A Theoretically Correct Mixing Rule for Cubic Equations of State

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A new mixing rule developed for cubic equations of state equates the excess Helmholtz free energy at infinite pressure from an equation of state to that from an activity coefficient model. Use of the Helmholtz free energy insures that the second virial coefficient calculated from the equation of state has a quadratic composition dependence, as required by statistical mechanics. Consequently, this mixing rule produces the correct low- and high-density limits without being density-dependent.

As a test, the mixing rule is used for ternary mixiutes of cyclohexane + benzene + water, ethanol + benzene + water and carbon dioxide + n-propane + water, and all the constituent binaries. The new mixing rule and a simple cubic equation of state can be used for the accurate correlation of vapor-liquid and liquid-liquid equilibria for binary mixiures. Using the parameters obtained from binary systems, the phase behavior of ternary mixiures can be predicted. Also, unlike previous empirical mixing rules, this theoretically based mixing rule is equally applicable and accurate for simple mixtures containing hydrocarbons and inorganic gases and mixtures containing polar, aromatic and associating species over a wide range of pressures. This mixing rule makes it possible to use a single equation of state model with equal accuracy for mixtures usually described by equations of state and for those traditionally described by activity coefficient models. It is the correct bridge between these two classes of models.

Introduction

the low-density limit they are inconsistent with the statistical mechanical result that the second virial coefficient must be a quadratic function of composition. Further, these rules may fail for simple mixtures (Shibata and Sandler, 1988).

To correct these problems, attempts have been made to develop density-dependent mixing rules (for example, Luedecke and Prausmitz, 1985; Panagiotopoulos and Reid, 1986b) so that the correct low-density limit is recovered. Such an approach, however, is ad hoc and does not preserve the cubic nature of equation of state when used for mixtures. Moreover, as pointed out by Michelsen and Kistenmacher (1990), some of the new mixing rules that have been proposed lead to inconsistencies when a component is split into two or more identical fractions. There have been ongoing efforts to correct such tical fractions. There have been ongoing efforts to correct such

Cubic equations of state are used widely for phase equilibrium calculations. To use such equations to model complexibnes behavior of highly nonideal mixtures, however, mixing rules other than the commonly employed van der Waals one-fluid mixing rules are required. Several authors (Panagioropoulos and Reid, 1986s; Adachi and Sugie, 1986; Sandoval et al., 1989; Schwartzentruber et al., 1989) have proposed various modified forms of the van der Waals mixing rules that use composition-dependent binary interaction parameters. While such mixing rules have been used successfully for some highly nonideal mixtures, they are not generally applicable; since in

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and Reid (1986) was to make ky composition-dependent rebinary pair. The modification proposed by Panagiotopoulos where k_{ϕ} is an interaction parameter characteristic of each

$$a_{ij} = \sqrt{a_{ij}a_{j}}(1-k_{ij})$$
 $b_{ij} = \frac{\lambda}{\lambda}$ (3)

Further, the following combining rules are generally assumed:

fluid" mixing model is: mixing rules. For example, the traditional van der Waals "oncrameters a,, and b,, are made functions of compositions using To apply this equation to mixtures, the equation of state pa-

$$\frac{\sqrt{\Lambda}}{B} - \frac{q - \overline{\Lambda}}{18} = d$$

The van der Waals equation of state is given by:

eter cubic equations of state. pendix; a similar derivation can be used for other two-paramequation for the Peng-Robinson equation is given in the Apvation of the new mixing rule and the fugacity coefficient rule here for the van der Waals equation of state. The deri-For simplicity, we present the derivation of the new mixing

Треогу

and high-density limits without being density-dependent. ing rules produce results that are correct at both the low-density second virial coefficient. Consequently, this new class of mixously, the correct quadratic composition dependence of the excess Helmholtz energy at infinite pressure and, simultaneof any cubic equation of state are determined to give the correct energy parameter a_m and the excluded volume parameter b_m new class of density-independent mixing rules, in which the of considering the Helmholtz free energy, we can develop a mechanical theory. Here we show that by the simple expedient of composition, and therefore is inconsistent with statistical ment that the second virial coefficient be a quadratic function

The Huron-Vidal mixing rule does not satisfy the requirerule for the equation of state parameter bm. free energies instead. This allows a wider choice for the mixing an assumption is not necessary if one equates excess Helmholtz to be used. It was pointed out by Sheng et al. (1992) that such linear mixing tule for the excluded volume parameter b, had otherwise, the PV^E term becomes infinite. Consequently, a require that the excess volume at infinite pressure be zero; from an equation of state be finite, Huron and Vidal had to that the excess Gibbs free energy at infinite pressure computed free energy (activity coefficient) model for liquids. In order culated from an equation of state to that of an excess Oibbs equating the excess Gibbs free energy at infinite pressure calproposed by Huron and Vidal (1979). This method involves An alternative approach for developing mixing rules was

al., 1991), but with little success. deficiencies (Schwartzentruber and Renon, 1991; Mathias et

 $\left(1 - \frac{RL}{q}\right)^{n} = \frac{2}{\left(p^{n} - \frac{RL}{q}\right) + \left(p^{n} - \frac{RL}{q}\right)} \left(1 - p^{n}\right)$

to those of the pure components by: The cross second virial coefficient of Eq. 7 can be related though they may be devoid of any physics.

