continuum-fluid descendants of the seminal GC approach developed by Prausnitz and co-workers³⁹ within a quasichemical (lattice-based) framework for the activity coefficient of mixtures⁴⁰. A discussion of the different versions of the SAFT EOS and the adequacy of representing the thermodynamics and fluid-phase equilibria of a broad variety of fluids and fluid mixtures with the methodology has now appeared in a number of excellent reviews^{7,41–45}. It is clear from the chronology of this list that since the inception of SAFT in the late 1980s there has been continuous effort to improve the theory. An important recent development in this regard is the SAFT-VR Mie EOS⁴⁶, and its GC offspring, SAFT- γ Mie³⁷.

The SAFT-VR Mie EOS⁴⁶ is an extension of the SAFT-VR^{22,23} approach in which the underlying Barker and Henderson high-temperature perturbation expansion of the monomer free energy is incorporated to third order and, in particular, where the interactions between

the monomeric segments are represented using Mie potentials⁴⁷, *i.e.*, generalized Lennard-Jones (LJ) forms (with variable repulsive and attractive exponents). The refinements to the theory result in an EOS with substantially improved performance for the fluid properties as compared with SAFT-VR prior to its reformulation^{22,23,48–50}, greatly enhancing the accuracy of the description of the near-critical region, while retaining the accurate modelling of second-derivative properties provided by the use of the Mie potential^{48–50}. Combined with a novel treatment of the association contribution to the free energy⁵¹, these advances allow for a high-fidelity global representation of the thermodynamic properties and fluid-phase equilibria of associating molecules and their mixtures.

The Mie $(\lambda_{\rm r}-\lambda_{\rm a})$ potential for two spherical segments separated by a distance r is given by $u^{\rm Mie}(r) = \mathcal{C}(\lambda_{\rm r},\lambda_{\rm a})\varepsilon \left[\left(\frac{\sigma}{r}\right)^{\lambda_{\rm r}} \left(\frac{\sigma}{r}\right)^{\lambda_{\rm a}} \right]. \tag{1}$

$$\mu^{\text{Mie}}(r) = \mathcal{C}(\lambda_{\text{r}}, \lambda_{\text{a}}) \varepsilon \left[\left(\frac{\sigma}{r} \right)^{\lambda_{\text{r}}} \left(\frac{\sigma}{r} \right)^{\lambda_{\text{a}}} \right].$$
(1)

Here, just as with the LJ (12-6) potential, σ is a size parameter representing the diameter of each spherical monomeric core, ε is an energetic parameter representing the depth of the potential well, $\lambda_{\rm r}$ and $\lambda_{\rm a}$ are the repulsive and attractive exponents, respectively, and the parameter $C(\lambda_{\rm r}, \lambda_{\rm a})$, which takes a value of 4 for the LJ potential, is defined as

$$\mathcal{C}(\lambda_{\rm r},\lambda_{\rm a}) = \frac{\lambda_{\rm r}}{\lambda_{\rm r} - \lambda_{\rm a}} \left(\frac{\lambda_{\rm r}}{\lambda_{\rm a}}\right)^{\frac{\lambda_{\rm a}}{\lambda_{\rm r} - \lambda_{\rm a}}},\tag{2}$$

such that the minimum of the potential well is at $-\varepsilon$. The two exponents of the Mie potential constitute two additional parameters as compared with the LJ potential (for which $\lambda_r = 12$ and $\lambda_a = 6$ are fixed) or one additional parameter as compared with the variable-range square-well (SW) potential (the interaction considered with the most frequently used formulation of SAFT-VR^{22,23}). Although the fluid-phase behaviour can be accurately described using a single fixed form of interaction, such as the LJ (12-6) potential, often one cannot obtain an accurate simultaneous description of the second-derivative properties, which are very sensitive to the precise nature of the repulsive interactions. Accordingly, the added versatile form provided by the use of the Mie potential offers a distinct advantage over the use of the simple LJ model.

On the other hand, the introduction of additional parameters raises issues relating to the possibility of model degeneracy: in parameterising the force field for a given fluid, one may

encounter a variety of different models that provide essentially equivalent performance. Issues of model degeneracy have already been recognized in the context of a SAFT description of associating fluids that require extra parameters to characterize the association interactions (see, *e.g.*, References^{52,53}). To best exploit the power of the SAFT-VR Mie EOS it is necessary to assess the nature of any model degeneracy and how it may be resolved, for example with an awareness of the sensitivities of the property predictions to variations in the parameter values or by taking advantage of known physics to assign parameter values; the mathematically optimum model may not always provide the best physical model⁵². For our current work with SAFT-VR Mie, it is of particular importance to gain an understanding of the role of the repulsive range parameter, λ_r .

The success of SAFT in general, and the main source of the predictive capability it provides, is predicated on its solid foundation in statistical mechanics, which has allowed the incorporation of more of the molecular-level physics of the system than was previously possible (for example, using cubic EOS). This is partly facilitated by the perturbative treatment of the free energy as a sum of individual contributions, which can be related to physical attributes such as molecular shape, or association interactions. The formulation of SAFT-VR for the Mie potential allows, in principle, even more detailed information relating to the physical interactions to be included. However, the danger is that one could, instead, lose contact with that physics by blindly manipulating the extra adjustable parameter(s) that have been introduced, thereby reducing the EOS simply to a very good correlative tool. In a separate publication⁵¹ this issue is examined in the context of associating fluids. The main purpose of our current contribution is to provide an understanding of the model-parameter space and how it relates to the performance of the SAFT-VR Mie EOS in describing thermodynamic properties of non-associating fluids, thereby avoiding the trap of simply providing correlations, and exploiting the tremendous predictive capabilities of this new EOS.

THEORY

The precise form developed for the SAFT-VR Mie EOS has been described in detail elsewhere⁴⁶; here we simply collect together the important expressions. The underlying molecular model of a non-associating fluid is that of a chain of m tangentially bonded segments, interacting via a Mie potential, as given in Equation (1). The general SAFT EOS

for a non-associating fluid is expressed as

$$A = A^{\text{ideal}} + A^{\text{monomer}} + A^{\text{chain}}, \qquad (3)$$

where A represents the Helmholtz free energy. Conceptually, to build up the equation one imagines first an ideal gas of particles. The second (monomer) term on the right-hand side of Equation (3) represents the change in free energy brought about by introducing interactions between the monomeric segments (thereby conferring size on them), and the final (chain) term represents the change in free energy brought about by grouping the segments as molecular chains (each comprising m monomeric segments). This grouping confers shape on the model molecule, so that the ideal contribution to the free energy (the first term) is not simply that of the point-particle ideal gas; rather, it represents the free energy of an ideal gas of the molecules, represented by $A^{\text{ideal}} = Nk_{\text{B}}T(\ln(\rho\Lambda^3) - 1)$, where N is the number of molecules, k_{B} is Boltzmann's constant, T is the absolute temperature, $\rho = N/V$ with V the volume, and Λ^3 is taken to represent the de Broglie volume which incorporates all of the kinetic (translational, rotational, and vibrational) contributions to the partition function. (We do not consider associating molecules in our current work, hence no term appears to account for the free energy of association.)

Monomer term

As with the original formulation of SAFT-VR²², the Helmholtz free energy of the monomer fluid is expressed as a high-temperature perturbation expansion, but this time taken to third order⁴⁶:

$$\frac{A^{\text{monomer}}}{Nk_{\text{B}}T} = m\left(a^{\text{HS}} + \frac{a_1}{k_{\text{B}}T} + \frac{a_2}{(k_{\text{B}}T)^2} + \frac{a_3}{(k_{\text{B}}T)^3}\right).$$
(4)

For pure components, the free energy of the reference HS system is obtained from the well-known Carnahan and Starling⁵⁴ EOS as

$$a^{\rm HS} = \frac{4\eta - 3\eta^2}{\left(1 - \eta\right)^2},\tag{5}$$

where $\eta = \rho_{\rm s} \pi d^3/6$ is the packing fraction of the reference hard-sphere system, and $\rho_{\rm s} =$

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 $N_{\rm s}/V$ is the segment density of the Mie fluid, with $N_{\rm s} = Nm$ the total number of segments. The Barker and Henderson⁵⁵ effective hard-sphere diameter d(T) has been introduced as

$$d(T) = \int_0^\sigma 1 - \exp\left(\frac{u^{\text{Mie}}(r)}{k_{\text{B}}T}\right) \mathrm{d}r\,,\tag{6}$$

to represent the repulsive reference system.

The first-order perturbation contribution of the SAFT-VR Mie EOS is given by

$$a_{1} = \mathcal{C}(\lambda_{\mathrm{r}}, \lambda_{\mathrm{a}}) \left[x_{0}^{\lambda_{\mathrm{a}}} \left(a_{1}^{\mathrm{S}}(\eta; \lambda_{\mathrm{a}}) + B(\eta; \lambda_{\mathrm{a}}) \right) - x_{0}^{\lambda_{\mathrm{r}}} \left(a_{1}^{\mathrm{S}}(\eta; \lambda_{\mathrm{r}}) + B(\eta; \lambda_{\mathrm{r}}) \right) \right], \tag{7}$$

where $x_0 = \sigma/d$,

$$B(\eta;\lambda) = 12\eta\varepsilon \times \left(\frac{(1-\eta/2)}{(1-\eta)^3} \left(\frac{1-x_0^{3-\lambda}}{\lambda-3}\right) - \frac{9\eta(1+\eta)}{2(1-\eta)^3} J_\lambda(\lambda)\right),\tag{8}$$

and

$$J_{\lambda}(\lambda) = -\frac{x_0^{4-\lambda}(\lambda-3) - x_0^{3-\lambda}(\lambda-4) - 1}{(\lambda-3)(\lambda-4)}$$
(9)

for both $\lambda = \lambda_{\rm a}$ and $\lambda = \lambda_{\rm r}$. The term $a_1^{\rm S}$ appearing in Equation (7) is the first-order perturbation term for a Sutherland fluid, which can be obtained as

$$a_1^{\rm S}(\eta;\lambda) = -12\varepsilon\eta \left(\frac{1}{\lambda-3}\right) \frac{(1-\eta_{\rm eff}(\eta;\lambda)/2)}{(1-\eta_{\rm eff}(\eta;\lambda))^3},\tag{10}$$

with

$$\eta_{\text{eff}}(\eta;\lambda) = c_1(\lambda)\eta + c_2(\lambda)\eta^2 + c_3(\lambda)\eta^3 + c_4(\lambda)\eta^4, \qquad (11)$$

where the coefficients c_1 , c_2 , c_3 , and c_4 are given in equation (41) of Reference⁴⁶.

The second perturbation term is given by

$$a_{2} = \frac{K^{\mathrm{HS}}}{2} (1+\chi) \varepsilon \mathcal{C}^{2}(\lambda_{\mathrm{r}}, \lambda_{\mathrm{a}}) \left[x_{0}^{2\lambda_{\mathrm{a}}}(a_{1}^{\mathrm{S}}(\eta; 2\lambda_{\mathrm{a}}) + B(\eta; 2\lambda_{\mathrm{a}})) - 2x_{0}^{\lambda_{\mathrm{a}}+\lambda_{\mathrm{r}}}(a_{1}^{\mathrm{S}}(\eta; \lambda_{\mathrm{a}}+\lambda_{\mathrm{r}}) + B(\eta; \lambda_{\mathrm{a}}+\lambda_{\mathrm{r}})) + x_{0}^{2\lambda_{\mathrm{r}}}(a_{1}^{\mathrm{S}}(\eta; 2\lambda_{\mathrm{r}}) + B(\eta; 2\lambda_{\mathrm{r}})) \right],$$

$$(12)$$

where K^{HS} is the isothermal compressibility of the hard-sphere reference fluid, obtained from

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and

the Carnahan and Starling^{54} expression for the compressibility factor as

$$K^{\rm HS} = \frac{(1-\eta)^4}{1+4\eta+4\eta^2-4\eta^3+\eta^4},\tag{13}$$

$$\chi = f_1(\alpha)\eta x_0^3 + f_2(\alpha) \left(\eta x_0^3\right)^5 + f_3(\alpha) \left(\eta x_0^3\right)^8.$$
(14)

The functions $f_i(\alpha)$ characterizing the correction factor χ are obtained as

$$f_i(\alpha) = \frac{\sum_{n=0}^3 \phi_{i,n} \alpha^n}{1 + \sum_{n=4}^6 \phi_{i,n} \alpha^{n-3}},$$
(15)

where the dimensionless van der Waals-like integrated attractive energy α is defined as

$$\alpha = \mathcal{C}(\lambda_{\rm r}, \lambda_{\rm a}) \left(\frac{1}{\lambda_{\rm a} - 3} - \frac{1}{\lambda_{\rm r} - 3}\right), \qquad (16)$$

and the coefficients $\phi_{i,n}$ are taken from table II of Reference⁴⁶.

