Examining the Adsorption (Vapor–Liquid Equilibria) of Short-Chain Hydrocarbons in Low-Density Polyethylene with the SAFT-VR Approach

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The versatility of the SAFT-VR approach in the prediction of fluid-phase equilibria of complex fluids is illustrated for systems of short and long hydrocarbons. In particular, we examine the vapor–liquid equilibria of a mixture of methane + hexatriacontane (C36H74) and the adsorption and coadsorption of light alkanes on low-density polyethylene. The molecules are modeled as chains of square-well segments, and the approach incorporates separate contributions to take into account the effect of monomer–monomer interactions and the bonding of the segments to form chains. In previous work (McCabe, C.; Jackson, G. Phys. Chem. Chem. Phys. 1999, 1, 2057), the phase behavior of pure long-chain n-alkane molecules was examined using the SAFT-VR approach; the intermolecular potential parameters were found to tend to a limiting value as the chain length of the n-alkane increases. Linear relationships with molecular weight for these parameters are used to model the polyethylene polymer in this work.

Introduction

The phase behavior and thermodynamics of hydrocarbon fluids are central to the petrochemical industry in the general context of oil recovery and in the subsequent production of waxes and polymers, which find widespread use. In particular, a knowledge of the phase behavior of mixtures of small and large hydrocarbons is crucial to the design and optimization of polymer production and processing. The equilibrium compositions of the reacting components determine the outcome of the polymerization reaction, and in downstream separation processes, the polymer product needs to be separated from unreacted monomers, solvents, and additives. To obtain a final product with the right properties, it is also important to ensure that the mixture remains in one phase during the polymerization process. Experimental studies under reactor conditions are, however, expensive to run, with safety considerations adding to the cost. In this context, predictive theoretical methods are particularly useful.

Mixtures of solvents and polymers (and of short and long hydrocarbons) generally exhibit a fluid-phase behavior of type IV or V in the scheme of Scott and van Konynenburg. Their phase diagrams are characterized by a discontinuous vapor–liquid critical line, reflecting a phase separation of two dense fluids, one polymer-rich and another solvent-rich, usually in the high-pressure regime; as the temperature is lowered at a fixed pressure, the two phases become miscible at a lower critical solution temperature (LCST). Type IV systems exhibit an additional liquid–liquid immiscibility and upper critical solution temperature (UCST) in the low-temperature region of the phase diagram. In most cases, however, solid phases pre-empt this liquid–liquid immiscibility, and only type V phase behavior is observed. For the largest differences in size between the two components, e.g. for a mixture of methane and hexatriacontane (C36H74), type III phase behavior is observed. Type III phase diagrams are characterized by a large region of liquid–liquid immiscibility, with the locus of UCSTs being found at relatively high temperatures, merging into the vapor–liquid critical line. LCSTs are not found in these systems, but instead, a critical line runs continuously from the critical point of the less volatile component (the polymer) to the locus of the UCSTs. A second small branch of the vapor–liquid critical curve runs from the critical point of the more volatile component (the small hydrocarbon) to the end of the three-phase line. This having been said, we shall not be concerned with critical lines here but rather will focus on the gas–liquid equilibrium in the system, which is of importance in the flash separation step of the polymer production process.

Reliable experimental data are readily available for the thermodynamic and phase equilibrium properties of light (small) hydrocarbons (molecular weight < 500),2–4 but those for heavier hydrocarbons are more scarce. Experimental vapor pressures and saturated liquid densities of n-alkanes up to and including hexatriacontane (C36H74) have been determined, although thermal instability makes accurate experimental measurement extremely difficult for chains longer than decane.7 As one might expect, experimental fluid-phase...
equilibrium data for polymer systems is even more limited, especially at high pressures. Furthermore, typical polymer products are not true pure fluids, but polydispersed mixtures of molecules of varying chain length and degree of branching. The development of an accurate theoretical framework for interpreting and ultimately predicting the thermodynamic properties and phase behavior of polymer systems is hence becoming increasingly important.8