(01)
$$ATA + \mu a_i x_i x_j = m a_i + F(x)$$
 $A_m = \sum_i x_i x_i x_j x_i + FTF(x)$

Many other algebraic solutions are possible, for example, is also a solution of Eq. 7.

where F(x) is an arbitrary function of composition vector x,

(6)
$$(x) \underline{A}^{\mu} q = {}^{\mu} v$$

pug

(8)
$$\frac{\sqrt{(x)^{q}}-d}{\sqrt{TR}-d}\sqrt{x^{2}x^{2}}\sqrt{\frac{x}{x^{2}}} = \pi d$$

As a disgression, for later reference, we note that: subject of this article.

solution to Eq. 7; other solutions are possible, which is the has been used at all densities. However, Eqs. 2 are only one Waals one-fluid model of Eq. 2. Traditionally this solution but not necessary, condition for satisfying Eq. 7 is the van der second virial coefficient of the equation of state. A sufficient, with $[b-a/(RT)]_{ij}$ being the composition-independent cross

$$p^{\omega} - \frac{q^{\omega}}{BT} = \sum_{i} \sum_{j} x_{i} x_{j} \left(p - \frac{BT}{BT} \right)_{ij}$$
 (7)

we must have that

$$\mathbf{B}_{m}(T) = \sum_{i} x_{i} x_{i} \mathbf{B}_{ij}(T)$$

of the second virial coefficient is quadratic: Since, from statistical mechanics, the composition dependence

$$B(T) = b - \frac{a}{RT}$$

virial coefficient B(T) and the equation of state parameters equation as a virial series, the relation between the second the cubic nature of the equation of state. If we expand the mixing rules being independent of density since this preserves composition dependence. However, there is an advantage to other theoretical restriction on their temperature, density or of the second virial coefficient on composition, there is no While mixing rules should lead to a quadratic dependence

(4)
$$\sum_{n} \sum_{i} x_i x_i \sqrt{a_i a_i} (1 - k_{ij} + x_j t_{ij})$$

solution the molecules are so closely packed that there is no free volume. This limit in an equation of state is:

$$\lim_{m \to \infty} \underline{Y}_{n} = b_{m}$$

$$\lim_{m \to \infty} \underline{Y}_{n} = b_{m}$$

Therefore, if we equate the excess Helmholtz free energy at infinite pressure from an equation of state to that of a liquid solution model we have:

(81)
$$\frac{d^{n}}{d} x + \frac{d^{n}}{d} = (x)^{\frac{3}{2}} \underline{b}$$

Equation 18 is analogous to the relation found by Huron and Vidal (1979), but since they used the excess Gibbs free energy at infinite pressure they had to make the additional assumption on the b parameter of Eq. 3.

Equations 7 and 18 completely define a_m and b_m in terms of $\mathcal{A}_{b_m}^{L}(x)$ (the high-density term) and k_y (the low-density term). These equations can be solved to obtain

(61)
$$\frac{\left(\frac{n}{L}\frac{n}{A}-d\right)_{i}x_{i}x_{i}}{\left(\frac{n}{L}\frac{n}{A}-d\right)_{i}x_{i}x_{i}x_{i}} + 1$$

pu

(02)
$$(3) \stackrel{\infty}{\sim} \sqrt{q} \cdot x \stackrel{\alpha}{\sqrt{q}} = \frac{m^{D}}{q}$$

Note that Eqs. 19 and 20 are in the form of Eqs. 8 and 9 with

$$V(x) = \sum_{i} x_i \frac{a_i}{a_i} - \frac{1}{A} \frac{a_i}{a_i} = (x)$$

Therefore, these mixing rules produce a second virial coefficient which has a quadratic composition dependence, and at low densities these mixing rules predict fugacities similar to a virial equation truncated at the second virial coefficient. At high density, however, the Helmholtz free energy of the solution is that of the chosen liquid activity coefficient model described by $\underline{A}_{\infty}^{E}(x)$, and could be of either a random or local composition form. Consequently, we have obtained a mixture equation form. Consequently, we have obtained a mixture equation of state model that is correct at both the low- and high-density limits without postulating a density-dependent mixing rule.