The third perturbation term is also expressed in terms of the functions $f_i(\alpha)$ as

$$a_3 = -\varepsilon^3 f_4(\alpha) \eta x_0^3 \exp\left(f_5(\alpha) \eta x_0^3 + f_6(\alpha) \eta^2 x_0^6\right).$$
(17)

Chain term

The chain term is expressed in the usual SAFT form as

$$\frac{A^{\text{chain}}}{Nk_{\text{B}}T} = -(m-1)\ln g^{\text{Mie}}(\sigma).$$
(18)

The radial distribution function (RDF) of the Mie fluid at contact is obtained as a second order expansion:

$$g^{\text{Mie}}(\sigma) = g_d^{\text{HS}}(\sigma) \exp\left[\frac{g_1(\sigma)}{g_d^{\text{HS}}(\sigma)}\frac{\varepsilon}{k_{\text{B}}T} + \frac{g_2(\sigma)}{g_d^{\text{HS}}(\sigma)}\frac{\varepsilon^2}{(k_{\text{B}}T)^2}\right].$$
(19)

The RDF of the reference fluid comprising hard-spheres of diameter d evaluated at σ is calculated from

$$g_d^{\rm HS}(\sigma) = \exp\left[k_0 + k_1 x_0 + k_2 x_0^2 + k_3 x_0^3\right],\tag{20}$$

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wherein the coefficients k_i are given by

$$k_0 = -\ln(1-\eta) + \frac{42\eta - 39\eta^2 + 9\eta^3 - 2\eta^4}{6(1-\eta)^3}, \qquad (21)$$

$$k_1 = \frac{-12\eta + 6\eta^2 + \eta^4}{2(1-\eta)^3}, \qquad (22)$$

$$k_2 = \frac{-3\eta^2}{8(1-\eta)^2},\tag{23}$$

and

$$k_3 = \frac{3\eta + 3\eta^2 - \eta^4}{6(1 - \eta)^3} \,. \tag{24}$$

The first-order perturbation term in Equation (19) is obtained from

$$g_{1}(\sigma) = \frac{1}{2\pi\varepsilon d^{3}} \left[3\frac{\partial a_{1}}{\partial \rho_{s}} - \mathcal{C}(\lambda_{r},\lambda_{a})\lambda_{a}x_{0}^{\lambda_{a}}\frac{a_{1}^{S}(\eta;\lambda_{a}) + B(\eta;\lambda_{a})}{\rho_{s}} + \mathcal{C}(\lambda_{r},\lambda_{a})\lambda_{r}x_{0}^{\lambda_{r}}\frac{a_{1}^{S}(\eta;\lambda_{r}) + B(\eta;\lambda_{r})}{\rho_{s}} \right], \qquad (25)$$

and the second-order perturbation term from

$$g_{2}(\sigma) = \frac{1+\gamma_{c}}{2\pi\varepsilon^{2}d^{3}} \left\{ 3 \frac{\partial a_{2}/(1+\chi)}{\partial\rho_{s}} - \varepsilon K_{\rm HS}\mathcal{C}^{2}(\lambda_{\rm r},\lambda_{\rm a}) \left(\lambda_{\rm r}x_{0}^{2\lambda_{\rm r}}\frac{a_{1}^{\rm S}(\eta;\lambda_{\rm r}) + B(\eta;\lambda_{\rm r})}{\rho_{\rm s}} + (\lambda_{\rm r}+\lambda_{\rm a})x_{0}^{\lambda_{\rm r}+\lambda_{\rm a}}\frac{a_{1}^{\rm S}(\eta;\lambda_{\rm r}+\lambda_{\rm a}) + B(\eta;\lambda_{\rm r}+\lambda_{\rm a})}{\rho_{\rm s}} + \varepsilon K_{\rm HS}\mathcal{C}(\lambda_{\rm r},\lambda_{\rm a})^{2}\lambda_{\rm a}x_{0}^{2\lambda_{\rm a}}\frac{a_{1}^{\rm S}(\eta;\lambda_{\rm a}) + B(\eta;\lambda_{\rm a})}{\rho_{\rm s}} \right) \right\},$$
(26)

where

$$\gamma_{c} = \phi_{7,0} \left(-\tanh\left(\phi_{7,1}(\phi_{7,2} - \alpha)\right) + 1\right) \eta x_{0}^{3} \\ \times \left(\exp\left(\frac{\varepsilon}{k_{\rm B}T}\right) - 1 \right) \exp\left(\phi_{7,3}\eta x_{0}^{3} + \phi_{7,4}\eta^{2} x_{0}^{6}\right) \,.$$
(27)

The coefficients $\phi_{7,i}$ (i = 0, 3, and 4) are available from table II of Reference⁴⁶.

Fluid-phase coexistence requires the conditions of thermal, mechanical and chemical equilibrium, which can be solved for a pure component by ensuring that the temperature, pressure, and chemical potential are equal in each phase. Having specified the explicit form of the Helmholtz free energy, the pressure $p = -(\partial A/\partial V)_{N,T}$, chemical potential $\mu = (\partial A/\partial N)_{V,T}$, and other thermodynamic properties can be obtained algebraically from the standard ther-

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modynamic relations⁵⁶. As well as the fluid-phase equilibrium properties, in our current paper we assess the adequacy of the SAFT-VR Mie EOS in describing second-order derivative properties including the isobaric heat capacity c_p , the speed of sound u, and the Joule-Thomson coefficient $\mu_{\rm JT}$.

METHOD

In the SAFT-VR Mie approach, one represents molecules as tangentially bonded spherical segments interacting through Mie pair potentials so that the parameters defining the model of a molecule (in the case of non-associating fluids) comprise the number of segments m forming the model molecule, and the parameters of the Mie potential, σ , ε , $\lambda_{\rm r}$, and $\lambda_{\rm a}$. As is customary in the development of models for use with an EOS, the values of these parameters are estimated using experimental pure-component information; one adjusts the values to minimize the difference between the experimental and calculated properties. We choose to minimize an objective function of the following form:

$$F_{\rm obj} = \sum_{X} \left(\frac{\omega_X}{n_X} \sum_{k=1}^{n_X} \left[\frac{X_k^{\rm exp} - X_k^{\rm calc}}{X_k^{\rm exp}} \right]^2 \right) , \qquad (28)$$

where n_X is the number of datapoints relating to a given property X, ω_X is the weight assigned to property X, and k is a specific datapoint; a Levenberg-Marquardt algorithm is employed^{57,58} for the minimization procedure. A convenient measure of the quality of the description with the model is the percentage absolute average deviation (%AAD) from experimental values of a property, calculated as:

$$\% AAD(X) = \frac{100}{n_X} \sum_{k=1}^{n_X} \left| \frac{X_k^{\exp} - X_k^{\text{calc}}}{X_k^{\exp}} \right|.$$
(29)

One can incorporate different properties in the objective function. Whereas parameter estimation for the previous formulation of SAFT-VR (using the SW potential) was mostly restricted to the description of vapour-liuqid equilibria including the vapour pressure and saturated-liquid density – as is common practice for model development in relation also to other EOSs – the ability of SAFT-VR Mie to provide quantitative agreement for different second-derivative properties such as the speed of sound and heat capacity^{46,48–50} allows one

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to include other properties in the objective function. A variation of the respective weights of the properties will result in different "optimal" estimated models, each tailored according to important features of the task at hand. One must choose not only the properties to include and their relative weights but, also, the range of experimental conditions (*e.g.*, in temperature and pressure); this will also impact the % AADs and, thereby, the model highlighted as "optimal", so should be selected with care. (We note in passing that it is important when using % AADs as a basis for comparison of different models, that the experimental data sets against which the % AADs are calculated are identical; if they are not, the comparison can be somewhat misleading.)

In order to investigate the parameter space we extend the method developed in Reference⁵² and used subsequently in Reference⁵³; both these studies were carried out with the previous formulation of SAFT-VR using the SW potential, given by

$$u^{\rm SW} = \begin{cases} +\infty & r < \sigma^{\rm SW} \\ -\varepsilon^{\rm SW} & \sigma^{\rm SW} \le r < \lambda^{\rm SW} \sigma^{\rm SW} \\ 0 & r \ge \lambda^{\rm SW} \sigma^{\rm SW} \end{cases}$$
(30)

where σ^{SW} , ε^{SW} , and λ^{SW} represent size, energy, and range parameters of the model. This method consists of performing multiple optimizations to obtain models for the same fluid with the same objective function, but using different fixed values for two of the parameters in each. Every point in the parameter-space plane formed by these two parameters then represents a model that is optimal in the remaining parameters, and different properties of the model, such as the % AAD or the objective function, can be easily represented using contour plots projected onto the plane defined by the two fixed parameters. In studies on water⁵², an almost identical representation of the properties is obtained with a number of different models, for broad intervals of the interaction energies; this degeneracy of parameters led to difficulty in identifying the best model. In the context of water models (of the type used in SAFT) one of the most useful characteristics is the ratio of the dispersion and associating contributions; accordingly, the plane selected was that defined by the parameters representing the depths of the dispersion and association $energies^{52}$. On the other hand, dos Ramos *et al.*⁵³ considered different pairs of parameters, (m, λ^{SW}) and $(\sigma^{SW}, \lambda^{SW})$ for non-assocating molecules, and $(\varepsilon^{\text{HB}}, \lambda^{\text{SW}})$ for water, *i.e.*, representing a geometric and an energetic parameter: by fixing one parameter (in this case m) to a specific value a more

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clearly identifiable optimal model can be obtained. In our current work, we extend the idea further, considering initially all ten of the parameter-space planes defined by pairs of the five model parameters of the molecules formed from Mie segments (m, ε , σ , $\lambda_{\rm a}$ and $\lambda_{\rm r}$). A case study in the development of such models is first undertaken for a carefully chosen fluid, to examine in detail the nature of the parameter space. Based on understanding gained in this case study, a subset of these planes is then chosen for the subsequent development of models of other fluids.

It remains only to select a suitable fluid on which to base the case study. The fluid selected for this purpose should be one for which the SAFT-VR Mie EOS is expected to provide an accurate representation of the thermophysical properties, for which one can readily judge the physical significance (or lack thereof) of model parameters, and for which reliable experimental data are plentiful. Methane is an ideal candidate for such a study, fulfilling all these criteria. A model for methane has already been developed for use with the SAFT-VR Mie EOS which provides an excellent description of the vapour-liquid equilibrium (VLE) and single-phase properties⁴⁶. The leading multipole moment of methane is the octupole; this is not expected to make a significant contribution to the intermolecular interactions, which are therefore expected to be of London-dispersion form, with $\lambda_{\rm a} = 6$. The spherical-top symmetry of methane suggests that it can be treated as a near-spherical molecule so that $m \sim 1$ is expected; moreover, from quantum-mechanical calculations⁵⁹ one expects an average molecular diameter $\sigma \sim 3.7$ Å. Although physically sensible values for the remaining two parameters, ε and $\lambda_{\rm r}$, cannot so readily be anticipated, one would intuitively expect $\varepsilon/k_{\rm B}$ to be of the order of a few hundred Kelvin. The ability to return physically reasonable parameter values close to these will provide a good test of our parameter estimation procedure based on the individual planes, and a means for the selection of the appropriate plane or planes to use in subsequent model development.

RESULTS

Developing models for methane: detailed investigation of the parameter space

Estimations of model parameters for methane are performed for the ten different planes defined by pairs taken from the five model parameters: $(\varepsilon, \lambda_{\rm r})$; $(\varepsilon, \lambda_{\rm a})$; $(\lambda_{\rm a}, \lambda_{\rm r})$; $(m, \lambda_{\rm r})$;

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 $(m, \lambda_{\rm a})$; $(\sigma, \lambda_{\rm r})$; $(\sigma, \lambda_{\rm a})$; (ε, m) ; (ε, σ) ; and (m, σ) . The values of the objective function (Equation (28)) and %AAD (Equation (29)) are calculated using the correlated experimental data from the National Institute of Standards and Technology (NIST)⁶⁰; different properties are chosen, including saturation properties (the vapour pressure and saturatedliquid density), the single-phase density, and the speed of sound (21 points are taken in the range T = 95 to 185 K for each of the VLE properties, and 30 points for each of the single-phase properties, at conditions ranging from T = 100 to 190 K and p = 10 to 50 MPa).

The objective-function landscape for the SAFT-VR Mie models of methane in relation to each of the planes is characterized by two distinct valleys, the minima of which correspond to the same two models irrespective of the plane. A selection of contour plots relating to these are provided for illustration in Figure 1; for this examination the objective function comprised the vapour pressure and saturated-liquid density (with equal weights, $\omega = 1$). Immediately it is apparent that one of these valleys corresponds to physically more-realisitic models, with $\sigma \sim 3.7$ Å and $m \sim 1$, while the models in the other valley feature $\sigma > 4$ Å and m < 0.8; the latter, in particular, is non-physical, since m = 1 (representing a single spherical segment) is a lower bound. Models in both valleys are characterized by values of $\lambda_{\rm a} \sim 5$; reassuringly, this is close to the London-dispersion value, although it is notable that the deviation, while small, leads to values lower and not higher than six.

In each of these estimations, two parameters defining the individual plane are fixed while the remaining three are free to adopt whatever values provide the optimal model. However, in certain circumstances the values of one or more of these parameters may be determined by physical constraints. For example, as already noted, to be commensurate with the approximate spherical symmetry of the methane molecule, m = 1 is appropriate. Having already observed that only one of the two valleys observed in each of the contour plots given in Figure 1 corresponds to $m \sim 1$, one would expect that assigning m = 1 would have the effect of removing one of the valleys, thereby simplifying the objective-function landscape.

In Figure 2 we provide a comparison between the SAFT-VR Mie models obtained in the case m is free (Figure 2(a)) and the case where m = 1 is assigned (Figure 2(b)); the behaviour of an expanded objective function comprising the vapour pressure, saturated liquid density, single-phase compressed-liquid density (each given weight $\omega = 1$ in Equation (28)), and the speed of sound (with weight $\omega = 0.25$) is projected onto the (λ_a, λ_r) plane. From a comparison of Figures 2(a) and (b) it is evident that the objective-function landscape is

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altered by fixing m, and that the resulting high-quality models are restricted to a narrower portion of parameter space, in accordance with the findings of dos Ramos *et al.*⁵³. Moreover, as expected, assigning m = 1 indeed simplifies the objective-function landscape, removing one of the valleys. Qualitatively similar results are obtained by fixing the attractive exponent to the London value of $\lambda_a = 6$ though, of course, one projects the landscape onto a different plane of parameter space.

In passing, it is interesting to note from Figure 2(a) that the inclusion of extra properties (namely, the single-phase density and speed of sound) in the objective function has resulted in one of the two valleys from the equivalent plot in Figure 1(b) being shifted to higher values of $\lambda_{\rm a}$, and that it now encompasses the London-dispersion value $\lambda_{\rm a} = 6$.

For our study of the SAFT-VR Mie models of methane we had in advance some sense of physically sensible parameter values, specifically m = 1 and $\lambda_{\rm a} = 6$, however this will certainly not be the case for all molecules, and more-general strategies for judging parameters are desirable. One such strategy would be to consider a representation of the molecules at the van der Waals' level, *i.e.*, in terms of the corresponding excluded volume and integrated energy. It is not straightforward to determine the excluded volumes for the chains of segments that comprise models of molecules in SAFT, however the volume occupied by the molecules is proportional to $\mathcal{V} = m\sigma^3$, whereby \mathcal{V} is a more convenient measure to consider. In Figure 3 the values for \mathcal{V} of methane models obtained in the examination of the $(\lambda_{\rm a}, \lambda_{\rm r})$ plane (in which m is not constrained) are illustrated; see also Figure 1(b) for the objective-function landscape corresponding to this examination. The variation in \mathcal{V} is less than 10% across the whole plane, demonstrating that no matter which Mie potential is used, the volume of all models of methane enabling a good description of the properties must lie within a narrow range. Moreover, by comparing this figure with Figure 1(b), within the valley corresponding to the best models, it can be seen that the variation of \mathcal{V} is less still. (The same conclusions can be drawn from an analogous inspection of molecular volumes corresponding to the landscape illustrated in Figure 2(a), relating to the objective function incorporating the single-phase liquid density and speed of sound for methane in addition to the saturation properties.)