At present, thermodynamic models for polymer solutions fall into two broad categories: activity coefficient models9–11 (e.g., the Flory–Huggins theory,12 polymer NRTL,13 and polymer UNIFAC or UNIFAC-FV14), and equations of state (e.g., Sanchez–Lacombe15–17 cubic equations of state such as polymer-SRK,18 and SAFT19,20). The classic lattice model of Flory and Huggins is still one of the most popular because of its simplicity. In the original approach all lattice sites are occupied, and so the mixture is incompressible. Sanchez and Lacombe15–17 developed a very similar model, with the important difference of allowing for empty lattice sites, in this way bringing in a density and pressure dependence. These models are better suited for studies of systems at liquidlike densities because they are based on a lattice. Cubic equations of state, such as the Soave–Redlich–Kwong (SRK) equation, combined with the Flory–Huggins activity coefficient model have been proposed to overcome this difficulty.21,22 However, the most promising models in terms of predictive capabilities are those derived from the thermodynamic perturbation theory. These approaches are applicable over a wide range of densities, from gas to liquid, and for small and large molecules. In particular, the statistical associating fluid theory19,20 (SAFT) was specifically developed to study chain and associating chain fluids.

The SAFT approach stems from the perturbation theory of Wertheim.23–26 It was originally developed in the late 1980s by Chapman and co-workers19,20 to model associating fluids as chains of Lennard-Jones segments with associating sites. In the equation, separate contributions are incorporated to take into account segment–segment interactions, segment bonding to form chains, association, and dispersion interactions. As SAFT explicitly takes into account molecular nonsphericity, it provides a powerful method for studying the phase behavior of chain molecules. Huang and Radosz27–30 have successfully correlated the equilibrium data of a vast range of pure substances and mixtures, including polymer fluids, using a version of the SAFT approach with a modified attractive term. In the original approach, and in that of Huang and Radosz, a fluid of hard-sphere monomers is taken as the reference system. More recently, a number of modifications have been proposed that incorporate a more accurate description of the reference Lennard-Jones monomer fluid.31–41 These approaches have been used to study alkane and polymer-like fluids, and comparisons have been made with experimental and computer simulation data.

The simplest version of the SAFT approach describes chains of associating hard-sphere (HS) segments with the dispersion interactions described at the van der Waals mean-field level (SAFT-HS).42,43 This approach is especially suited to studies of strongly associating systems, where the association dominates the attractive contribution and the dispersion forces can be adequately described by a mean-field interaction. For example, the critical behavior of n-alkanes,45 the high-pressure critical lines of mixtures of water + alkanes,45 mixtures containing hydrogen fluoride,44 and water + polyethylene surfactant mixtures have been modeled with SAFT-HS.

The SAFT approach has been generalized to describe chain molecules of attractive monomer segments incorporating attractive potentials of variable range (SAFT-VR).48,49 In this case, the long-range dispersion interactions are treated via a high-temperature perturbation expansion up to second order. An additional parameter λ, which characterizes the range of the attractive part of the potential, is used, resulting in a significant improvement over the treatment used in the SAFT-HS approach. In terms of the chain and association contributions, the contact distribution function of the fluid of attractive monomers is used instead of that of the hard-sphere system. Radosz and Addidharma50,51 presented a similar approach (termed SAFT1) for chains of square-well segments adding a term to correct for the truncation of the high-temperature expansion at second order. Using this approach they studied a number of alkane + polymer fluids finding good agreement with experimental data.52–54

In previous work, we have used the SAFT-VR approach extensively to describe the fluid-phase behavior of a wide range of industrially important substances and their mixtures (see, for example, refs 55–61). In particular, we have examined the fluid-phase equilibria of the n-alkanes and their binary mixtures.48,55,56,58 Short n-alkane chains were studied initially, and the change in phase behavior from type I to type V (according to the classification of Scott and van Konynenburg)58 as the molecular weight of one component is increased was illustrated.58 In a comprehensive study, n-alkanes from n-nonane (C9H20) to n-hexatriacontane (C39H80) were examined, and simple linear relationships with molecular weight were obtained for the intermolecular potential parameters.58 Similar prescriptions for the parameters of hydrocarbons have been proposed by Fu and Sandler62 for the simplified SAFT equation and Blas and Vega for the soft-SAFT approach. These relations allow for the determination of the potential model parameters of the longer members of the n-alkane homologous series for which no experimental data are available. The accuracy of this method was confirmed by comparing the SAFT-VR predictions for the saturated liquid density of octadecacontane (C38H80) with pseudo-experimental data from the Gibbs ensemble computer simulations of Smit et al.64,65 given the excellent agreement obtained between the SAFT-VR prediction and the simulation, we extend the approach to study polymeric systems. In particular, we examined the vapor–liquid phase behavior of short-chain n-alkane + low-density polyethylene (LDPE) mixtures. The aim of this paper is to show that, by using the linear relations obtained from the intermolecular potential parameters of the alkanes,58 we can easily determine suitable parameters to model the vapor–liquid phase behavior of polymer systems.