It should be noted that the mixing rule suggested by Huron and Vidal (1979) is, in effect, the zeroth-order truncation of a series expansion of Eqs. 19 and 20 with respect to ($1/\Upsilon$) and with k_{ψ} set equal to zero. However, since both the excluded volume parameter b and the second virial coefficient are of the order ($a/k\Upsilon$), the proper composition dependence for the second virial coefficient cannot be preserved unless the expansion is carried beyond the first-order term. In fact, there is no need to expand Eqs. 19 and 20 with respect to temperature.

where here k_0 is a second virial coefficient binary interaction

The Helmholtz free energy departure function, which is the difference between the molar Helmholtz free energy of pure species i and the ideal gas at the same temperature and pressure,

$$\left(\underline{A}b \stackrel{A}{q} \stackrel{A}{=} V\right) = (A, T)^{0} \stackrel{A}{\wedge} \underline{A} - (A, T) \stackrel{A}{\wedge} \underline{A}$$

$$\left(\underline{A}b \stackrel{A}{q} \stackrel{A}{=} V\right) -$$

$$\left(\underline{A}b \stackrel{A}{q} \stackrel{A}{=} V\right) -$$

$$(21)$$

for the van der Waals fluid this becomes:

$$\underline{A}_{i}(T, P) - \underline{A}^{iG}_{i}(T, P) = -RT \ln \left[\frac{P(\underline{V} - b_{i})}{RT} \right] - \frac{a_{i}}{L} \quad (13)$$

Similarly, the mixture Helmholtz free energy departure function, which is the difference between the molar Helmholtz free energy of a mixture, $\underline{\Lambda}_m$, and that of the same mixture as an ideal gas, $\underline{\Lambda}_m^{\text{IOM}}$, at the same temperature, pressure, and composition is:

$$= (x, P, x) - \underline{A}_{m}^{IGM}(T, P, x) = -\frac{A}{M} - \frac{A}{M} - \frac{A}{M} - \frac{A}{M} - \frac{A}{M} - \frac{A}{M}$$

$$= (A, P, M) - \frac{A}{M} - \frac{A}{M}$$

Finally, the excess Helmholtz free energy for mixing at constant temperature and pressure, $\underline{A}^{E}(T, P, x)$, is:

since

(61)
$$x_i = \sum_i X_i \Delta_i^{1/0} (T, P) = X T \sum_i X_i \ln x_i$$
 (61)

Expressions for the excess Helmholtz free energy of liquid mixtures have usually been derived using lattice models with the assumption that there are no free sites on the lattice. This is approximately equivalent to the assumption that in a liquid

The parameters in our mixing rule are cross virial coefficient in the expression coefficient and the coefficients in the expression used for the excess Helmholtz free energy. Thus, for example, there are three parameters per binary if the Wilson model is used, and tour parameters (including a) if the NRTL model is used.

In the following examples, we have used the Stryjek and Vera (1986) modification of the Peng-Robinson (1975) equation of state. To get the correct pure-component vapor presures, the equation of state constants for all pure component were computed using the correlations of Stryjek and Vera were computed using the correlations of Stryjek and Vera calculations here, though any other $\underline{\Lambda}^E$ (or \underline{O}^E) model could be used as well. Interaction parameters were obtained by teglession of data for binary mixtures at the temperatures of interest; their values and the results of the correlations are interest; their values and the results of the correlations are predictions using parameters, the temperatures of the provided here.

syndrome is evident from its derivation. rule does not suffer from the so-called Michelsen-Kistenmacher agree well with experimental data. The fact that our mixing ternary system using the mixing model proposed in this work as show in Figure 1, while liquid-liquid predictions for this solubility of water as a function of mixed solvent composition however, the BIP2 mixing rule predicts a maximum in the mixed solvent. With parameters determined from binary data, of water changes almost linearly with the composition of the Cyclohexane and benzene are very similar, and the solubility solubility of water in mixtures of cyclohexane and benzene. by Mathias et al. (1991) using the specific example of the tures containing two similar components. This has been shown tule (1986a) here called BIP2, perform poorly for ternary mixsuch as the density-independent Panagiotopoulos-Reid mixing rules that use two asymmetric binary interaction parameters, of a component in two or more identical components, mixing drome, in which a mixing rule is not invariant to the subdivision Due to the so-called Michelsen-Kistenmacher (1990) syn-

Another system considered here is the binary mixture of ethanol and benzene. When parameters are adjusted to obtain agreement with experimental VLE data for the ethanol-benzene mixture near the two high dilution ends near room temperature, the Panagiotopoulos-Reid mixing tule predicts the formation of a heterogeneous ascotrope and liquid-liquid phase separation at higher pressures. Experimental data, however, show that ethanol and benzene are completely miscible at this temperature. Figure 2 shows that our mixing rule results in much perature. Figure 2 shows that our mixing rule results in much better agreement with experiment, including the prediction of a homogeneous ascottope.

