Molecular fluids and mixtures

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<u>Abstract</u> - On the basis of a WCA-like perturbation theory various potentials for polyatomic molecules are discussed. These are multicenter potentials of the Lennard-Jones and the exp-6-8 type, and also other distributions than sitesite potentials. For mixtures, the sensitivity of excess properties to the form of the potential and to shape effects is demonstrated. Various combining rules are discussed for the calculation of the potential between unlike partners from pure component properties.

INTRODUCTION

The approach followed here is that of the perturbation theory. When the principle correctness is assured by a comparison to computer simulation, perturbation theory enables quicker and internally more consistent calculations than any other approach for the high density region. In this way pure components modelled by one center Lennard-Jones molecules (1CLJ), 2CLJ, 3CLJ, 4CLJ and 6CLJ have been dealt with [1-6], and excess properties of mixtures of 1CLJ+1CLJ, 1CLJ+2CLJ, 2CLJ+2CLJ, and 1CLJ+4CLJ type have been calculated [7-10]. On the basis of this experience a short review is given of the merits and shortcomings of the employed WCA-like perturbation theory (for short called the Bochum approach), and on the merits and shortcomings of the multicenter Lennard-Jones potential model. Two directions of modifying this potential model are discussed: (1) employing a different site-site potential, (2) using other distributions than site-site potentials in polyatomic molecules. For mixtures, a discussion of combining rules for the calculating of the potential between unlike partners from pure component properties follows.

THE STATISTICAL-MECHANICAL CONCEPT

The Bochum approach [1,2] is a WCA-like perturbation theory in the molecular frame. That means that variables are the center-center distances and the angles defining mutual orientations of the molecules. Hard fused sphere bodies are used as reference. The pair distribution function of the repulsive assembly is calculated via the Baxter method of solving the Ornstein-Zernike equation with the Percus-Yevick closure. Therefore, the procedure is as follows:

First, the pair potential is calculated for each mutual orientation of the two molecules and divided the WCA way at the minimum into a branch of attractive forces u (r, ω_1, ω_2) and a branch of repulsive forces u (r, ω_1, ω_2) (where ω_1 is a short notation for the orientation of molecule i). Then the angle

averages $\langle e^{-u^{\circ}/kT} \rangle$ and $\langle u^{1}e^{-u^{\circ}/kT} \rangle$ are formed. The residual Helmholtz energy can be written

$$\frac{A^{\star}}{NkT} = \left(\frac{A^{\star}}{NkT}\right)_{0} + \frac{\rho}{2} \int \langle g^{\circ} \frac{u^{1}}{kT} \rangle d\underline{r} , \qquad (1)$$

where u^1 and g° are dependent on the mutual orientation. The essential appro-ximation is now

$$\sigma^{\circ} = \hat{v} e^{-u^{\circ}/kT}$$
(2)

with an angle-independent background correlation function §. With that, the perturbation term of eqn. (1) becomes

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$$(\rho/2) \int \hat{y} < (u^{1}/kT) e^{-u^{*}/kT} > dr$$
 (3)

The background correlation function \hat{y} is calculated from the spherically symmetric potential \hat{u} given by

$$e^{-u/\kappa T} = \langle e^{-u'/\kappa T} \rangle$$
 (4)

Now the only problem left is to lead the Helmholtz energy of the repulsive assembly back to the Helmholtz energy of the hard fused sphere body, which is done by the blip condition

$$\left(\frac{A^{\star}}{NkT}\right)_{O} = \left(\frac{A^{\star}}{NkT}\right)_{H} + \frac{\rho}{2} \int \left(\langle e^{-u}H^{/kT} \rangle - \langle e^{-u^{\circ}/kT} \rangle\right) \hat{y} \, d\underline{r} \quad , \tag{5}$$

where

$$B = \int (\langle e^{-u} H^{/kT} \rangle - \langle e^{-u^{\circ}/kT} \rangle) \hat{y} d\underline{r}$$
 (5a)

is called the blip integral. Assuming that the centers of the spheres of the hard fused sphere body are identical with the centers of the sites in a multicenter site potential, the blip integral can be adjusted to zero by fixing the proper diameter of the spheres. $(A*/NkT)_H$ is then calculated from the Boublik-Nezbeda equation [11]. The mean curvature of the hard fused sphere body is calculated from the envelope, volume and surface of more complicated bodies are calculated using the formulae of Lustig [12].