The SAFT-VR Approach

In the SAFT approach, the free energy of a mixture of associating chain molecules can be written as the sum of four separate contributions

$$A_{\text{NKT}} = A_{\text{IDEAL NKT}} + A_{\text{MONO NKT}} + A_{\text{CHAIN NKT}} + A_{\text{ASSOC NKT}}$$
where $N$ is the number of molecules, $k$ is the Boltzmann constant, and $T$ is the temperature. $A^\text{ideal}$ is the ideal free energy, $A^\text{mono}$ is the contribution to the free energy due to the monomeric segments, $A^\text{chain}$ is the contribution due to the formation of a chain of $m$ monomers, and $A^\text{assoc}$ is the contribution due to association. In this work, we do not consider the association term because we are dealing with nonassociating systems. In the SAFT-VR approach, the monomer segments are modeled with an arbitrary attractive potential of variable range $\lambda$ and depth $\epsilon$

$$u(r) = u^\text{HS}(r;\sigma) - \phi(r;\lambda)$$

In the case of square-well monomers, the intermolecular potential is given by

$$u(r) = \begin{cases} +\infty & \text{if } r < \sigma \\ -\epsilon & \text{if } \sigma \leq r < \lambda \sigma \\ 0 & \text{if } r \geq \lambda \sigma \end{cases}$$

where $r$ is the distance and $\sigma$ defines the contact distance between the spherical segments. The monomer free energy is then written as a second-order high-temperature perturbation expansion

$$A^\text{mono} = A^\text{HS} + \frac{1}{NkT} A_1 + \frac{1}{(kT)^2} A_2$$

where $A^\text{HS}$ is the free energy of the reference hard-sphere fluid given by Boublik and Mansoori et al. $A_1$ is the mean attractive attractive energy, and $A_2$ describes the energy fluctuations (see refs 48 and 49 for more details).

As in other versions of the SAFT approach, the free-energy contribution due to chain formation is given by the number of contacts in the chain $(m-1)$ and the contact-pair distribution function of the reference monomer fluid, $g^\text{mono}(\sigma)$, which, for a pure fluid, is given by

$$A^\text{chain} = -(m-1) \ln g^\text{mono}(\sigma)$$

In the SAFT-VR approach, $g^\text{mono}(\sigma)$ is written as a first-order perturbation expansion

$$g^\text{mono}(\sigma) = g^\text{HS}(\sigma) + \frac{1}{kT} g_1(\sigma)$$

where $g^\text{HS}(\sigma)$ is the hard-sphere radial distribution function. A closed form for $g_1(\sigma)$ can be obtained from the pressure expression (the Clausius virial theorem) and the first derivative of the free energy with respect to the density (see ref 48). The pressure and chemical potential needed to evaluate phase equilibria are obtained from the free energy using the standard thermodynamic relations. Phase equilibria between phases I and II in mixtures requires that the temperature, pressure, and chemical potential of each component in each phase be equal, although in this work, we do not allow the polymer into the gas phase. Hence, the condition becomes

$$T^1 = T^II, \quad p^1 = p^II, \quad y_2 = 0, \quad \mu^1 = \mu^II$$

Table 1. SAFT-VR Potential Model Parameters

<table>
<thead>
<tr>
<th>molecule</th>
<th>$M_w$ (g mol$^{-1}$)</th>
<th>$m$</th>
<th>$\lambda$</th>
<th>$\sigma$ (Å)</th>
<th>$\epsilon/k$ (K)</th>
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<tr>
<td>CH$_4$</td>
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<td>3.670</td>
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<td>1.501</td>
<td>3.887</td>
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<td>1.505</td>
<td>3.931</td>
<td>265.0</td>
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<tr>
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<td>12.67</td>
<td>1.565</td>
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<tr>
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<td>1810</td>
<td>1.614</td>
<td>4.026</td>
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</tr>
<tr>
<td>LDPE</td>
<td>248 700</td>
<td>5921</td>
<td>1.614</td>
<td>4.026</td>
<td>264.8</td>
</tr>
</tbody>
</table>

$^a$ $M_w$ is the molecular weight of the molecule, $m$ is the number of spherical segments in the model, $\lambda$ is the square-well range, $\sigma$ is the diameter of each segment, and $\epsilon$ is the square-well depth.

Table 1. SAFT-VR Potential Model Parameters

where $y$ is the mole fraction in the gas phase and the subscripts 1 and 2 denote the short-chain and polymer components of the mixture, respectively.