The integrated van der Waals attractive energy for a chain of m segments can be char-

acterized in terms of the parameter α introduced earlier in Equation (16) as

$$E_{\rm int} = m^2 \sigma^3 \varepsilon \alpha. \tag{31}$$

The energy $E_{\rm int}$ spans over a wider range than \mathcal{V} (~ 100%), mainly because α varies almost threefold in this region. The importance of α can be seen in Figure 4, wherein contours representing the % AAD of the experimental isobaric heat capacity c_p from the corresponding values calculated with the SAFT-VR Mie EOS are plotted in the (λ_a, λ_r) plane along with contours representing constant values of α . The close correspondence between the two sets of contours indicates that models providing equivalent representations of the heat capacity correspond to similar values of α . In fact, by comparison of the contours in Figure 4 with those in Figures 1(a) and 2(a) and (b), it is clear that the curvatures of the lines of constant α also broadly follow the contours of the optimum-parameter-space landscape. Single-segment Mie fluids with the same value of α have been shown to be conformal,⁶¹ even though Mie fluids with different values of one or both of the exponents λ_r and λ_a are not generally conformal. This suggests that one can use α to inform the selection of an appropriate form of the Mie potential to characterize a model; for example, should a parameter estimation return an otherwise excellent model that has an unphysical value of one or both of the exponents, it indicates that this model can be replaced by another more-physical model, without serious detriment to the description of c_p or, indeed to performance of the model in general, provided it has the same value of α . In Reference⁶¹ a relation between α and the range of the fluid as described by the ratio of the critical-point temperature to that of the triple-point, is shown for the case of spherical models; its use to develop coarse-grained potentials is highlighted in that work.

As a final observation in relation to Figure 4, one can discern from a comparison with Figure 1(b) (in which the same objective function is used) that the models providing the very best description of the heat capacity are not those that give the best description of saturation properties. Importantly, however, there is some overlap in which both sets of properties are well described – this means that although one should tailor one's model if a property such as c_p is of the utmost importance, one clearly does not need to sacrifice a good representation of the vapour-liquid equilibria, nor vice versa.

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At this point, it is helpful to collect the insights gained in the foregoing discussion. In particular:

- As expected, the SAFT-VR Mie EOS provides for an excellent overall description of many of the thermodynamic properties of methane.
- Allowing the two exponents, λ_r and λ_a, to vary freely leads to an increasing complexity of the optimization.
- An appropriate form of the Mie potential can be identified by considering the integrated energy parameter α , as illustrated in Figure 4.
- The objective function surface is very sensitive to the properties included, together with their respective weights, and to whether or not any parameters are fixed. As a consequence, the objective function surface can be very simple with a single minimum or very complex with multiple local minima.
- Each plane in the multidimensional parameter space yields essentially equivalent results; consequently, one need investigate only a single plane for a pair of parameters in selecting a intermolecular-potential model.
- Different intermolecular-potential models are required to provide the very best descriptions of different properties, although parameter sets obtained by targeting a particular property may nevertheless perform well in respect of other properties.

In the remainder of the discussion we will focus on obtaining models that are able to provide for a good description of vapour-liquid equilibrium properties, *i.e.*, the models are selected based on their ability to represent the vapour pressure and saturated-liquid density of the pure components. A good description of the vapour pressure is an extremely important precursor to modelling multi-component mixture phase behaviour since the phase envelope of the mixture tends towards the vapour-pressure curve in the pure-component limit; any inaccuracy in the vapour pressure will result either in a poor description of the mixture phase behaviour or lead to unrealistic mixture parameters to compensate for bad pure-component models.

Developing SAFT-VR Mie models for other fluids

Models based on LJ (12-6), Mie (λ_r -6), and generic Mie (λ_r - λ_a) potentials

To assess the overall performance of the SAFT-VR Mie EOS, we now develop intermolecularpotential models for a wide range of compounds. The experimental data used in the parameter estimation are taken from the NIST⁶⁰ correlations in most cases except for the alkyl benzenes where the data are taken from References⁶²⁻⁶⁶. The conformal behaviour of single-segment Mie fluids with the same value of α presented in Reference⁶¹ is again confirmed in Figure 4, for methane; this suggests that one can reproduce the behaviour of the generic Mie (λ_r - λ_a) fluid using a Mie fluid of the same α with one of the exponents fixed to any value. In addition to our investigations in which both exponents are left free, we choose also to focus on the Mie (λ_r -6) family of fluids, characterized by a fixed value of the attractive exponent $\lambda_a = 6$. The value is chosen to match the scaling exponent of the London dispersion interaction⁶⁷, and can be reasonably taken as physically relevant for many fluids. To provide a reference for comparison, we consider also the more-familiar LJ (12-6) form of the Mie potential. We select the (λ_a, λ_r) plane as that on which to base our search for models. This choice enables us to highlight the advantages of the generic Mie potential over the more-widely used LJ (12-6) potential.

We focus first on carbon dioxide. The three models of carbon dioxide developed are based on a Lennard-Jones (12-6), a Mie (λ_r -6), and a generic Mie (λ_r - λ_a) potential form. The parameters for each model are estimated using the same objective function (comprising the vapour pressure and saturated-liquid density of carbon dioxide, with equal weights, $\omega = 1$); in each case the number of segments, m, is excluded from the parameter estimation and is assigned the fixed value m = 1.6936, which is estimated from quantum mechanical calculations for the size and non-sphericity parameters⁵⁹.

The performance of the different models used to represent carbon dioxide in relation to saturation properties is illustrated in Figure 5. The model featuring a Lennard-Jones (12-6) potential does not perform as well as the Mie (λ_r -6) model with a fixed attractive exponent of six, which itself performs less well than the Mie (λ_r - λ_a) model for which both exponents are estimated. One would, of course, expect such a result simply on the basis of the number of parameters: the model with the most adjustable parameters performs best. Accordingly,

it is important to consider the respective adequacies of the models to assess whether the improvement in performance is due solely to the use of extra adjustable parameters.

The Mie $(\lambda_r \cdot \lambda_a)$ model obtained with variable repulsive λ_r and attractive λ_a exponents provides an excellent description of the VLE of carbon dioxide, but the physical significance of the exponents is not clear. This model is characterized by a value of $\lambda_a = 5.055$; the two leading contributions to the attractive intermolecular interactions of carbon dioxide are the London dispersion (which scales as $1/r^6$) and the quadrupole-quadrupole interaction (which scales as $1/r^8$), hence it is difficult to account for an attractive exponent lower than six. A simple analysis of this type is however complicated by the fact that carbon dioxide is not spherical and possesses three atomic centres. Overall the Mie (λ_r -6) model model offers a good compromise between the quality of the representation of the thermodynamic properties, the number of adjustable parameters, and the physical significance of these parameters.

In relation to the comparison between the different underlying potentials, similar results to those seen for carbon dioxide are obtained for other fluids. Accordingly, we favour in general those based on the Mie (λ_r -6) potential, which carry the London-dispersion attractive exponent. The SAFT-VR Mie parameters for the models obtained for the different fluids examined are presented in Tables I to III, along with the % *AAD* values for different properties. The models relating to the LJ (12-6) and generic Mie (λ_r - λ_a) potentials are presented in an appendix (Tables V to X).

Fluid models based on Mie (λ_r-6) potentials

The representation of the experimental data obtained using the SAFT-VR Mie EOS is found to be very good. This is illustrated for the case of Mie (λ_r -6) models for the VLE (saturation vapour and liquid densities, vapour pressure and enthalpy of vaporization) of *n*-alkanes from methane to *n*-decane and *n*-dodecane in Figure 6, and for other common fluids in Figure 7 (the model parameters can be found in Tables I to III). The quality of the description of the enthalpy of vaporization (Figures 6(*c*) and 7(*c*)) is particularly gratifying. We note also that the versatility of the models and theory is evident from the diversity of molecules that are very well described.

A noteworthy feature common to all the models is a relatively small overestimate of the critical point, compared to that obtained with the previous formulations of SAFT-VR

and with other SAFT EOS; this improved performance in the near-critical region enables a much better description of the saturation densities. This feature will simplify the modelling of mixtures where one or more components is just supercritical. Conditions at which the real fluid is supercritical but at which the model fluid is subcritical necessarily create difficulties in the description of mixtures; for example, phase diagrams in composition space have a fundamentally different shape when one or more component becomes supercritical.

It is interesting to examine the variation in the repulsive exponent, $\lambda_{\rm r}$, obtained for the different fluids; the exponent takes values ranging from less than 9 for O₂ to ~37 for SF₆. The very hard repulsive nature of the potential obtained for SF₆ is rationalized by considering the highly electronegative nature of fluorine, whereby one expects very strong repulsions between fluoride ligands on separate molecules; in similar fashion, a very steeply repulsive potential was required to obtain a good representation of CF₄⁴⁶. The values of $\lambda_{\rm r}$ for the remaining molecules lies generally in the range (9 $\leq \lambda_{\rm r} \leq 20$). The exception is dimethyl propane, for which $\lambda_{\rm r} \sim 32$ is obtained; here the hardness of the potential may reflect the steric hindrance to molecular deformation of the closely positioned methyl groups on the molecule, although we note that the shape parameter was constrained to a value m = 1.0 during the parameter estimation so the anomalously high value of $\lambda_{\rm r}$ could also indicate that this restriction is too severe.

For many years it has been standard practice to employ the standard LJ (12-6) potential, almost by default. With this in mind, if one compares the models presented in Tables I to III with those in Tables V to VII (in the Appendix), for which the LJ (12-6) form is enforced, some of the Mie (λ_r -6) models perhaps appear surprising in terms of the magnitudes of λ_r and ε one encounters. However, this does not mean that our featured Mie models are non-physical. On the contrary; the improved predictive power of the models evidenced by the reduction in AAD% from experimental data for second-derivative properties (in particular, for speeds of sound) suggests that the models have a firmer physical foundation. As a final comment on the values of obtained for λ_r it is interesting to find that, with the exception of neon (which is influenced by quantum effects and imperfectly described using LJ (12-6) without quantum correction⁶⁸⁻⁷⁰) the noble gases are generally well described using a potential that is essentially of the LJ (12-6) form.

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Versatility of models obtained using vapour-liquid saturation properties

As illustrated in previous work^{46,48,49,71}, the SAFT-VR Mie EOS offers the simultaneous capability of an accurate description of the vapour-liquid equilibrium and second-derivative properties (such as speed of sound or heat capacity) of fluids. This feature is very useful as it widens the applicability of the EOS, as well as providing confidence that the theory and the resulting models are physically sound. Although the (λ_r -6) Mie models we obtain are based on an objective function comprising the vapour pressures and saturated-liquid densities (with equal weight), a good overall description of the single-phase densities and second-derivative properties of many different fluids is provided, as can be seen from the % AAD from experimental data⁶⁰ given in Tables I to III (analogous tables relating to the LJ (12-6) and generic Mie (λ_r - λ_a) models are given in the appendix).

Further demonstration is provided graphically in Figures 8, 9, and 10, in which descriptions of single-phase isobaric properties of carbon dioxide, xenon, and isopentane (respectively) are provided for illustration; the properties examined include the single-phase density, the Joule-Thomson coefficient, the isobaric heat capacity, and the speed of sound. (Essentially equivalent performance is found using the models obtained for the other fluids.) As can be seen from these figures, the descriptions of the various properties provided by these models are excellent throughout, notwithstanding that none of these properties were included in the objective functions used in parameter estimation; as discussed earlier for methane, an enhanced description may be obtained if the properties are targeted in the parameter estimation (though probably at the expense of a slightly poorer description of the VLE). It is also interesting to note that, in separate work⁷² employing models for *n*-butane and *n*-decane developed previously in Reference⁴⁶ (also provided in Table I for completeness), excellent predictions were obtained for the fluid-phase equilibria of these two compounds in both the temperature-entropy (*T*, *S*) and pressure-enthalpy (*p*, *H*) projections.

Homologous series: trends in the characteristic parameters of the models

When appropriately combined, the parameters used in SAFT-VR Mie for molecules in homologous series show clear physical trends, reflecting the molecular basis of the theory. This was first explored within the context of SAFT by Huang and Radosz¹⁶, who examined

the molecular weight (MW) dependence of the molecular length m, segment volume, and segment energies for alkanes and polynuclear aromatics. The idea was exploited to obtain parameters for polyethylene by correlating the trends observed for low-MW alkanes and extrapolating to high MW. Similar procedures have been adopted to procure model parameters for polymers within other SAFT frameworks, for example, in References⁷³ and⁷⁴ (for SAFT-VR),²⁶ and⁷⁵ (for PC-SAFT), and⁷⁶ (for soft-SAFT). The numerous successful studies of thermodynamic properties of polymers in which such models have been used collectively demonstrate the veracity of the concept; these include, for example, the studies presented in References^{26,73–88}.

In our case we focus on the *n*-alkanes using the Mie (λ_r -6), Lennard-Jones (12-6), and generic Mie $(\lambda_r - \lambda_a)$ models presented in Tables I, V, and VIII, respectively, and the *n*-alkyl benzenes using the Mie ($\lambda_{\rm r}$ -6) models given in Table II (note that we have not developed LJ or generic Mie models for the alkyl benzenes). The individual parameters themselves do not necessarily follow clear relationships, although there is an increasing trend in the values of $\lambda_{\rm r}$ as evidenced in table I for the *n*-alkanes, excluding methane. The fundamental physics that allows these models to perform so well reappears when considering appropriate combinations of parameters. A useful combination of parameters is the molecular volume, characterized by $\mathcal{V} = m\sigma^3$, which is found to be linear with the number of carbon atoms for the three families of models for the *n*-alkanes and also for the Mie(λ_r -6) models of the *n*-alkyl-benzenes, as seen in figure 11a. This volumetric parameter takes into account the very strong coupling between the chain length m and the segment diameter σ , and is found to be quite constant for a given component, as seen previously in Figure 3. The other physically meaningful parameter combination of interest is the integrated van der Waals-like attractive energy E_{int} of the molecule, defined in Equation (31). This energetic parameter is found to vary as a second-degree polynomial of the number of carbons, as can be seen in Figure 11b: such a quadratic dependence is expected from the dependence of $E_{\rm int}$ on the square of the molecular length, m^2 . Although one cannot directly determine a model using these relationships as there are more parameters than equations, they can be used to determine a good starting guess for the estimation. More importantly, the existence of the trends is a further testament to the manner in which theory and models capture the physical characteristics of the molecules.