Before quoting the comparisons of the Bochum approach to computer simulation, other approaches with anisotropic references should be mentioned. There is the approach of Boublik and his group [13], which might be termed a perturbation theory in a Kihara frame, as the variable is the shortest distance between a pair of molecular envelopes. The reference here is a hard convex body, and the Kihara frame pair distribution function (which cannot be traced back to a molecular pair distribution function) is given by a semi-empirical extension of the pair distribution function of hard spheres. Another approach [14] uses the site-site frame (with the site-site distance as variable), and calculates the site-site pair distribution function via RISM. For simple linear molecules, the results of the different approaches are about equivalent. Whereas the Kihara frame theory has more potential to deal approximately with more complicated molecules, the molecular frame theory has the best potential for an extension to anisotropic molecules with electric moments. (For the extensive work almost isotropic molecules with electric moments, cf. of Gubbins, Stell at al. [15,16]).

There exist extensive tests of the Bochum perturbation approach by computer simulation. The first concerns the Helmholtz energy of nitrogen at about the triple point [17]. Then the chemical potential of the 2CLJ liquid with an elongation L = 0.63 has been compared [18] to applications of the Widom test method [19,20]. The internal energy of 2LCJ liquids agreed very well with computer simulations up to an elongation of L = 0.505, but perturbation theory gave 4% too high values at L = 0.67 and 7% too high values at L = 0.793 [21], which is ascribed to the approximation of using an angle-independent background correlation function. Lustig [22] could receive an almost perfect agreement in the thermodynamic properties of 3CLJ propane, calculated by both perturbation theory and computer simulation. Also, in the pair correlation of 4CLJ CF₄ an excellent agreement was observed [6]. Further comparisons were made by Gupta and Coon [23] for 2CLJ liquids and by Sediawan et al. [24] for the Gaussian overlap model. The agreement was always satisfactory. In the last case the Gaussian overlap potential had to be adjusted to make the blip width sufficiently small.

THE PAIR POTENTIAL

It is very important that the parameters of the pair potential are fitted to the liquid state properties. A fit to the second virial coefficients, e.g., can easily result in big errors in the liquid state properties. The reason is that the characteristic potential parameters are less interrelated for the liquid state properties, where the orthobaric liquid density is mainly determined by the size parameter and the vapour pressure is mainly determined by the energy parameter. An appropriate geometric model (the elongation of an anisotropic molecule) is reflected in the slope of the vapour pressure curve. Most experience is accumulated for the multicenter Lennard-Jones model (nCLJ), which has been remarkably successful in reproducing thermodynamic properties of real liquids in spite of the known shortcomings of the Lennard-Jones potential. The shortcomings are essentially: (1) a too steep repulsion, (2) a too shallow minimum, (3) too negative long range tail. Thermodynamic consequences are: (1) The compressibility factor Z has the tendency to become

A /1. m

... 0 /1-00

40.0







Fig. 1. The H potential in compa- Fig. 2. Same as Fig. 1, but rison to the LJ and the Barker-Fisher-Watts potential (BFW) for argon at medium distances.

at short distances.

ZLJ ΖH 6 Zexp Z ŧ 5 300 100 200 400 500 17K

Fig. 3. The compressibility factor Z of methane for the isochore with $\rho = 30 \mod 1$ as function of temperature, in comparison to calculations with the LJ and H potential.