Mathias et al. (1991) have shown that mixing rules with two asymmetric binary interaction parameters, such as that of Panagiotopoulos and Reid, fail to predict a plait point for the ternary system ethanol + benzene + water (experimental data reported in Sørenson and Arlt, 1980c), not will their proposed empirical modification that rectifies the Michelsen-Kistenmacher syndrome. This is probably due to the poor correlation of the ethanol + benzene data, which results from using those of the ethanol + benzene data, which results from using those anixing rules. Figure 3 shows that our model contently predicts a liquid-liquid plait point, though the predicted phase envelope is larger than that found experimentally and contains less water. It also shows the results of the NRTL activity coefficient model prediction using reported parameters obtained from binary prediction using reported parameters obtained from binary

If, instead of our mixing rule, the van der Waals one-fluid mixing model is used, the form of excess Helmholtz free energy of the liquid has then been dictated to be:

Alternatively, if one uses this excess Helmholtz free energy expression in our mixing rule, the van der Waals one-fluid mixing rule is recovered at all densities. However, the mixing rule developed here allows a much more flexible choice of liquid solution model so that, if necessary, local composition the approach developed here is theoretically correct and ean be easily extended to other two-parameter cubic equations of state. For example, the expressions of mixture parameters and fugacities of components derived for the Peng-Robinson equation of state (Peng and Robinson, 1975) are given in the Appendix.

Results and Discussion

In this section we demonstrate the applicability of the mixing tule we have developed by examining experimental vapor-liquid uid, liquid-liquid and vapor-liquid-liquid equilibrium data for several binary and ternary systems at both low pressures and fright pressures, some of which have presented difficulty in other almost ideal to highly nonideal mixtures and include some systems that are traditionally described by equations of state, and others for which activity coefficient models are used. It should be noted that since we can always choose Eq. 22 for should be noted that since we can always choose Eq. 22 for should be noted that since we can always choose Eq. 22 for should be noted that since we can always choose Eq. 22 for should be noted that since we can always choose Eq. 22 for should be noted that since we can always choose Eq. 22 for below, our mixing rule can never be worse than the one-constant van der Waals one-fluid model and therefore will describe below, our mixing rule is also applicable to mixtures that previously could not be accurately described by an equation of state.

In the following, we will use models for the excess Helmholtz free energy, which are normally used for the excess Gibbs free energy. The relation between the two is:

$$\overline{C}_{E} = \overline{V}_{E} + \mathbf{b} \overline{\Lambda}_{E} \tag{53}$$

 $(\lambda x) \quad (\infty = A, x, T)^{2} \underline{h} =$

Excess Gibbs free energy (activity coefficient) models are used generally as low pressure, and \underline{V}^{ϵ} is usually small, so that the difference between \underline{C}^{ϵ} and \underline{A}^{ϵ} is small. Further, if one examines the derivation of \underline{G}^{ϵ} models, it is evident that it is really a model for \underline{A}^{ϵ} which has been derived. Also, due to the absence of the \underline{P}^{ϵ} term, \underline{A}^{ϵ} is much less pressure-dependent than \underline{G}^{ϵ} of the \underline{P}^{ϵ} term, \underline{A}^{ϵ} is much less pressure dependent than \underline{G}^{ϵ} .

$$(W_0) = A_1 \cdot x$$
, $P = low) = \underline{A}^E(T, x, P = low)$

Therefore, in what follows, we will use the same functional form for \underline{A}^k at infinite pressure as is now used for \underline{O}^k at low pressures. It needs to be emphasized that it is not necessary that dependence of \underline{A}^k which is important; it is not necessary that for the liquid phase of any specific mixture to be stable over the whole composition range 0 < x < 1 to use our mixing rule.

Table I. Binary Data Sources, Parameters and Average Absolute Deviation in K Values

BIP2				NRTL						
K & VD	/ _K	дЯ	K ‰ VD	# ₄	ħ _L	n _{ro}	K ⁽⁾	77KJ	Type	məisig
5.1	S Þ '0	ÞI 0 -	4,1	95'6	7L"L	07.0	17.0	867	те	chejo-
										water*
9"1	62.0	21.0 -	9°7	7 0°9	LE"S	07-0	22.0	867	TLE	+ anaznac
2.9	21.0	\$0.0	8.1	PE-0	2.55	9£,0	12.0	867	ALE	vater ^b
m								000		ionati'
₱"L	40.0	41.0-	€€	84.1	60.03	0.30	87.0	867	ALE	thanol + vater ^a
6,€	60.0	40.0	0,2	28.1	4.03	Sp.0	91.0	313	ALE	nobsuc +
	***			31.0	.0 :	U JU	or u	344	ALE	stpou setpsuol
€,₽	81.0	81.0	1,2	91°0 -	20.1 21.1	0£.0	04.0 0.35	116	~3 ~7 A	+ spixoil
€.£	Þ1'0	0'1¢ 0'1¢	£.5	81.0	97 1	06.0	75.0	872) snaqor
L"L	80.0	\$0.0	1.5	72.0	49.0	62.0	8£.0	313	ALE	nodia
										+ əbixoil
										Methanol

*Sørensen and Arli (1980a)

*Sørensen and Arli (1980b).