DISCUSSION AND CONCLUSIONS

It was demonstrated in Reference ⁴⁶, using detailed comparisons with molecular simulation, that the new formulation of SAFT-VR Mie performs extremely well in describing the physics of Mie (generalized Lennard-Jones) fluids; the close agreement between thermodynamic properties calculated with the theory and those obtained in simulations is remarkable. However, although excellent models for real fluids were developed, a detailed examination of the model parameter space was not conducted and, accordingly, it remained to gain the insight necessary to be able to best exploit the enhanced capability of the new theory in terms of developing models to represent real fluids. In our current work this has been addressed in relation to non-associating fluids; our extension of this investigation to associating fluids is presented in a separate publication⁵¹.

To identify the most-appropriate models for real fluids one requires an appreciation of the sensitivity of calculated properties to changes in the values of the intermolecular potential model parameters; in related fashion, one needs also to understand and be able to resolve any issues associated with model degeneracy, whereby different models provide apparently equivalent performance. Degeneracy has been reported previously with respect to models based on the square-well potential (with three adjustable parameters) 52,53 , and is therefore expected when using models based on the Mie potential, which is characterized by four parameters. Here we have carried out a detailed examination of the parameter space for models of non-associating molecules, as exemplified in contour plots of the landscapes of objective functions (based on the square residual differences between experimental and calculated properties) projected onto different planes of parameter space. Even for an apparently simple molecule such as methane, the parameter space is seen to be quite complex, and therefore identifying "the best" model for a compound is not straightforward. Model degeneracy is indeed encountered; the objective-function landscape features valleys within which many high-fidelity models may be located. The number of these valleys, and their location, depends on the properties included in the objective function; consequently, even though performance is generally good over a wide range of properties, the optimal descriptions of particular properties are obtained using property-specific models. Additionally, these valleys may extend over large regions of the landscape and although a mathematically optimal parameter set may be found within (as judged by the value of the objective

function), this model may be characterized by physically unreasonable values of molecular parameters, whereas the valleys may contain other only marginally poorer models characterized by physically sensible parameter values – models that may themselves have been "optimal" given a slightly different choice of the form of the objective function. Thus it is important not to rely blindly on a simple optimization to estimate parameters for an intermolecular potential model of a fluid. Rather, one should utilize knowledge of the physical nature of the molecule to inform the estimation procedure; for example, in the case of a simple molecule such as methane a physically reasonable model is expected to be characterized by an attractive exponent of $\lambda_a = 6$, following London dispersion, and m = 1, *i.e.*, a single spherical core – and we have seen that assigning these values does indeed assist in discriminating between the many good models of methane. From a different perspective, this means that one can assign parameters based on physical considerations and still obtain excellent models.

To examine the objective-function landscape in every possible plane of the parameter space is a labour-intensive and time-consuming task. From our examination of the parameter space of models of methane, we find that it is sufficient to consider only a single plane; this conclusion is born out in subsequent studies of other fluids. We favour the (λ_a, λ_r) plane defining the attractive and repulsive exponents of the Mie potential form, which facilitates simpler comparisons with models based on the more-familiar Lennard-Jones (12-6) potential. It has been shown in Reference⁶¹ that two spherical Mie fluids of the same α (see equation 16) have the same free energy, and hence the same thermodynamic properties; this was also demonstrated in our current work, in Figure 4. Accordingly a suitable procedure for developing models can be summarized as comprising a detailed examination of the parameter space in the (λ_a, λ_r) plane, using an objective function tailored to include properties that are of the greatest importance for the purpose at hand, while utilising knowledge of the physical nature of the molecule and the conformal behaviour of Mie fluids of the same α to discriminate between near-degenerate models.

Using the strategy we have prescribed, we have developed models for a wide range of fluids, based not only on the generic Mie (λ_r - λ_a) potential (in which both exponents are free to vary), but also on the Mie (λ_r -6) form (for which the attractive exponent, λ_a , is constrained to the London-dispersion value of six), and the Mie (12-6) form, commonly known as the Lennard-Jones potential; the experimental data used in the parameter-estimation procedure

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comprised saturation properties including the vapour-pressure and saturated-liquid density, chosen since in our current work, the suitability of the models for use in mixture phaseequilibria problems is of key importance. (For more-general work, one might prefer models in which some single-phase properties are targetted, such as the liquid density or heat capacity.) In conjunction with the SAFT-VR Mie EOS⁴⁶, the resulting models provide an excellent description of a variety of thermophysical properties of the chosen fluids, including single-phase and second-derivative properties not included in the model development.

When comparing models based on the different potentials, generic Mie $(\lambda_r - \lambda_a)$ models are seen to provide the best performance though the difference in quality between these and the Mie (λ_r -6) models is not very significant – which is consistent with our earlier observation of conformality of Mie fluids (which implies that one can always find a Mie (λ_r -6) model with the same integrated energy parameter α as the generic Mie form, that will provide similar performance). Of the three classes of model, those based on the LJ (12-6) potential performed least well on average. Some fluids, exemplified by the Mie models of SF_6 or dimethylpropane, are not well represented with LJ models; SF_6 , in particular, requires an very repulsive potential. On the other hand, in some particular cases, such as the noble gases argon, krypton and xenon, the performance of the LJ models was just as good as that of the generic Mie potential. (Given the ubiquitous use of the LJ potential in both theory and simulation, it is reassuring to see how well LJ models can work in comparison with the moreversatile Mie models but, at the same time, it is very useful to be able to assess when such a model may not be appropriate.) Based on these comparisons and the foregoing discussion, for non-associating molecules of the type considered here we favour models based on the Mie (λ_r -6) form, since the potential offers an excellent compromise between the number of adjustable parameters, a respect of the physical nature of the molecules, and overall performance of the model.

The adequacy of our models in accurately capturing the saturation properties is seen to be outstanding, in the context of modelling using a theoretically-based equation of state (as opposed to a detailed mathematical correlation, such as the Span and Wagner⁸⁹ EOS for carbon dioxide). There is discernibly a significant improvement compared with the performance available using the previous formulation of SAFT-VR²² with available models. Especially noteworthy is the greatly reduced "overshoot" of the critical point; previously, such good performance simultaneously for both the sub-critical and near-critical regions

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using an analytical equation of state had been possible only with the incorporation of a cross-over treatment, such as those of References⁹⁰⁻⁹³.

For most problems of practical interest, one seeks to model not only pure-component fluids, but mixtures – often containing many components. The excellence of the description of pure-component saturation properties provides confidence that the fluid-phase equilibria of mixtures, based on these pure-component models, will in turn be well described. For example, one cannot obtain a good representation of the isothermal pressure-composition or isobaric temperature-composition phase diagrams of binary mixtures if one cannot first describe well both pure-component limits, *i.e.*, the vapour-pressure curves. On this basis alone, one can anticipate that the modelling of the fluid-phase equilibria of mixtures with the SAFT-VR EOS will be of equivalent quality to that of the pure components. However, by setting out a clearly prescribed strategy for pure-component models from which mixture models may later be developed are themselves developed in a consistent manner, thereby providing extra confidence that one will obtain a coherent and high-quality description of the properties of the fluid mixture.

Although the models developed in our current work are homonuclear models based on molecular segments described with the Mie potential, it is important to note that they may be used seamlessly in the SAFT- γ Mie group-contribution approach^{37,38}, in which generic heteronuclear molecular models formed from different types of fused Mie segments are considered. For example, the CO₂ model developed here can be used in a group-contribution methodology in modelling mixtures with hydrocarbons. Moreover, the generality of our models extends beyond their use in equation-of-state modelling. The faithful representation of the thermodynamic properties of Mie fluids with the SAFT-VR Mie⁴⁶ allows the same models to be employed in coarse-grained molecular simulations; thereby interfacial and transport properties, or other properties that are not directly accessible using the equation of state, may be determined. Successful examples of coarse-grained simulations based on SAFT-VR Mie and SAFT- γ Mie models are described in, for example, References^{71,94-100}.

In the introduction to our paper, we noted the trend in modern engineering practice towards a demand for equations of state that provide a simultaneous description of a wider variety of thermophysical properties, *i.e.*, an all-encompassing representation of the thermodynamics of a fluid, and we return to this point in closing. Although our models were

developed using only saturation properties, their performance in respect of the description they allow for other thermodynamic properties is nevertheless excellent. This has been exemplified in our current work for the single-phase compressed liquid density, enthalpy of vaporization, Joule-Thomson coefficient, speed of sound, and isobaric heat capacity, and, in a recent publication⁷², for a representation of the saturation properties in the (T, S)and (p, H) projections of the vapour-liquid equilibria. The capacity of the new SAFT-VR Mie EOS to simultaneously provide such high-quality descriptions of numerous thermodynamic properties in a simultaneous manner represents a tremendous advance in generalized equation-of-state modelling of fluids.

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Figure Captions

FIGURE 1

Behaviour of an objective function (Equation (28)) comprising the saturation properties of methane including the vapour pressure and saturated-liquid density (with equal weights, $\omega = 1$) obtained with the SAFT-VR Mie EOS⁴⁶ for planes of pairs of intermolecular parameters: (a) the (ε, λ_r) plane; (b) the (λ_a, λ_r) plane; (c) the (m, λ_a) plane; (d) the (m, λ_r) plane; (e) the (σ, λ_a) plane; and (f) the (σ, λ_r) plane. Contours are projected onto each plane; blue areas correspond to lower and red to higher values of the objective function (the quantification of the colour scale, common to each frame, is indicated in (b)). Two distinct valleys are present no matter the plane considered, with minima corresponding to the same models.

FIGURE 2

Behaviour of an objective function (Equation (28)) comprising the saturation and single phase properties of methane including the vapour pressure, saturated-liquid density, singlephase liquid density (each with weight $\omega = 1$), and speed of sound (with weight $\omega = 0.25$) obtained with the SAFT-VR Mie EOS⁴⁶ for the plane comprising the pair of Mie exponents $(\lambda_{\rm a}, \lambda_{\rm r})$; in (a) m is estimated freely, while in (b) m = 1 is prescribed. Blue areas correspond to lower and red to higher values of the objective function; note that the colour scales are chosen to provide the best contrast in each figure, whereby that in (b) differs from that in (a).

FIGURE 3

Molecular volume of the SAFT-VR Mie model for methane as characterized with the parameter $\mathcal{V} = m\sigma^3$ (in units of Å³) corresponding to optimizations in the plane comprising the pair of Mie exponents (λ_a, λ_r). Blue areas correspond to lower and red areas to higher volumes. The parameter estimations are carried out using an objective function (see Equation (28)) comprising the vapour pressure and saturated-liquid density, each with equal weight ($\omega = 1$). (The objective-function landscape corresponding to this plane is given in Figure 1(b).)

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FIGURE 4

Behaviour of the % AAD (Equation (29)) of the experimental isobaric heat capacity c_p of methane from the corresponding values calculated with the SAFT-VR Mie EOS⁴⁶ (blue areas correspond to lower and red to higher AAD values). The dashed white curves correspond to contours of the integrated energetic parameter α . The close coincidence in the shapes of the two sets of contours demonstrates that models yielding similar heat capacities are characterized by similar values of α . (In the parameter estimation, the objective function (Equation (28)) comprised the vapour pressure and saturated-liquid density with equal weights ($\omega = 1.0$).)

FIGURE 5

Performance of the different SAFT-VR Mie models based on the Lennard-Jones (12-6), Mie (λ_r -6), and generic Mie (λ_r - λ_a) potentials in relation to saturation properties of carbon dioxide: (a) vapour pressure; (b) saturation vapour and liquid densities; and (c) enthalpy of vaporization. The symbols represent the correlated experimental data from NIST⁶⁰, and the curves represent the description with the SAFT-VR Mie EOS⁴⁶.

FIGURE 6

Performance of the SAFT-VR Mie EOS⁴⁶ in relation to saturation properties of the *n*-alkanes: (a) vapour pressure; (b) saturation vapour and liquid densities; and (c) enthalpy of vaporization. The symbols represent correlated experimental data from NIST⁶⁰, and the curves represent the description with the Mie (λ_r -6) models presented in Table III.

FIGURE 7

Performance of the SAFT-VR Mie EOS⁴⁶ in relation to saturation properties of various compounds: (a) vapour pressure, (b) saturation vapour and liquid densities, and (c) enthalpy of vaporization. The symbols represent correlated experimental data from NIST⁶⁰, and the curves represent the description with the Mie (λ_r -6) models presented in Table III.

FIGURE 8

Prediction of single-phase isobaric properties of carbon dioxide with the SAFT-VR Mie EOS⁴⁶: (a) compressed-liquid density, (b) Joule-Thomson coefficient, (c) isobaric heat ca-

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pacity, and (d) speed of sound, at p = 10, 20, 30, 40 and 50 MPa. The symbols represent correlated experimental data from NIST⁶⁰, and the curves represent the description with the Mie (λ_r -6) model of carbon dioxide presented in Table III).