### Intermolecular Potential Parameters and Results

The alkane molecules are modeled as homonuclear chains formed from $m$ tangent square-well segments of hard-core diameter $\sigma$. A simple empirical relationship developed in earlier work is used to determine the number of spherical segments $m$ from the number of carbon atoms $C$: $m = \frac{3}{2}(C - 1) + 1$. For polymer molecules, the number of carbon atoms is obtained from the average molecular weight. In general, pure-component parameters in the SAFT-VR approach ($\sigma$, $\epsilon$, and $\lambda$) are obtained by fitting to experimental vapor pressure and liquid density data from the triple point to the critical point using a simplex method. The parameters are combined to give an expression for the integrated van der Waals attractive energy $\alpha$; in the case of a square-well potential, the relation is given by $\alpha = 2\pi\epsilon\sigma^3(\lambda^3 - 1)/3$. These parameter values have been determined in previous work for $n$-alkanes from methane to n-hexadecane (C$_{36}$H$_{74}$). Simple linear relations can be used for the longer $n$-alkanes for which limited experimental data is available

$$m\epsilon = 0.039 00M_w + 0.8730$$

$$m\epsilon^3 = 1.566M_w + 24.02$$

$$m\epsilon^3/k = 6.343M_w + 76.38$$

where $M_w$ is the molecular weight of the chain molecule. Using these relations, the parameters of any polyethylene polymer can be determined from its molecular weight. As expected, the parameters tend to limiting values as the molecular weight and chain length of the molecules increase. Average limiting values of the parameters are used for all of the polyethylene polymers; the different polymers are characterized in terms of their size with the chain-length parameter $m$ (see Table 1 for details). To study the fluid-phase behavior of the mixtures, the unlike parameters $\alpha_{ij}$, $\epsilon_{ij}$, and $\lambda_{ij}$ need to be determined. In this work, the unlike size and energy parameters are determined using the Lorentz–Berthelot combining rules, and the unlike range parameter is determined from the arithmetic mean

$$\lambda_{ij} = \frac{\lambda_{ii}\lambda_{jj} + \lambda_{ij}^2\lambda_{jj}^2}{\lambda_{ji} + \lambda_{ij}}$$

This corresponds to the MX3b mixing rule of ref 49. We find that there is no need to readjust the unlike
parameters for the alkane mixtures as the approach accounts explicitly for the molecular size through the parameter $m$ and the segment–segment interactions are of the same nature in the alkane and polyethylene components. However, we also examine the adsorption of an unsaturated short-chain alkane (ethene) in LPDE. As could be expected, in this case the unlike intermolecular parameters need to be readjusted. The optimized intermolecular parameters for ethene are shown in Table 1.

As an extension of earlier work, we concentrate here on mixtures of short- and long-chain (up to polymer-like) alkane molecules. The first system we have studied is the methane ($1) + n$-hexatriacontane ($2$) binary mixture. Four constant-temperature $p_x$slices of the phase diagram in which the SAFT-VR predictions are compared with the experimental data of Marteau et al. are shown in Figure 1. The SAFT-VR approach gives a very good description of the experimental phase behavior, especially in the low-pressure region. As expected for an analytical equation of state, the critical point is overestimated; better agreement in this region can be obtained by rescaling the model parameters to the critical points of the pure components in a mixture (see, for example, refs 55–57), although this is always at the expense of poorer agreement at lower temperatures and pressures. Together with the $p_x$slices, we also compared the SAFT-VR predictions with experimental data for constant-temperature pressure–density $p_F$ diagrams (Figure 2). Excellent agreement is again obtained.