*Gmebling and Onken (1977)

*Coslivel-Solastiouk et al (1986)

[Reamer et al. (1951).

*Reamer et al. (1951).

*Ohgaki and Katayama (1976).

(for example, dePablo and Prausnitz, 1988). Since our mixing rule uses a classical activity coefficient model to describe the nonideality of the high-density phase, it is not surprising to observe similar behavior.

Schwartzentruber et al. (1987) have pointed out that erronneous liquid-phase splitting is generally predicted when equation of state models with various mixing rules are used to

vapor-liquid and liquid-liquid equilibrium data (Gmehling and Onken, 1977; Gmehling et al., 1988; Sørensen and Arlt, 1980a,b). The equation of state model using our mixing rule gives predictions that are similar to, if not slightly better than, those obtained from the activity coefficient approach. The shortcoming of classical activity coefficient models, especially in the plait point region, is well documented in the literature

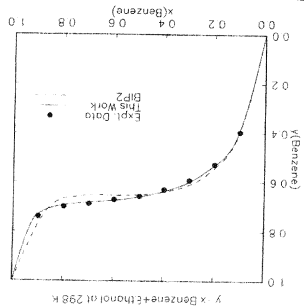


Figure 2. Measured and correlated x-y curves for the benzene-ethanol system at 298 K.

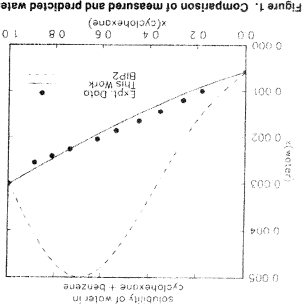


Figure 1. Comparison of measured and predicted water solubility in the benzene and cyclohexane mixture at 298 K.

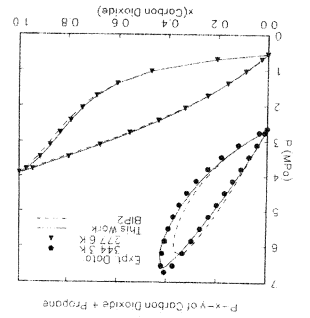


Figure 5. Measured and correlated P-x-y curves for the carbon dioxide-propane mixture at 277.6 and

proposed by Schwarzentruber et al. (1987) avoids the prediction of liquid-liquid separation, but produces a homogeneous ascotrope with a maximum in bubble point pressure near the experimental data that supports the existence of such an ascotrope. On the other hand, our mixing rule predicts the correct phase behavior, in good agreement with the experimental data, without showing false local maxima and minima in the bubble out showing false local maxima and minima in the bubble point pressure curve.

Shibsta and Sandler (1979) have pointed out that some miximg models useful for complex mixtures are unsatisfactory for simple mixtures such as the inorganic gases (nitrogen and carbon dioxide) with hydrocarbons, especially near the critical mixing rule for such systems we have correlated the experimental data for such systems we have correlated the experimental data for such systems we have correlated the experimental data for such on dioxide + propane reported by Reamer et al. (1951) using the mixing rule developed here and, for comparison, the BIP2 model. Figure 5 shows that the difference between the two models is negligible at temperatures below the critical temperature of carbon dioxide (for example, 277.6 K). However, at 344.3 K, the BIP2 model predicts a critical pressure for the binary mixture that is substantially lower than pressure for the binary mixture that is substantially lower than that found experimentally, while the performance of our mixture found experimentally, while the performance of our mix-

We next compared the predictions of our mixing rule with the vapor-liquid equilibrium data for the ternary system carbon dioxide + propane + methanol of Galivel-Solastiouk et al. (1986). In this calculation the vapor-liquid equilibrium data of Ohgaki and Katayama (1976) were used to obtain the interaction parameters for the carbon dioxide + methanol binary system; the other binary mixtures have been discussed above. The experimental data and the predictions using our new mixthe experimental data and the predictions using our new mixthe experimental data and the predictions using our new mixthe experimental data and the predictions using our new mixthe experimental data and the predictions using our new mixthe experimental data and the predictions using our new mixthe experimental data and the BIP2 model at 313.1 K for both 1.21 MPa and 1.71 MPa are compared in Figures 6a and 6b, respectively.

ing model near the critical region is much better.