(6)

too positive at high pressures and temperatures; (2) The second virial coef-ficient becomes too positive at low temperatures; (3) The calculated characteristic energy becomes about twice the ionization potential. The last statement should be explained in some detail. According to London [25] the dispersion energy is given by

$$u_{\rm disp} = \frac{3}{4} \frac{\alpha^2}{r^6} h v ,$$

where α is the polarizability and $h\nu$ a characteristic energy, which should be about equal to the ionization potential. As for the Lennard-Jones potential, the attractive term (given by the dispersion energy) is -2 ϵ (ϵ being the well depth) at the minimum of the pair potential, and as the minimum distance is $2^{10}\sigma$, we have

$$\varepsilon = \frac{3}{16} \frac{\alpha^2}{\sigma^6} h \nu , \qquad (7)$$

so that hv can be calculated from the potential parameters and the polarizability.

An attempt has been made [26] to construct a two-parameter potential with an exponential repulsive term and an additional r^{-8} attractive term. In order to limit the number of parameters to two, the exponent in the repulsive term and the C $_{0}/C_{6}$ ratio have been fixed so as to give good orthobaric properties to the rare gase liquids. The resulting potential is

$$\frac{u(r)}{\hat{\epsilon}} = 374887 \exp(-11.2 \frac{r}{\sigma}) - 2.56314 (\frac{\sigma}{r})^6 - 2.56314 (\frac{\sigma}{r})^8$$
(8)

and 2 show how this potential - for short called the Hermann (H) Figures 1 potential - compares to the Lennard-Jones potential and the Barker-Fisher-Watts potential for argon.

The improvements are shown in Figure 3 for the compressibility factor of methane at high temperatures and pressures; in Table 2 for the second virial coefficients of various substances (with the parameters fitted to the orthobaric properties of the liquid and given in Table 1); and in Table 3 for the ratio of the calculated characteristic energy (eqn. 7) to the experimental ionization potential.

TABLE 1. Pair potential parameters adjusted for a fit of the orthobaric properties of the liquid. The first line gives the reduced elongation $L = 1/\sigma$, the second gives σ/A , the third gives $(\epsilon/k)/K$. For substances with L = 0, ϵ/k is one fourth of the CLJ value.

sub- stance	2CLJ	2CH	SSR-LJ	sub- stance	2CLJ	2CH	SSR-LJ
Ar	0.0 3.4039 29.425	0.0 3.4003 31.193	0.0 3.4030 29.434	co	0.39 3.2717 42.282	0.39 3.2862 44.612	0.26 3.3845 39.657
Kr	0.0 3.6272 41.008	0.0 3.6226 43.473	0.0 3.6325 40.972	F2	0.505 2.8317 53.472	0.43 2.9091 51.350	0.29 3.0181 45.720
CH4	0.0 3.7310 37.480	0.0 3.7243 39.775	0.0 3.7247 37.524	с ₂ н ₆	0.67 3.5120 139.81	0.67 3.5208 149.35	0.30 3.9038 100.927
Xe	0.0 3.9517 56.933	0.0 3.9475 60.348	0.0 3.9567 56.867	Cl ₂	0.73 3.2618 201.31	0.73 3.2727 215.45	0.29 3.6921 134.428
°2	0.22 3.2104 38.003	0.22 3.2127 39.990	0.16 3.2604 37.257	^с 2 ^н 4	0.74 3.3268 137.73	0.74 3.3350 147.35	0.30 3.7590 92.923
^N 2	0.3292 3.3078 36.673	0.33 3.3256 38.61	0.22 3.4072 34.537	co ₂	0.793 2.9376 161.83	0.86 2.9051 183.93	0.45 3.2308 134.45

TABLE 2. Second virial coefficients $B_2/cm^3 mol^{-1}$ for various substances, calculated with the 2CLJ, 2CH, SSR-LJ, and Kihara potential. The parameters are from Table 1 or from Boublik [13].