Following the study of methane + $n$-hexatriacontane, it is possible to examine in the same way the vapor–liquid adsorption equilibrium of a short-chain hydrocarbon molecule in low-density polyethylene (LDPE). In the SAFT approach, the extension to longer molecules is straightforward. As mentioned earlier, a relation between the number of carbons in the alkane chain (or the molecular weight of a polymer) and the number of segments $m$ forming the corresponding chain model molecules is used. As the size of the heavier molecule increases, its vapor pressure decreases. As a result, the vapor phase contains practically no polymer (i.e., only the short-chain alkane molecule is found in the vapor phase). To illustrate this point, we compare (in Figure 3) our previous calculation of the phase behavior of methane ($1) + n$-hexatriacontane ($2$) with hexatriacontane in both phases (Figure 1) with the result of a calculation in which $n$-hexatriacontane is restricted to the liquid phase. As can be seen from Figure 3, the calculated phase boundaries of the liquid are almost identical in both cases for this mixture; in the case of heavier polymer molecules, the slight difference observed will become insignificant. In Figure 4, the adsorption equilibrium of $n$-pentane ($1$) in a melt of LDPE of molecular weight 76 000 g/mol ($2$) is examined. The SAFT-VR calculations are compared with the experimental data of Surana et al. at $T = 150 \degree C$. The SAFT-VR approach clearly provides an excellent description of the phase behavior of the system at this temperature. It is especially encouraging to note that no unlike adjustable parameters were used in these theoretical predictions. Note, however, that because we only examine vapor–liquid equilibria in this work, at
T = 150 °C, one should not extend the current calculations to polymer weight fractions of less than 0.5 for this system (the corresponding vapor pressure here is p = 1.6 MPa); liquid–liquid separation is expected beyond this point. As mentioned in the Introduction, liquid–liquid phase separation occurs in mixtures of short and long alkanes as a result of the size difference between the two components. At a constant temperature, the upper pressure limit of the immiscibility corresponds to a critical point, while the lower pressure limit is a point of three-phase coexistence (i.e., where the liquid–liquid region meets the vapor–liquid region, see Figures 3 and 5 of ref 56). This three-phase coexistence typically occurs at pressures just below the pure vapor pressure of the more volatile component (the short alkane), and in the case of pentane, the vapor pressure at T = 150 °C is p = 1.615 MPa. Similarly, we examine the adsorption vapor–liquid equilibrium of ethene (1) in a LDPE of molecular weight 248 700 g/mol (2) (see Figure 5). Good agreement between the SAFT-VR prediction and the experimental data is again obtained for the two constant-temperature pw slices studied, although for the adsorption of ethene, we have used an adjustable binary interaction parameter for the unlike attractive dispersion interaction [ξ = 0.94, where α_{12} = ξ(α_{11} α_{22})^{0.5}]. ξ was obtained by comparison with the experimental vapor–liquid equilibria data at T = 155 °C. It is not surprising that an adjustable parameter is required for this system as the interactions between unsaturated and saturated alkanes are expected to be different than those between saturated alkanes.

Additionally, we have carried out a purely predictive theoretical study of the coadsorption of two short-chain alkanes in LDPE. We consider the different fraction-
The short-chain alkanes is pressure of 2 MPa, the ratio of the weight fractions of and pentane) are more evenly adsorbed in this case, as ever, it can clearly be seen that the two alkanes (butane increases. In terms of the amount of short-chain alkane decrease as the amount of pentane in the gas phase previous one is obtained; the vapor pressure is seen to complete phase diagrams, including also the liquid—liquid equilibria (cloud curves) commonly found in polymer systems.

![Figure 7](image-url)

**Figure 7.** (a) Constant-temperature pw slices of the phase diagram for pentane (1) + butane (2) + LDPE (M_w = 76 000) (3) phase diagram at T = 150 °C. The curves correspond to the calculated polymer weight fractions w_p in the liquid phase for different compositions y of the gas phase. (b) Weight fractions w of each component in the liquid phase for a 50:50 gas composition of the two short-chain alkanes in the pentane (1) + methane (2) + LDPE (M_w = 76 000) (3) ternary system at T = 150 °C. The dotted curve corresponds to pentane, the dashed curve to butane, and the continuous curve to LDPE.

In conclusion, we have used the SAF-VR approach to study the phase behavior of short and long hydrocarbons. Using simple linear relations developed from the intermolecular potential parameters for the n-alkanes from methane to hexatriacontane, we are able to determine parameters for LDPE chains of a given molecular weight. We have examined the adsorption of a number of monomers in LDPE and find excellent agreement with experimental data. In this work, we have concentrated on the vapor—liquid adsorption equilibria of these systems; soon, we plan to examine the complete phase diagrams, including also the liquid—liquid equilibria (cloud curves) commonly found in polymer systems.

Acknowledgment

C. M. thanks Sheffield University for the award of a UGC Scholarship, A. G. thanks the EPSRC for the award of an Advanced Fellowship. M. N. G. thanks BP—Amoco for funding a Research Fellowship. We also acknowledge support from the European Commission, the Royal Society, and the J REI (GR/M94427) and ROPA (GR/N20317) Initiatives of the EPSRC for the provision of computer hardware.

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Received for review February 12, 2001
Revised manuscript received June 15, 2001
Accepted June 20, 2001

IE0101386