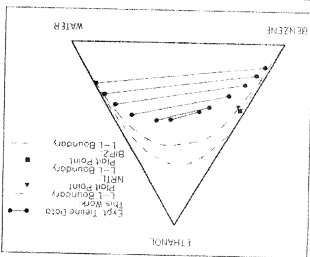


Figure 3. Measured tie lines for the benzene-ethanol-water system at 298 K and predicted phase boundaries and plait points using our mixing rule, the NRTL model, and the Pangiotopoulos-Reid model (which does not predict a plait point).

correlate vapor-liquid equilibrium data for the propane-methanol binary mixture at 313.1 K. The experimental data of Galivel-Solastiouk et al. (1986) in Figure 4 show this system to be highly nonideal. In particular, the bubble point pressure first rises sharply and then abruptly levels off as propane is added to methanol. Schwarzentruber et al. (1987) found that for this system the Huron-Vidal mixing model incorrectly predicts liquid-liquid immiscibility. Figure 4 shows that a simple diets liquid-liquid immiscibility. Figure 4 shows that a simple asymmetric mixing model such as BIP2 gives similar etroneous nearlies. A modified version of the asymmetric mixing model such as BIP2 gives similar etroneous

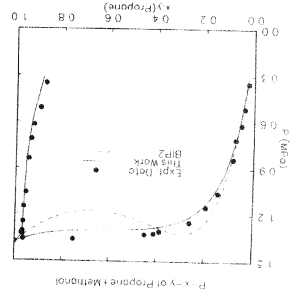
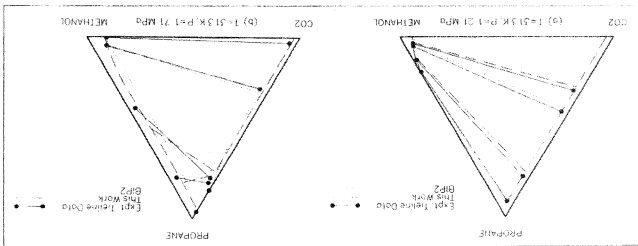


Figure 4. Measured and correlated P-x-y curves for the propane-methanol mixture at 298 K.

p



at 313 K: (a) P= 1.21 MPa; (b) P= 1.71 MPa. Figure 6. Measured and predicted phase boundaries and tie lines for the carbon dioxide-propane-methanol system

Note that in Figure 6a, since the predictions of the two models are similar, only one prediction is shown for each tie line. However, both model

de singiff ni nwork are enoitaibare

it largely predictive. We are now exploring this possibility. 1975) can be incorporated directly into our mixing rule making contribution method such as UNIFAC (Fredenslund et al., calculation of phase behavior. This provides hope that a group same model is used directly in an activity coefficient $(\gamma \! - \! \phi)$ free energy model are quite similar to those found when the

parameter for the second virial coefficient. it should be, since in our development, k_w is a binary interaction consistent with observed virial coefficient behavior. This is as the parameter k_{ψ} may appear large, at first glance, they are ficients. It is noteworthy, however, that while the values of of state to predict quantitatively accurate second virial coefdata for the alcohols and/or the inability of cubic equations result of the generally poor accuracy of second virial coefficient from our correlation of VLE data. This difference may be the carbon-dioxide-methanol system, while we find a value of 0.38 from second virial coefficient data k_{θ} should be 0.70 for the favorably with the value of 0.35 reported in Table 1. However, ide-propane mixture at about 315 K is 0.32, which compares the data in Dymond and Smith (1980), k_{θ} for the carbon dioxcoefficient behavior for nonideal systems. For example, using k_∥ parameter are large, as is to be expected from second virial study. Further, we note from Table 1 that the values of the for $\underline{A}_{\omega}^{\varepsilon}$ provides a reasonable description of the system under of the free energy model used, provided the expression used this, we have found that the value of k_{y} is relatively independent describes second virial coefficient behavior. Consistent with From our derivation, the binary interaction parameter, k,,

Conclusions

of temperature and pressure with few parameters. While simple critical point; and (3) to have models that cover a wide range which can be used at high pressures and close to a vapor-liquid states and hypothetical phase conditions; (2) to have models Among the reasons for this are: (1) to avoid the use of standard activity coefficient models for mixtures with equations of state. A goal of applied thermodynamicists has been to replace