FIGURE 9

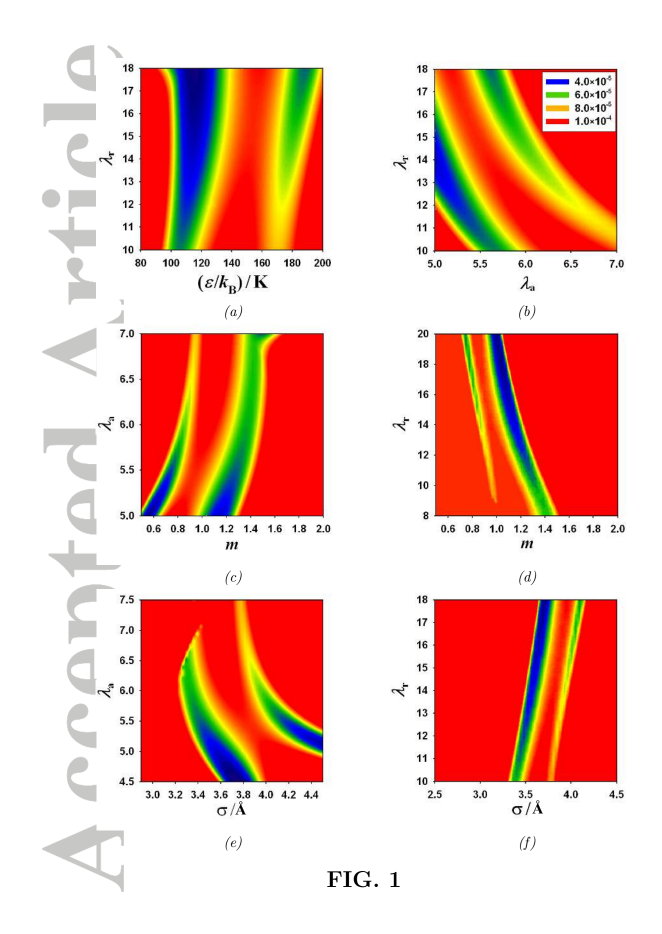
Prediction of single-phase isobaric properties of xenon with the SAFT-VR Mie EOS⁴⁶: (a) compressed-liquid density, (b) Joule-Thomson coefficient, (c) isobaric heat capacity, and (d) speed of sound, at p = 5, 10, 15, 20, and 30 MPa. The symbols represent correlated experimental data from NIST⁶⁰, and the curves represent the description with the Mie (λ_r -6) model of xenon presented in Table III).

FIGURE 10

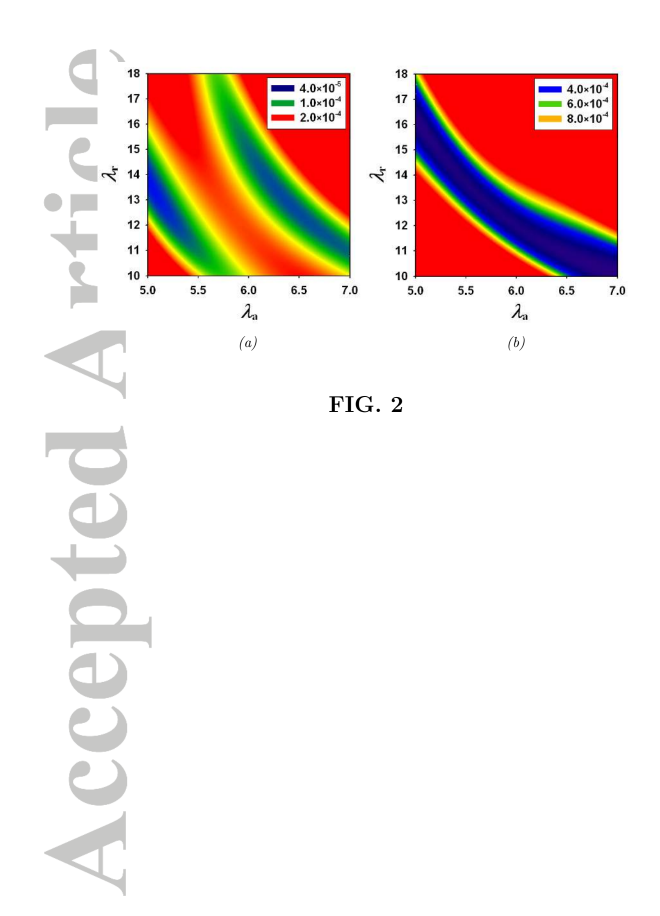
Prediction of single-phase isobaric properties of isopentane with the SAFT-VR Mie EOS⁴⁶: (a) compressed-liquid density, (b) Joule-Thomson coefficient, (c) isobaric heat capacity, and (d) speed of sound, at p = 2, 3, 4, 5, and 10 MPa. The symbols represent correlated experimental data from NIST⁶⁰, and the curves represent the description with the Mie (λ_r -6) model of isopentane presented in Table III).

FIGURE 11

Trends in (a) the molecular volume $\mathcal{V} = m\sigma^3$, and (b) the integrated van der Waals-like attractive energy $E_{\rm int} = m^2 \sigma^3 \epsilon \alpha$ for the models of the *n*-alkanes and *n*-alkyl benzenes; blue corresponds to Lennard-Jones (12-6) models, red to Mie ($\lambda_{\rm r}$ -6) models, and green to models based on the generic Mie ($\lambda_{\rm r}$ - $\lambda_{\rm a}$) potential for the *n*-alkane family; purple corresponds to the Mie ($\lambda_{\rm r}$ -6) models for the *n*-alkyl benzene family. The values of R^2 represent the square of the Pearson product-moment correlation coefficient for the fits represented by the dashed curves (linear for the molecular volume, second degree polynomial for the integrated energy).



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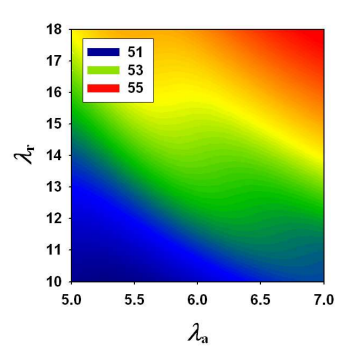


FIG. 3

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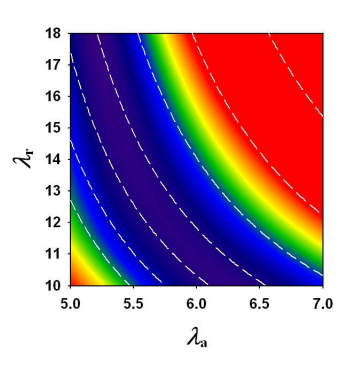
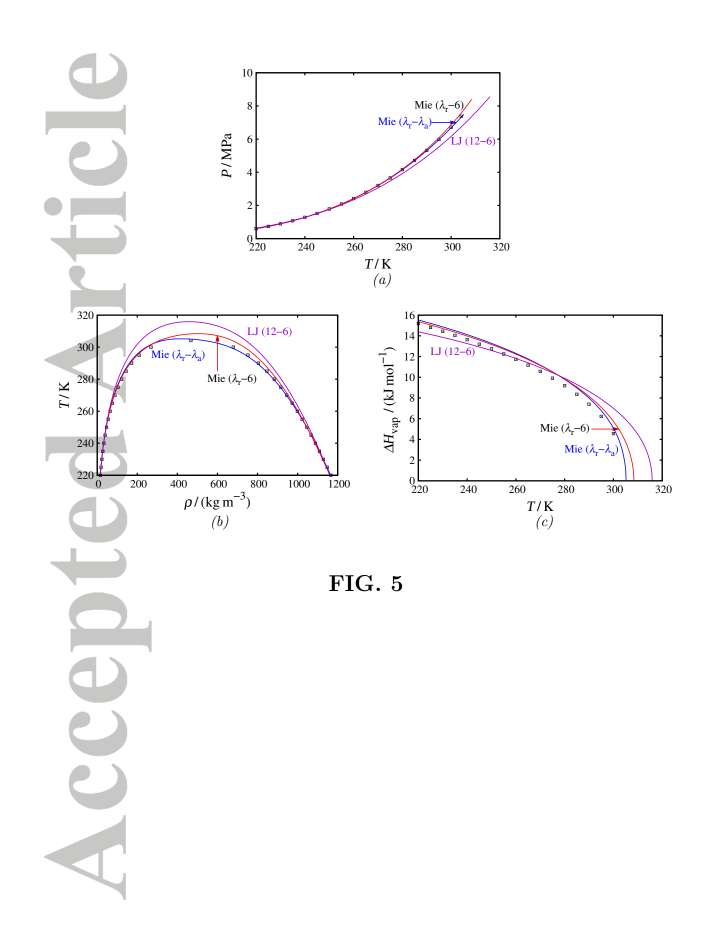
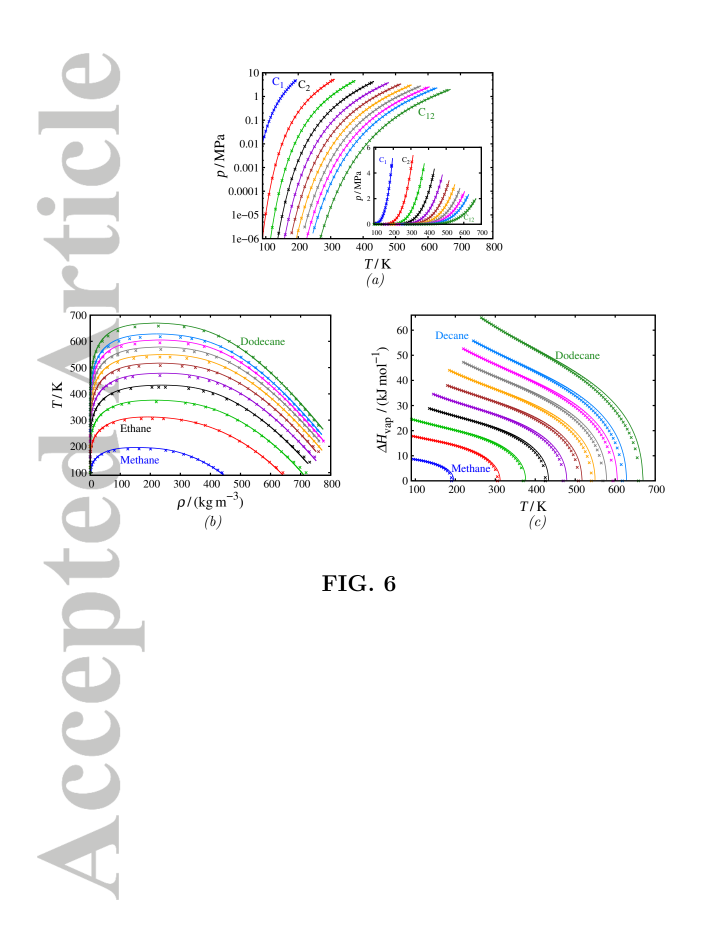


FIG. 4

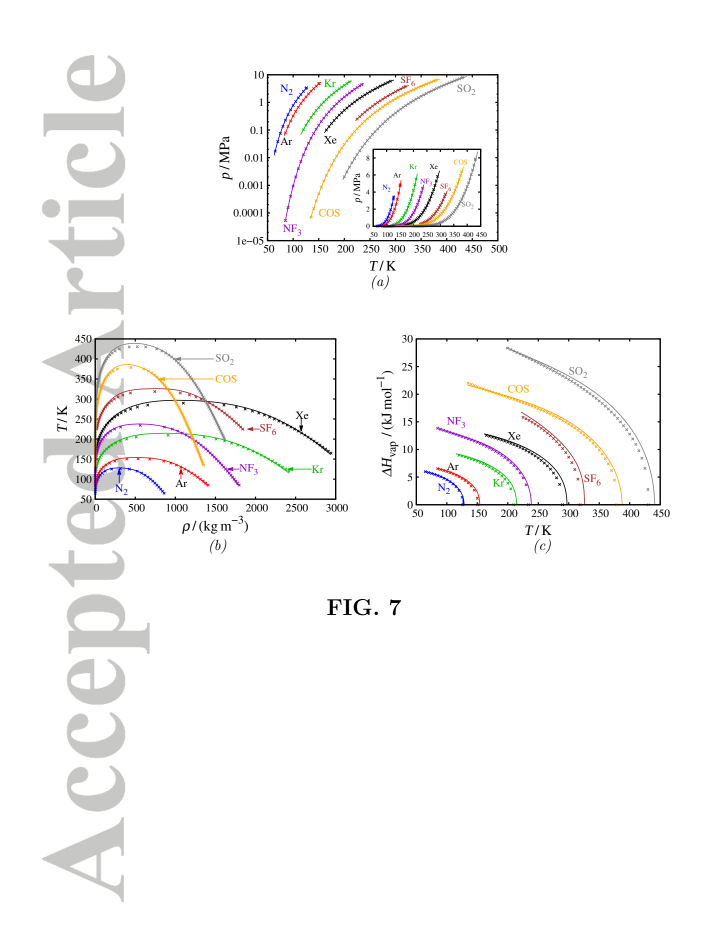
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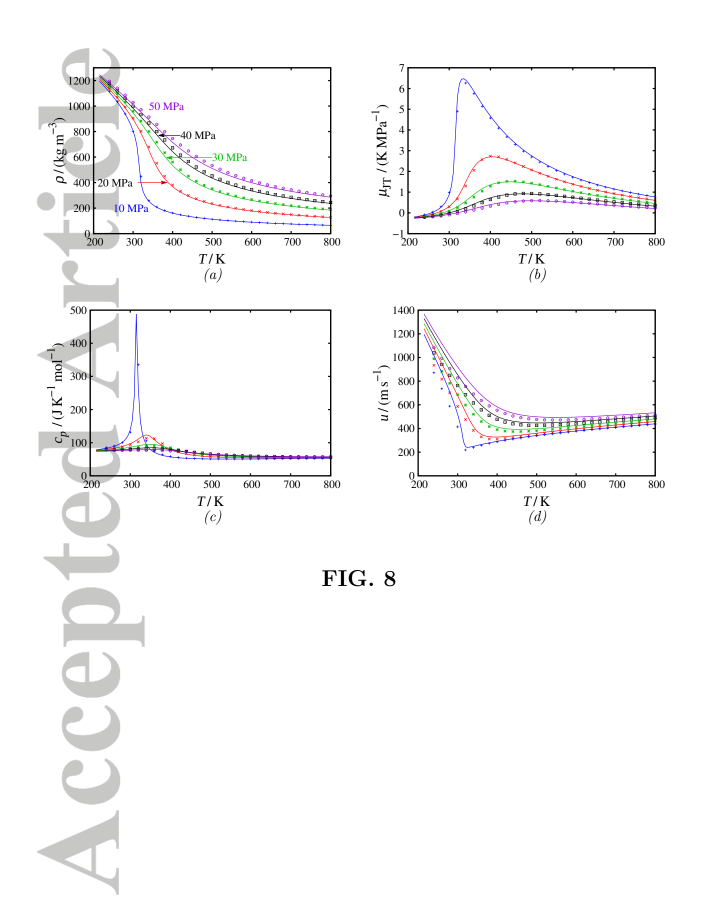