> indistinguishable and agree well with experimental data tic lines calculated by the two mixing models are virtually binary limits as shown in Figure 6a. The phase boundaries and tially methanol) phase boundaries run smoothly between the (essentially carbon dioxide and propane) and liquid- (essenand propane + methanol binaries. The equilibrium vaporvapor-liquid equilibrium exists in the carbon dioxide + methdioxide and propane are gaseous and completely miscible, while pane is about 1.3 MPa, so that at 1.21 MPa mixtures of carbon anol is a liquid at both pressures. The vapor pressure of pro-At 313.1 K, carbon dioxide is a supercritical fluid, while meth-

is able to reproduce these characteristics and certainly much any model to a high degree of accuracy. Nonetheless, our model the tie lines change slope abruptly, is difficult to predict with direction to end on the CO2-propane axis. The region, in which shifting toward the propane-methanol axis, and then changing in slope, starting by being parallel to the CO2-methanol axis, propane concentration is increased, the tie lines change rapidly Note from the experimental data in Figure 6b that as the agreement with the experimental data than the BIP2 model. however, produces a liquid-phase boundary that is in better are correct for each of the binary systems. Our mixing model, correctly predict this feature of the phase diagram, since they as was the case at the lower pressure. Both mixing models pane boundary, rather than the propane + methanol boundary gioxide + methanol boundary to the carbon dioxide + prothe vapor- and liquid-phase boundaries extend from the carbon and carbon dioxide + methanol binary mixtures. Therefore, liquid-phase splitting occurs in the carbon dioxide + propane anol form completely miscible liquid mixtures, and vapor-At L71 MPa, pure propane is a liquid, propane and meth-

in this work. We note that the values of the parameters in the the values and behavior of the parameters that we have found and prediction of phase behavior. It is useful to comment on the mixing rule developed is very good for both correlation For mixtures, which vary from slightly to highly nonideal,

better than the BIP2 model.

- Creek letters
- α = NRTL model parameter, Eq. A17
- r = NRTL model binary interaction parameter, Eq. A17
- y = activity coefficient φ = fugacity coefficient

x = mole fraction vector

x = mole fraction

sidiussuadns

E = excess property

ideal gas = 91

ideal gas mixture = WOI

IM = ideal mixture

siduosans

w = wixinic i. i = molecular species

∞ = infinite pressure state

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= molar volume A = temperature = 892 constant ¥ ∂ quadratic sum of second virial coefficients, Eq. A8 b = bicssureleasty interaction coefficient

p =ednation of state "excluded volume" parameter

= local composition factor for the NRTL model in Eq. A18

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that could not previously be described by equations of state. mixtures and phase behavior, and most importantly, systems

 $\mathcal{A}^{\mathfrak{L}}_{\infty}$, our mixing rule can be used to describe a wide variety of

Since we have complete freedom in choosing the expression

and ternary systems for the diverse systems we have considered. in describing both simple and complex phase behavior of binary

correct, but as we have shown here, it is reasonably accurate by one or the other. Further, not only is the model theoretically encompass mixtures that previously could only be described combining equations of state and excess free energy models to

This mixing rule provides a unified and consistent way of

the combination of the mixing rule and the equation of state.

ence, since the mixing rule is density-independent, but rather

not the mixing rule alone that produces this density depend-

is quadratic in composition. We need to emphasize that it is a virial equation of state with a second virial coefficient that

prescribed by the user at high density and at low densities to equation of state, it converges to the activity coefficient model

while it is independent of density when combined with an

to all mixtures. A unique seature of this mixing rule is that,

is applicable to other equations of state as well), which applies

mixing rule for cubic equations of state (though the procedure In contrast, here we have derived a theoretically correct

mixtures, and/or violate the low-density quadratic composi-

are even density-dependent, apply to only certain classes of

derived Usually, the mixing rules are complicated, perhaps

sequently, many empirical, ad noc mixing rules have been used to obtain the mixture equation of state parameters. Conscribed reasonably well, these failures are due to the models

for example, polar fluids. Since the pure components are de-

poor results are generally obtained with mixtures containing,

tures (mixtures of inorganic gases and hydrocarbons), while

dictions, or at least correlations, are obtained for simple mixsuccess with nonideal mixtures has been limited. Good preto describe the PVT and VLE behavior of pure components,

equations of state with suitably adjusted parameters are able

binary interaction coefficient

O = summation term defined in Eq. A9

C = numerical constant defined in Eq. A4

molat Gibbs free energy

arbitrary function

B = second virial coefficient

a molar Helmholtz free energy a = equation of state "energy" parameter

.... ¥ = O = 8

= 4

Notation

of Delaware.