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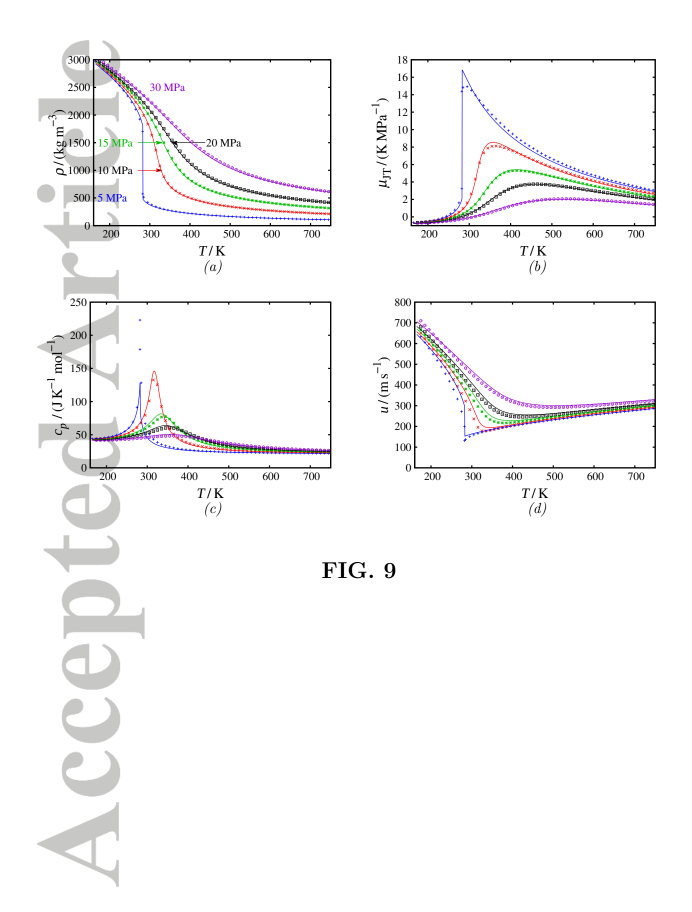


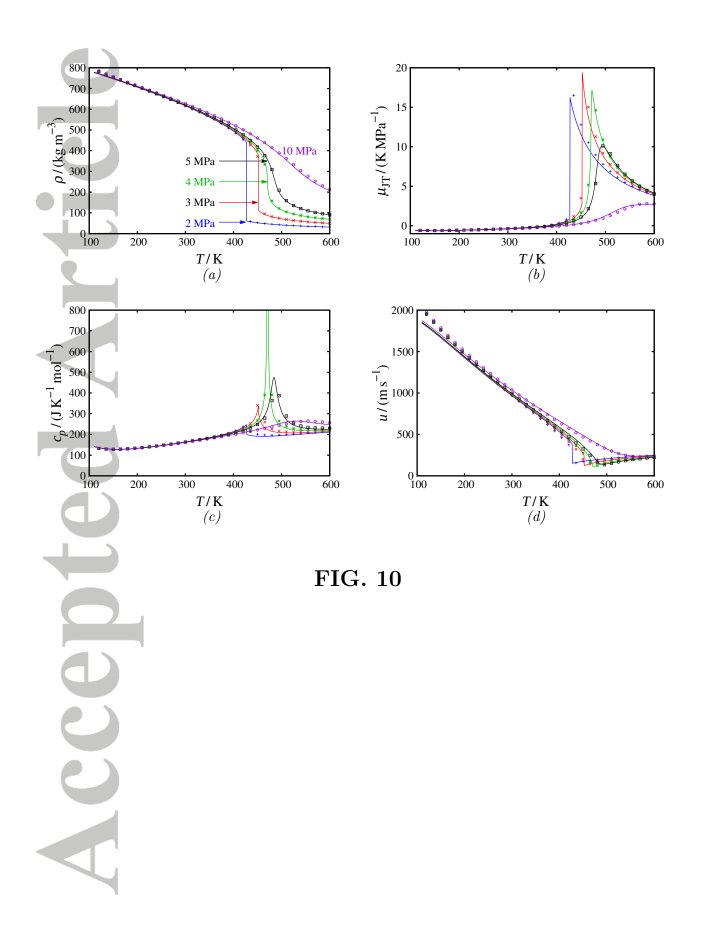
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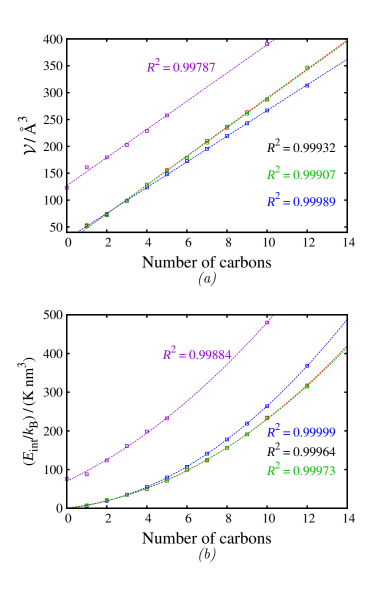


FIG. 11

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Cubatanao		Mole	Molecular Parar	ameters					% AAD			
antrance	m	$\sigma / \text{\AA}$	$\lambda_{ m r}$	$\lambda_{ m a}$	$(\varepsilon/k_{\rm B}) \ / \ { m K}$	$d_{ m liq}$	n	$d_{ m sat}$	p_{sat}	c_p	$T_{\rm c}$	$\Delta H_{\rm vap}$
Methane	1.0000^{\dagger}	3.7366	12.319	6.0^{\dagger}	151.45	0.54	1.81	0.75	0.40	1.23	2.69	2.69
Methane *	1.0000^{\dagger}	3.7412	12.650	6.0^{\dagger}	153.36	0.74	1.97	0.75	0.63	2.03	2.41	2.87
Ethane	1.7230	3.4763	10.121	6.0^{\dagger}	164.27	0.53	5.75	0.40	0.44	2.59	2.27	1.49
Ethane *	1.4373	3.7257	12.400	6.0^{\dagger}	206.12	0.37	3.87	0.55	0.53	1.69	0.49	1.73
Propane	1.8068	3.7943	12.106	6.0^{\dagger}	221.96	0.59	9.39	0.67	0.77	2.24	1.68	1.44
$\operatorname{Propane}^*$	1.6845	3.9056	13.006	6.0^{\dagger}	239.89	0.51	4.76	0.93	0.44	2.40	1.44	1.41
Butane	1.6791	4.2476	15.453	6.0^{\dagger}	306.52	0.34	2.45	0.58	0.09	0.93	1.82	1.46
Butane *	1.8514	4.0887	13.650	6.0^{\dagger}	273.64	0.31	1.35	0.54	0.39	1.56	1.80	1.55
Pentane	1.8594	4.3759	16.438	6.0^{\dagger}	336.74	0.19	2.05	0.34	0.16	0.90	2.36	1.39
Pentane *	1.9606	4.2928	15.847	6.0^{\dagger}	321.94	0.32	1.65	0.52	0.60	0.68	2.85	1.67
Hexane	2.2549	4.2968	15.069	6.0^{\dagger}	321.81	0.29	1.83	0.29	0.16	0.87	2.47	1.68
${ m Hexane}^*$	2.1097	4.4230	17.203	6.0^{\dagger}	354.38	0.46	1.43	0.25	1.19	0.32	3.26	2.23
Heptane	2.2413	4.5427	18.252	6.0^{\dagger}	381.42	0.37	0.92	0.19	0.27	0.78	3.26	1.65
${ m Heptane}^*$	2.3949	4.4282	17.092	6.0^{\dagger}	358.51	0.39	0.67	0.45	0.91	0.91	3.52	1.79
Octane	2.4777	4.5708	18.654	6.0^{\dagger}	391.87	0.61	1.79	0.36	0.29	0.86	3.62	1.59
Octane *	2.6253	4.4696	17.378	6.0^{\dagger}	369.18	0.54	0.75	0.52	0.96	1.04	3.60	1.58
Nonane	2.6665	4.6236	19.116	6.0^{\dagger}	404.83	0.47	0.60	0.41	0.34	0.72	3.53	1.48
Nonane *	2.8099	4.5334	18.324	6.0^{\dagger}	387.55	0.43	0.45	0.60	0.82	0.63	3.77	1.60
Decane	3.0058	4.5727	18.403	6.0^{\dagger}	396.17	0.65	1.55	0.46	0.41	0.70	3.29	1.74
Decane *	2.9976	4.5890	18.885	6.0^{\dagger}	400.79	0.79	1.14	0.59	0.96	0.46	3.67	1.86
Dodecane	3.2483	4.7430	20.872	6.0^{\dagger}	438.2	0.68	1.22	0.43	0.56	0.95	3.52	1.78
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that the set that the the the the the the the the the th		$_{ m c}$ $\Delta H_{ m vap}$	37 230
TABLE II: Intermolecular potential models developed for various hydrocarbons using Mie (λ_r -6) potentials; [†] denotes that the highlighted parameter is fixed at the indicated value during the parameter estimation. The percentage absolute average deviation % <i>AAD</i> of the various properties calculated with the SAFT-VR Mie EOS ⁴⁶ from the correlated experimental data of NIST ⁶⁰ are also indicated: single-phase-liquid density d_{iq} ; single-phase-liquid speed of sound u ; saturated-liquid density d_{sat} ; vapour pressure p_{sat} ; single-phase-liquid heat capacity c_p ; critical temperature T_c ; enthalpy of vaporization ΔH_{vap} . Also included for comparison are models developed in Reference ⁴⁶ ; these models were developed using p_{sat} , d_{sat} , T_c , and u in the parameter estimation. *	% AAD	$c_p T_c$	059 347 099 035 939 107 930
Mie $(\lambda_{\rm r}$ -6) pol The percentage orrelated experimated curve urated-liquid c tion $\Delta H_{\rm vap}$. Al	7 %	$d_{ m sat}$ $p_{ m sat}$	0 22 0 3
carbons using r estimation. ¹ S ⁴⁶ from the c of sound u ; sat by of vaporizat ng p_{sat} , T_{sat} , $T_{\text{Reference}^{46}}$.		$d_{ m liq}$ u	0.52 3.47
developed for various hydrocarbons usion of value during the parameter estimation with the SAFT-VR Mie EOS ⁴⁶ from the single-phase-liquid speed of sound w ; tical temperature T_c ; enthalpy of vaporitical temperature developed using p_{sat} , d_{sat}^{46} , denotes models taken from Reference ⁴⁶ .		$(\varepsilon/k_{ m B}) \ / \ { m K}$	907 53
s developed for ed value during with the SAF ⁱ u; single-phase ttical temperat se models were denotes mode	Molecular Parameters	$\lambda_{ m r}$ $\lambda_{ m a}$	504 6 0 [†]
otential models at the indicate ties calculated quid density $d_{\rm li}$ capacity c_p ; cri eference ⁴⁶ ; thes	Molecula	$\sigma / m \AA$	<u>9 9785 3 7806 11 504 6 0†</u>
termolecular potenti ameter is fixed at th various properties c single-phase-liquid c se-liquid heat capac leveloped in Referen		m	9.9785
TABLE II: Intermolecular potential models developed for various hydrocarbons using Mie (λ_r -6) potentials; [†] denotes that the highlighted parameter is fixed at the indicated value during the parameter estimation. The percentage absolute average deviation % <i>AAD</i> of the various properties calculated with the SAFT-VR Mie EOS ⁴⁶ from the correlated experimental data of NIST ⁶⁰ are also indicated: single-phase-liquid density $d_{\rm ind}$; single-phase-liquid speed of sound u ; saturated-liquid density $d_{\rm sat}$; vapour pressure $p_{\rm sat}$; single-phase-liquid heat capacity c_p ; critical temperature T_c ; enthalpy of vaporization $\Delta H_{\rm vap}$. Also included for comparison are models developed in Reference ⁴⁶ ; these models were developed using $p_{\rm sat}$, $d_{\rm sat}$, T_c , and u in the parameter estimation. *	Cubatanao	AULANDA	Banzana

Substance		Molecula		r Parameters					% AAL			
antrajerance	m	$\sigma / \text{\AA}$	$\lambda_{ m r}$	$\lambda_{ m a}$	$(\varepsilon/k_{\rm B}) \ / \ {\rm K}$	$d_{ m liq}$	n	$d_{ m sat}$	$p_{ m sat}$	c_p	$T_{\rm c}$	$\Delta H_{ m vap}$
Benzene	2.2785	3.7806	11.594	6.0^{\dagger}	297.53	0.52	3.47	0.22	0.35	2.32	1.97	2.30
${ m Benzene}^*$	1.9163	4.0549	14.798	6.0^{\dagger}	372.59	0.44	2.33	0.34	0.89	1.95	2.80	2.46
Cyclohexane	2.1641	4.1071	12.412	6.0^{\dagger}	313.04	0.56	4.79	0.17	0.30	2.31	1.53	3.28
Cyclopropane	1.8769	3.4830	10.507	6.0^{\dagger}	213.11	0.63	4.97	0.02	0.07	2.38	2.56	4.54
Dimethylpropane	1.0000^{\dagger}	5.6491	31.792	6.0^{\dagger}	504.25	0.93	5.81	0.14	0.65	2.56	2.97	3.27
Ethylene	1.7972	3.2991	9.6463	6.0^{\dagger}	142.64	0.54	3.56	0.37	0.52	4.18	2.07	1.66
Isobutane	1.7186	4.2177	14.612	6.0^{\dagger}	281.12	0.43	8.43	0.62	0.36	1.70	2.11	1.48
Isohexane	2.2759	4.2839	14.756	6.0^{\dagger}	307.74	0.53	4.80	0.40	1.48	4.74	3.18	1.53
Isopentane	1.7464	4.4714	16.688	6.0^{\dagger}	339.95	0.28	2.27	0.37	0.27	0.85	2.40	1.17
$\operatorname{Propylene}$	2.0060	3.5392	10.643	6.0^{\dagger}	190.13	0.62	5.01	0.55	0.78	2.57	1.57	1.50
$\operatorname{Propyne}$	2.3038	3.2076	10.667	6.0^{\dagger}	200.11	0.31	6.45	0.16	0.53	7.00	2.89	4.54
Toluene	1.7112	4.5487	19.125	6.0^{\dagger}	474.13	0.35	3.44	0.46	0.20	1.15	1.74	1.82
Toluene *	1.9977	4.2777	16.334	6.0^{\dagger}	409.73	0.28	2.25	0.46	1.46	1.91	2.96	2.15
Ethyl-benzene	2.3501	4.2436	15.241	6.0^{\dagger}	386.09	ı	ı	2.85	0.60	ı	ı	ı
Propyl-benzene	2.7357	4.1999	12.853	6.0^{\dagger}	342.41	ı	ı	2.66	0.17	ı	ı	ı
Butyl-benzene	2.9762	4.2522	13.237	6.0^{\dagger}	350.48	ı	ı	3.50	0.19	ı	ı	ı
Pentyl-benzene	3.0488	4.3881	15.594	6.0^{\dagger}	394.46	ı	ı	1.47	0.50	ı	ı	ı
Decyl-benzene	4.0808	4.5745	17.746	6.0^{\dagger}	431.37	I	I	3.29	0.42	I	I	I