Acknowledgment

tion boundary condition

lowing expressions for equation of state parameters a_m and b_m : Using the same development as earlier, we obtain the fol-

$$(9V) \qquad \qquad \frac{Q}{(Q-1)} = mQ$$

pug

$$\frac{G}{(G-1)} Q = \frac{m^{D}}{TR}$$

with Q and D defined as:

pue

(64)
$$\frac{1}{\sqrt[m]{p'}} + \frac{CBL}{\sqrt[m]{p'}} \times \sum_{i=1}^{l} -C$$

calculated. The fugacity coefficient is computed from: The thermodynamic properties of a mixture can now be

(01A)
$$\left(\frac{\underline{\underline{Y}}q}{\underline{T}R}\right)$$
 at $-\underline{\underline{Y}}b\left[\frac{1}{\underline{\underline{Y}}} - \frac{q_b}{(n_b)^T}\right]_x^{\infty} = \sqrt[3]{n}$

of mixing rules for am and bm, we have: For the Peng-Robinson equation of state and an arbitrary set

$$\left(1 - \frac{1}{TA} \frac{Q}{Q} \left(\frac{1}{A} \frac{d}{d} \frac{d$$

(11A)
$$\left[\frac{(\overline{\zeta}\sqrt{-1})_m d + \underline{\lambda}}{(\overline{\zeta}\sqrt{+1})_m d + \underline{\lambda}}\right] \text{al}$$

The partial derivatives of a_m and b_m are:

(SIA)
$$\left(\frac{Qn6}{n6} - 1\right)^{\frac{1}{2}(Q-1)} - \left(\frac{Q^{2}n6}{n6} \frac{1}{n}\right) \frac{1}{(Q-1)} = \frac{ndn6}{n6}$$

(EIA)
$$\frac{Qn\delta}{n6} md + \frac{mdn\delta}{n6} Q = \left(\frac{mu^2n\delta}{n} \frac{1}{n}\right) \frac{1}{TR}$$

with the partial derivatives of Q and D given by:

$$(4.14) \qquad \frac{n}{\sqrt{2}} \int x \int_{0}^{\pi} dx \, dx = \int_{0}^{\pi} \frac{1}{\sqrt{2}} \int_{0}$$

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cation of the van der Waals equation of state: Peng and Robinson (1975) proposed the following modifi-

(IA)
$$\frac{RT}{(Y-b)} - \frac{L^2 + 2bY - b^2}{V^2 + 2bY - b^2} = q$$

same as Eq. 5. The Helmholtz free energy departure function series, the expression for the second virial coefficient is the If the compressibility factor PV(RT) is expanded in a virial

sure and composition is: for the Peng-Robinson equation at a given temperature, pres-

(2A)
$$\left[\frac{d(\overline{\zeta} \sqrt{-1}) + \underline{y}}{d(\overline{\zeta} \sqrt{-1}) + \underline{y}} \right] \operatorname{nl} \frac{d}{D} \frac{d}{D} + \frac{d}{D} \frac{$$

In the limit of pressure going to infinity this decomes:

(\$A)
$$\Im \frac{a}{TRd} = \frac{(^{3l}\underline{h} - \underline{h})}{TR} \min_{\alpha = -q}$$

with the constant C being:

$$C = \frac{1}{\sqrt{2}} \ln(\sqrt{2} - 1)$$
 (A4)

:2i (TA)\\&A Therefore, the excess Helmholtz free energy at infinite pressure

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(TIA)

(81A)
$$(a_{ij}\mathbf{x} = a_{ij}\mathbf{x}) \qquad (a_{ij}\mathbf{r}_{ij}\mathbf{x} -)\mathbf{q}\mathbf{x} = a_{ij}\mathbf{x}$$

$$\frac{\partial n}{\partial r} + \frac{\partial n}{\partial r} + \frac{\partial n}{\partial r} = \frac{\partial n}{\partial r}$$

Mith

In this case, the partial derivatives of $\underline{A}_{\infty}^{\mathbb{R}}\backslash(RT)$ with respect to the mole number of each species, which is the logarithm of the species activity coefficient, is given by:

ln
$$\gamma_{\infty}$$
 = $\frac{1}{RT} \frac{\partial n \underline{\Lambda}_{\infty}^E}{\partial n_i}$ (A16)

(61A)
$$\left(\frac{\sqrt{3}\sqrt{3}\sqrt{2}}{\sqrt{3}\sqrt{3}} - \sqrt{1}\right) \frac{\sqrt{3}\sqrt{2}}{\sqrt{3}\sqrt{2}} \sum_{i} + \frac{\sqrt{3}\sqrt{2}\sqrt{2}}{\sqrt{3}\sqrt{2}} = i \infty \gamma \text{nl}$$

Though any free energy model could have been used, we used the NRTL model (Renon and Prausnitz, 1965) here for $\underline{A}_{\infty}^{L}(RT);$

 $\left(\frac{1/3 \eta^2 L_1^2 \chi^2 \sum_{i}}{1/3 \eta^2 L_1^2 \chi^2 \sum_{i}}\right)_{i} \chi \stackrel{\sim}{\sum} = \frac{1}{2} \frac{3}{L_1}$

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