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hat the leviation ST ⁶⁰ are pressure nparison neter		$\Delta H_{ m vap}$	3.09	1.27	3.08	1.84	1.25
enotes th average (ata of NI ; vapour he paran		$T_{ m c}$	2.19	1.54	2.69	2.30	1.58
tials; † d absolute nental dansity $d_{\rm sat}$ includeo nd u in t	-	c_p	1.77	2.64	3.45	3.46	2.04
6) poten centage a d experir iquid den vap. Alsc vap. diq, a	% AAD	$p_{ m sat}$	0.21	0.08	0.12	0.74	0.02
Mie ($\lambda_{\rm r}$ - The pervorted pervorted to ΔH turrated-1 tion ΔH tion $\Delta H_{\rm va}$ to ce^{46} .		$d_{ m sat}$	0.66	0.15	0.3	0.38	0.25
ds using imation. com the com the complexity $u_{\rm sat}$, $d_{\rm sat}$, $d_{$		n	2.13	2.39	16.86	3.96	5.73
s compour ameter est ie EOS^{46} fi peed of sou nthalpy of bed using p taken fron		$d_{ m liq}$	0.49	0.32	0.85	0.59	0.36
TABLE III: Intermolecular potential models developed for various compounds using Mie (λ_r -6) potentials; [†] denotes that the highlighted parameter is fixed at the indicated value during the parameter estimation. The percentage absolute average deviation % AAD of the various properties calculated with the SAFT-VR Mie EOS ⁴⁶ from the correlated experimental data of NIST ⁶⁰ are also indicated: single-phase-liquid density $d_{\rm inq}$; single-phase-liquid speed of sound u ; saturated-liquid density $d_{\rm sat}$; vapour pressure $p_{\rm sat}$; single-phase-liquid heat capacity c_p ; critical temperature T_c ; enthalpy of vaporization $\Delta H_{\rm vap}$. Also included for comparison are models developed in Reference ⁴⁶ ; these models were developed using $p_{\rm sat}$, T_c , $\Delta H_{\rm vap}$, $d_{\rm inq}$, and u in the parameter estimation. * denotes taken from Reference ⁴⁶ .		$(\varepsilon/k_{ m B}) \ / \ { m K}$	117.84	72.11	235.73	186.38	80.803
ls develo d value o with the i; single- tical tem se models ion. * de	meters	$\lambda_{ m a}$	6.0^{\dagger}	6.0^{\dagger}	6.0^{\dagger}	6.0^{\dagger}	6.0^{\dagger}
ntial mode ne indicate salculated density $d_{\rm li}$ city c_p ; crit ince ⁴⁶ ; thes estimat	Molecular Parameters	$\lambda_{ m r}$	12.085	9.7420	18.067	9.5435	9.9255
lecular poten is fixed at th s properties c phase-liquid c pid heat capac ped in Referen	Mole	$\sigma \ / \ { m \AA}$	3.4038	3.0928	3.0465	3.2978	2.8118
I: Intermole barameter is he various J d: single-ph bhase-liquid els develope		m	1.0000^{\dagger}	1.5556	1.6936	1.8746	1.5094
TABLE IIhighlighted I $\%$ AAD of talso indicate p_{sat} ; single-Iare mode	Substance	DUIDUCUUC	Ar	CO	CO_2	COS	${\mathbb H}_2$

Substance		Mole	Molecular Para	meters					% AAD			
antrance	m	$\sigma / \text{Å}$	$\lambda_{ m r}$	$\lambda_{\rm a}$	$(\varepsilon/k_{ m B}) \ / \ { m K}$	$d_{ m liq}$	n	$d_{ m sat}$	p_{sat}	c_p	$T_{\rm c}$	$\Delta H_{\rm vap}$
$\overline{\mathrm{Ar}}$	1.0000^{\dagger}	3.4038	12.085	6.0^{\dagger}	117.84	0.49	2.13	0.66	0.21	1.77	2.19	3.09
CO	1.5556	3.0928	9.7420	6.0^{\dagger}	72.11	0.32	2.39	0.15	0.08	2.64	1.54	1.27
CO_2	1.6936	3.0465	18.067	6.0^{\dagger}	235.73	0.85	16.86	0.3	0.12	3.45	2.69	3.08
COS	1.8746	3.2978	9.5435	6.0^{\dagger}	186.38	0.59	3.96	0.38	0.74	3.46	2.30	1.84
F_2	1.5094	2.8118	9.9255	6.0^{\dagger}	80.803	0.36	5.73	0.25	0.02	2.04	1.58	1.25
$\mathrm{F}_{2}^{}$ *	1.3211	2.9554	11.606	6.0^{\dagger}	96.268	0.31	4.52	0.50	0.44	1.47	0.61	1.51
Kr	1.0000^{\dagger}	3.6359	11.985	6.0^{\dagger}	163.27	0.57	2.33	0.69	0.19	0.89	2.35	3.32
N_2	1.4214	3.1760	9.8749	6.0^{\dagger}	72.438	0.46	2.71	0.15	0.07	2.31	0.06	1.43
$ m N_2O$	2.0526	2.8440	10.930	6.0^{\dagger}	163.69	0.51	7.20	0.09	0.04	2.03	2.13	2.99
Ne	1.0^{\dagger}	2.8019	9.6977	6.0^{\dagger}	29.875	0.45	6.57	0.06	0.43	4.56	1.69	2.02
NF_3	1.8196	3.2069	10.764	6.0^{\dagger}	128.73	0.48	7.20	0.31	0.34	2.81	1.67	1.81
O_2	1.4283	2.9671	8.9218	6.0^{\dagger}	81.476	0.55	5.08	0.39	0.24	2.76	1.89	1.04
${ m SF}_6$	1.0000^{\dagger}	4.8768	36.696	6.0^{\dagger}	381.99	0.96	5.78	0.07	0.41	6.25	2.29	4.29
SO_2	2.4599	2.8511	11.865	6.0^{\dagger}	225.73	0.69	15.17	0.11	0.16	3.10	1.96	2.88
Xe	1.0000^{\dagger}	3.9612	12.275	6.0^{\dagger}	229.47	0.63	2.86	0.74	0.19	1.16	2.55	3.63
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TABLE IV: Ranges of temperature T and pressure p of the experimental data incorporated in the parameter estimations during the development of models, and for the determined of the percentage absolute average deviation % AAD of the various properties calculated with the SAFT-VR Mie EOS⁴⁶ from the correlated experimental data for each fluid. Also provided are the numbers of data points used. The correlated experimental data are taken from the NIST database⁶⁰, except for the alkyl benzenes, for which the sources of data are indicated by the cited references.

		d equilibrium		npressed-liquid	-
substance	T range (K)	\sharp (data points)	T range (K)	P range (MPa)	$\sharp(\text{data points})$
Methane	95 - 185	21	100 - 190	10 - 50	30
Ethane	95 - 275	37	100 - 300	10 - 50	33
Propane	90 - 330	49	100 - 360	10 - 50	42
Butane	135 - 380	50	140 - 420	10 - 50	44
Pentane	145 - 420	56	150 - 450	10 - 50	31
Hexane	180 - 455	56	200 - 500	10 - 50	32
Heptane	185 - 485	61	200 - 530	10 - 50	35
Octane	220 - 510	59	230 - 560	10 - 50	35
Nonane	220 - 535	64	230 - 560	10-50	34
Decane	245 - 555	63	300 - 600	10-50	33
Dodecane	265 - 590	66	290 - 650	10 - 50	38
Cyclohexane	280 - 495	44	300 - 560	10 - 50	38
Cyclopropane	275 - 355	17	290 - 390	10 - 28	30
Dimethylpropane	260 - 390	27	270 - 430	10 - 50	44
Ethylene	105 - 250	30	110 - 280	10 - 50	53
Isobutane	115 - 365	51	125 - 405	10 - 35	45
Isohexane	125 - 445	65	130 - 490	10-50	38
Isopentane	115 - 410	60	120 - 460	10-50	53
Propylene	100 - 325	46	120 - 360	10 - 50	39
Propyne	275 - 360	18	280 - 400	10 - 30	36
Benzene	280 - 505	46	300 - 560	10-50	41
Toluene	180 - 530	71	190 - 590	10-50	62
$Ethyl-benzene^{64,65}$	183 - 518	24	-	-	-
$Propyl-benzene^{64,65}$	223 - 543	15	-	-	-
$Butyl-benzene^{64,65}$	223 - 583	13	-	-	-
$Pentyl-benzene^{64,65}$	223 - 593	10	_	-	-
Decyl-benzene ^{62,63,65,66}	318 - 633	12	-	-	-
Ar	85 - 135	11	90 - 150	10 - 50	20
CO	70 - 115	10	75 - 130	10 - 50	35
CO_2	220 - 270	22	220 - 300	10 - 50	50
COS	135 - 340	42	155 - 375	10 - 50	36
F_2	55 - 125	15	60 - 140	10 - 20	27
Kr	120 - 185	14	120 - 200	10 - 50	24
N_2	65 - 110	10	75 - 125	10 - 100	23
N_2O	185 - 275	19	195 - 305	10-50	34
Ne	24 - 40	16	29 - 45	10-50	20
NF_3	85 - 210	26	95 - 225	10-50	41
O_2	55 - 135	17	60 - 150	10-50	30
SF_6	225 - 285	13	235 - 315	10 - 55	22
SO_2	200 - 385	3\$3	220 - 420	10 - 35	33
Xe	165 - 260	20	175 - 285	10-50	35

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Appendix A: Models Based on Lennard-Jones (12-6) and the Generic Mie $(\lambda_r - \lambda_a)$ Potentials

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TABLE V: Intermolecular potential models developed for the n-alkanes using Lennard-Jones (12-6) potentials; † denotes that the highlighted parameter is fixed at the indicated value during the parameter estimation. The percentage absolute average deviation % AAD of the various properties calculated with the SAFT-VR Mie EOS⁴⁶ from the correlated experimental data of NIST⁶⁰ are also indicated: single-phase-liquid density $d_{\rm liq}$; single-phase-liquid speed of sound u; saturated-liquid density $d_{\rm sat}$; vapour pressure p_{sat} ; single-phase-liquid heat capacity c_p ; critical temperature T_c ; enthalpy of vaporization ΔH_{vap} .

Chateneo		Molec	Molecular Para	umeters					% AAL	~		
annaisana	m	$\sigma / \text{Å}$	$\lambda_{ m r}$	$\lambda_{\rm a}$	$(\varepsilon/k_{ m B}) \ / \ { m K}$	$d_{ m liq}$	n	$d_{\rm sat}$	$p_{\rm sat}$	c_p	$T_{\rm c}$	$\Delta H_{ m vap}$
Methane	1.0000^{\dagger}	3.7340	12.0^{\dagger}	6.0^{\dagger}	149.49	0.51	1.79	0.80	0.49	0.71	2.94	2.44
Ethane	1.4702	3.6905	12.0^{\dagger}	6.0^{\dagger}	199.90	0.67	4.09	0.90	0.24	1.82	0.41	1.64
Propane	1.8236	3.7810	12.0^{\dagger}	6.0^{\dagger}	219.70	0.58	9.33	0.64	0.82	2.25	2.21	1.44
Butane	2.0904	3.8965	12.0^{\dagger}	6.0^{\dagger}	238.91	0.45	4.97	0.27	0.69	2.19	1.85	1.66
Pentane	2.4361	3.9312	12.0^{\dagger}	6.0^{\dagger}	248.10	0.64	6.10	0.45	1.20	2.01	2.09	1.68
Hexane	2.7048	3.9946	12.0^{\dagger}	6.0^{\dagger}	258.62	0.75	4.72	0.50	0.43	1.35	0.68	1.71
Heptane	3.1197	3.9723	12.0^{\dagger}	6.0^{\dagger}	260.32	1.18	6.70	1.03	1.10	2.00	0.60	1.99
Octane	3.4364	3.9966	12.0^{\dagger}	6.0^{\dagger}	265.57	1.38	5.82	1.25	0.69	1.76	0.18	1.83
Nonane	3.7694	4.0093	12.0^{\dagger}	6.0^{\dagger}	269.03	1.48	8.05	1.46	0.94	1.51	0.22	1.73
Decane	4.0744	4.0309	12.0^{\dagger}	6.0^{\dagger}	272.85	1.59	7.81	1.50	0.83	0.87	3.42	1.82
Dodecane	4.7992	4.0266	12.0^{\dagger}	6.0^{\dagger}	275.65	1 77	х 7.	1 74	1 00	1 30	3,02	1 95

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TABLE VI: Intermolecular potential models developed for various hydrocarbons using Lennard-Jones (12-6) potentials; † denotes that the highlighted parameter is fixed at the indicated value during the parameter estimation. The percentage absolute average deviation % AAD of the various properties calculated with the SAFT-VR Mie EOS⁴⁶ from the correlated experimental data of $NIST^{60}$ are also indicated: single-phase-liquid density d_{liq} ; single-phase-liquid speed of sound u; saturated-liquid density d_{sat} ; vapour pressure p_{sat} ; single-phase-liquid heat capacity c_p ; critical temperature T_c ; enthalpy of vaporization ΔH_{vap} .

Chatamaa		Molecular	ular Parê	ameters					% AAL			
anustance	m	$\sigma / \text{Å}$	$\lambda_{\rm r}$	$\lambda_{\rm a}$	$(\varepsilon/k_{\rm B}) \ / \ { m K}$	$d_{\rm liq}$	n	$d_{\rm sat}$	$p_{\rm sat}$	c_p	$T_{\rm c}$	$\Delta H_{\rm vap}$
Benzene	2.2125	3.8253	12.0^{\dagger}	6.0^{\dagger}	308.59	0.41	3.05	0.18	0.41	2.34	1.95	2.35
Cyclohexane	2.2265	4.0607	12.0^{\dagger}	6.0^{\dagger}	302.32	0.67	5.33	0.25	0.26	2.30	1.69	3.24
Cyclopropane	1.6575	3.6592	12.0^{\dagger}	6.0^{\dagger}	247.79	0.75	7.82	0.40	0.14	1.84	0.63	4.72
Dimethylpropane	1.0000^{\dagger}	5.4536	12.0^{\dagger}	6.0^{\dagger}	363.81	3.11	6.37	2.47	11.7	9.49	9.97	8.49
Ethylene	1.4768	3.5526	12.0^{\dagger}	6.0^{\dagger}	183.63	0.44	2.87	0.75	0.68	3.55	0.89	1.97
Isobutane	2.0645	3.9267	12.0^{\dagger}	6.0^{\dagger}	229.24	0.50	11.57	0.36	1.14	1.84	1.57	1.55
Isohexane	2.7808	3.9638	12.0^{\dagger}	6.0^{\dagger}	246.75	0.99	8.68	1.07	3.66	5.11	2.24	1.42
Isopentane	2.4487	3.9236	12.0^{\dagger}	6.0^{\dagger}	239.03	0.92	7.86	1.02	4.43	3.23	0.35	1.37
Propylene	1.7920	3.6948	12.0^{\dagger}	6.0^{\dagger}	218.08	0.62	3.91	0.91	0.61	2.57	1.76	1.65
$\operatorname{Propyne}$	2.0611	3.3561	12.0^{\dagger}	6.0^{\dagger}	228.75	0.23	6.36	0.53	0.45	6.28	2.14	4.69
Toluene	2.5926	3.8539	12.0^{\dagger}	6.0^{\dagger}	304.36	0.74	5.42	0.54	2.53	2.93	1.37	2.48

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TABLE VII: Intermolecular potential models developed for various compounds using Lennard-Jones (12-6) potentials; † denotes that the highlighted parameter is fixed at the indicated value during the parameter estimation. The percentage absolute average deviation % AAD of the various properties calculated with the SAFT-VR Mie EOS⁴⁶ from the correlated experimental data of $NIST^{60}$ are also indicated: single-phase-liquid density d_{liq} ; single-phase-liquid speed of sound u; saturated-liquid density d_{sat} ; vapour pressure p_{sat} ; single-phase-liquid heat capacity c_p ; critical temperature T_c ; enthalpy of vaporization ΔH_{vap} .

$\begin{array}{c c} Ar & 1.0 \\ Ar & 1.0 \\ CO & 1.5 \\ CO & 1.6 \\ CO & 1.6 \\ F_{0} & 1.5 \end{array}$	$rac{m}{.0000^{\dagger}}$ L.2903	× /							TVV 0/			
	000† 2903	$\sigma \ \ A$	$\lambda_{ m r}$	$\lambda_{ m a}$	$(\varepsilon/k_{\rm B}) \ / \ { m K}$	$d_{ m liq}$	n	$d_{ m sat}$	$p_{ m sat}$	c_p	$T_{ m c}$	$\Delta H_{ m vap}$
	2903	3.4033	12.0^{\dagger}	6.0^{\dagger}	117.41	0.48	2.15	0.67	0.19	1.76	2.24	3.02
		3.3186	12.0^{\dagger}	6.0^{\dagger}	91.515	0.40	1.66	0.62	0.21	2.92	0.55	1.63
	1.6936	3.0189	12.0^{\dagger}	6.0^{\dagger}	190.60	0.99	8.51	0.92	2.67	13.94	1.00	2.71
	5262	3.5645	12.0^{\dagger}	6.0^{\dagger}	243.24	0.44	4.11	0.77	0.93	2.61	0.82	2.18
	1.2684	2.9998	12.0^{\dagger}	6.0^{\dagger}	100.44	0.40	4.50	0.62	0.11	1.46	0.26	1.51
Kr 1.0	1000†	3.6360	12.0^{\dagger}	6.0^{\dagger}	163.38	0.57	2.33	0.69	0.20	0.91	2.34	3.34
	1.1959	3.3879	12.0^{\dagger}	6.0^{\dagger}	90.216	0.30	1.18	0.55	0.10	2.13	0.21	1.81
N_2O 1.8	8872	2.9405	12.0^{\dagger}	6.0^{\dagger}	181.47	0.55	9.20	0.35	0.12	2.29	1.73	3.10
	000¢	2.8099	12.0^{\dagger}	6.0^{\dagger}	33.916	0.83	1.88	0.49	2.69	11.8	0.03	4.45
NF_3 1.6	3517	3.3283	12.0^{\dagger}	6.0^{\dagger}	145.16	0.37	6.02	0.54	0.32	2.40	1.05	1.92
	0817	3.2824	12.0^{\dagger}	6.0^{\dagger}	116.45	0.77	2.50	1.04	0.24	1.65	1.68	1.51
—	$.0000^{\dagger}$	4.7035	12.0^{\dagger}	6.0^{\dagger}	262.89	2.49	7.97	1.56	7.18	12.04	8.23	9.17
$SO_2 \qquad 2.4$	2.4368	2.8619	12.0^{\dagger}	6.0^{\dagger}	228.41	0.69	15.44	0.13	0.15	3.09	0.79	2.89
Xe 1.0	$.0000^{\dagger}$	3.9593	12.0^{\dagger}	6.0^{\dagger}	226.80	0.59	2.81	0.78	0.22	1.28	2.72	3.37

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highlighted parameter is fixed at the indicated value during the parameter estimation. The percentage absolute average deviation TABLE VIII: Intermolecular potential models developed for the n-alkanes using generic Mie (λ_r - λ_a) potentials; [†] denotes that the % AAD of the various properties calculated with the SAFT-VR Mie EOS⁴⁶ from the correlated experimental data of NIST⁶⁰ are also indicated: single-phase-liquid density $d_{\rm liq}$; single-phase-liquid speed of sound u; saturated-liquid density $d_{\rm sat}$; vapour pressure p_{sat} ; single-phase-liquid heat capacity c_p ; critical temperature T_c ; enthalpy of vaporization ΔH_{vap} .

dubaten eo		Mol	Molecular Para	ameters					% AAL			
annstance	m	$\sigma / \text{Å}$	$\lambda_{\rm r}$	$\lambda_{\rm a}$	$(\varepsilon/k_{\rm B})$ / K	$d_{ m liq}$	n	$d_{\rm sat}$	$p_{\rm sat}$	c_p	$T_{\rm c}$	$\Delta H_{\rm vap}$
Methane	1.0000^{\dagger}	3.7710	19.650	4.4487	113.05	0.50	3.77	0.26	0.25	3.77	1.80	2.55
Ethane	1.8073	3.4133	8.3898	6.7055	158.31	0.57	6.67	0.39	0.41	2.61	2.43	1.47
Propane	1.7610	3.8329	12.954	5.8542	226.30	0.58	9.57	0.69	0.76	2.25	1.32	1.45
Butane	1.6772	4.2495	15.513	5.9910	306.65	0.34	2.43	0.58	0.09	0.93	1.82	1.46
Pentane	1.9068	4.3309	15.075	6.1999	333.12	0.21	2.50	0.34	0.13	0.96	2.44	1.39
Hexane	2.2747	4.2817	14.545	6.0880	320.91	0.28	1.97	0.30	0.14	0.88	2.52	1.69
Heptane	2.3879	4.4257	13.439	6.8824	377.36	0.26	2.15	0.14	0.09	0.93	3.36	1.73
Octane	2.5913	4.4882	13.061	7.1807	399.20	0.38	1.39	0.19	0.09	1.00	3.53	1.71
Nonane	2.7930	4.5379	13.123	7.2319	412.06	0.24	1.71	0.14	0.06	0.89	3.46	1.67
Decane	3.0670	4.5389	13.311	7.1398	412.68	0.35	2.17	0.15	0.08	0.95	3.76	1.95
Dodecane	3.2923	4.7212	15.678	6.8784	454.37	0.40	1.65	0.22	0.24	1.13	3,49	2.07

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that the highlighted parameter is fixed at the indicated value during the parameter estimation. The percentage absolute average deviation $\tilde{\gamma}_0 \tilde{A}AD$ of the various properties calculated with the SAFT-VR Mie EOS⁴⁶ from the correlated experimental data of TABLE IX: Intermolecular potential models developed for various hydrocarbons using generic Mie (λ_r - λ_a) potentials; [†] denotes $NIST^{60}$ are also indicated: single-phase-liquid density d_{liq} ; single-phase-liquid speed of sound u; saturated-liquid density d_{sat} ; vapour pressure p_{sat} ; single-phase-liquid heat capacity c_p ; critical temperature T_c ; enthalpy of vaporization ΔH_{vap} .

auhatan ao		Mol	Molecular Par	Parameters					% AAL			
oubstatice	m	$\sigma / \text{Å}$	$\lambda_{ m r}$	$\lambda_{ m a}$	$(\varepsilon/k_{\rm B})$ / K	$d_{ m liq}$	n	$d_{\rm sat}$	$p_{ m sat}$	c_p	$T_{\rm c}$	$\Delta H_{\rm vap}$
Benzene	1.9897	3.9905	16.59	5.3795	329.67	0.47	1.87	0.23	0.31	2.40	1.21	2.28
Cyclohexane	1.8592	4.3660	18.447	5.3693	349.99	0.37	2.43	0.17	0.25	2.32	2.06	3.26
Cyclopropane	1.0290	4.4323	19.327	6.7466	445.93	1.27	13.58	0.29	0.26	4.71	3.37	5.86
Dimethylpropane	1.0000^{\dagger}	5.6596	50.000	5.2201	462.53	1.09	7.07	0.27	0.41	1.79	2.33	3.50
Ethylene	1.2428	3.7865	10.16	7.8693	242.67	0.46	2.23	0.62	0.23	1.97	0.12	1.44
Isobutane	1.624	4.3159	17.164	5.7007	290.71	0.40	7.40	0.62	0.33	1.68	1.96	1.48
Isohexane	2.2025	4.3415	16.406	5.7892	313.38	0.53	4.08	0.42	1.47	4.68	3.13	1.49
Isopentane	1.8246	4.392	14.632	6.2913	332.07	0.29	3.07	0.36	0.29	0.94	2.57	1.18
Propylene	1.8798	3.6285	11.843	5.8588	204.23	0.57	4.10	0.67	0.74	2.60	1.54	1.58
Propyne	2.4395	3.1374	8.5420	6.7752	190.07	0.32	6.98	0.15	0.52	6.81	0.34	4.51
Toluene	1.8278	4.4263	15.327	6.5140	461.73	0.27	2.09	0.44	0.11	1.28	2.04	1.82

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Accepted Article	TABLE X: Intermolecular potential models developed for various compounds using generic Mie (λ r- λ_a) potentials; ' denotes that the highlighted parameter is fixed at the indicated value during the parameter estimation. The percentage absolute average deviation % AAD of the various properties calculated with the SAFT-VR Mie EOS ⁴⁶ from the correlated experimental data of NIST ⁶⁰ are also indicated: single-phase-liquid density $d_{\rm liq}$; single-phase-liquid speed of sound u ; saturated-liquid density $d_{\rm sat}$; vapour pressure $p_{\rm sat}$; single-phase-liquid heat capacity c_p ; critical temperature T_c ; enthalpy of vaporization $\Delta H_{\rm vap}$. * denotes model taken from Reference ⁴⁶ .
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C. hat an an		Mol	Molecular Para	ameters					% AAD			
annshanne	m	$\sigma / \text{Å}$	$\lambda_{ m r}$	$\lambda_{\rm a}$	$(\varepsilon/k_{\rm B}) \ / \ {\rm K}$	$d_{ m liq}$	n	$d_{\rm sat}$	$p_{\rm sat}$	c_p	$T_{\rm c}$	$\Delta H_{\rm vap}$
Ar	1.0000^{\dagger}	3.4273	17.713	4.6312	93.381	0.47	4.03	0.20	0.17	4.99	1.64	2.93
CO	1.5528	3.0950	9.7981	5.9827	72.21	0.31	2.35	0.15	0.08	2.65	1.61	1.27
CO_2	1.6936	3.0490	26.408	5.055	207.89	1.30	18.80	0.06	0.27	4.99	0.35	3.04
$CO_2 *$	1.5000	3.1916	27.557	5.1646	231.88	0.53	2.40	0.15	0.07	2.64	1.84	3.28
COS	1.2607	3.8273	10.269	7.8823	327.62	0.59	3.57	0.70	0.21	1.82	0.26	1.52
F_2	1.5397	2.7905	9.2354	6.2446	79.713	0.36	6.16	0.25	0.02	1.92	1.74	1.23
Kr	1.0000^{\dagger}	3.6667	18.219	4.5072	123.77	0.43	4.24	0.20	0.16	4.42	1.68	3.16
$ m N_2$	1.4836	3.1243	8.2863	6.6863	70.256	0.59	3.54	0.14	0.05	2.45	1.84	1.39
N_2O	2.0204	2.8616	11.442	5.8897	165.83	0.53	7.62	0.09	0.04	2.05	2.09	3.00
Ne	1.0000^{\dagger}	2.7991	8.8941	6.4568	30.626	0.44	6.90	0.04	0.43	4.31	1.73	2.02
NF_3	1.5434	3.4171	13.456	5.8529	155.22	0.39	5.38	0.67	0.25	2.22	1.04	1.98
O_2	1.4595	2.9427	8.0813	6.4062	80.36	0.57	5.47	0.38	0.22	2.76	2.30	1.03
${ m SF}_6$	1.0000^{\dagger}	4.8811	50.000	5.4185	358.00	1.05	6.66	0.08	0.35	5.81	2.16	4.33
SO_2	2.6207	2.7813	9.7327	6.5569	213.44	0.63	13.23	0.10	0.14	3.13	0.55	2.90
Xe	1.0000^{\dagger}	3.9941	18.910	4.4840	172.29	0.52	5.85	0.20	0.14	5.07	1.85	3.41