sub- stance	T/K	exp	2CLJ	2CH	SSR-LJ	Kihara
Ar	81 100 150 300 600 1000	$\begin{array}{r} -276 \pm 5 \\ -183.5 \pm 1 \\ -82.2 \pm 1 \\ -15.5 \pm 0.5 \\ +12 \pm 0.5 \\ +22 \pm 1 \end{array}$	-241 -167.4 -81.8 -14.4 +12.6 +21.3	-250.1 -171.4 -82.4 -14.5 +11.8 +20.0		
CH4	110 150 200 300 600	$\begin{array}{rrrr} -330 & \pm 10 \\ -182 & \pm 3 \\ -105 & \pm 2 \\ -41 & \pm 1 \\ +8.5\pm 1 \end{array}$	-285 -166 -99 -41.1 +7.5	-293.2 -168.8 -99.5 -41.1 +6.9		
N ₂	75 100 150 200 300 500 700	$\begin{array}{rrrr} -275 & \pm 8 \\ -160 & \pm 3 \\ -71.5\pm 2 \\ -35.2\pm 1 \\ -4.2\pm 0.5 \\ \pm 16.9\pm 0.5 \\ \pm 24.0\pm 0.5 \end{array}$	-257 -152 -69.7 -34.9 -4.1 +17.4 +25.3	-267 -156 -70.8 -35.5 -4.8 +16.1 +23.6	-255 -152 -35.6 -4.9 +16.4 +24.3	-257.4 -151.4 -68.0 -33.1 -2.4 +18.9 +26.7
F ₂	80 100 150 200 250 300	$\begin{array}{r} -240 & \pm 40 \\ -156 & \pm 10 \\ -70.9 \\ -37.6 \\ -20 \\ -9.5 \end{array}$	-229 -150 -68.7 -36.1 -18.7 -8.0	-236 -153 -69.5 -36.8 -19.4 -8.8	-228 -149 -69.1 -36.8 -19.6 -8.9	-231.1 -150.0 -67.3 -34.4 -16.9 -6.1
с ₂ н ₄	198.77 223.22 250 300 350 450	-315 -251 -201 <u>+2</u> -138 <u>+1</u> -99 <u>+1</u> -51.7 <u>+</u> 1	-314 -250 -200 -137.4 -97.7 -50.0	-327 -258 -205 -140.3 -99.3 -50.8	-294 -237 -191 -134 -96.8 -52.0	-289.3 -232.8 -187.8 -131.1 -94.3 -49.6
с ₂ н ₆	200 260 300 400 600	$\begin{array}{r} -410 & \pm 10 \\ -243 & \pm 2 \\ -182 & \pm 2 \\ -96.0 \pm 1 \\ -24.5 \pm 0.5 \end{array}$	-400 -240 -180 -96.3 -25.3	-418 -246 -184 -97.6 -25.9	-381 -232 -176 -96.0 -27.7	-375.5 -228.6 -173.0 -93.2 -25.0

TABLE 3. The ratio of the characteristic energy to the experimental ionization potential, $h\nu/I$, for various potential models. The parameters are from TAble 1 or from Boublik [13].

substance	2CLJ	2CH	SSR-LJ	Kihara	substance	2CLJ	2CH	SSR-LJ	Kihara
Ar	2.01	1.36	2.01	2.01	$\overset{N}{\underset{\substack{\text{C}_{2}H}\\ \text{C}_{2}H_{4}}{\overset{H}{\underset{\substack{\text{C}_{2}}}}}$	1.87	1.30	2.10	1.23
Kr	1.98	1.34	1.98	1.98		1.79	1.24	2.06	1.07
CH ₄	2.19	1.47	2.19	2.19		2.10	1.46	2.86	1.34
Xe ⁴	2.03	1.38	2.03	2.03		1.80	1.25	2.53	1.28
O ₂	2.48	1.68	2.67	1.40		1.96	1.35	2.90	0.82

Another question concerns the pair potential of polyatomic mole-At present, there are three different suggestions. The most anisotrocules. pic choice is the Kihara potential, then comes the site-site potential, and the least anisotropic is Lucas' SSR-MPA potential [27], where only the repulsive term is distributed to the sites, whereas the attractive term origi-nates from the molecular center. We will make a systematic comparison on the basis of the Lennard-Jones potential acting either between the nearest molecular distances, or between the sites, or with the attractive term between molecular centers. The anisotropy of the dispersion energy, taken into account by Lucas [27], is neglected here. This simplified potential should be called SSR-LJ. As for this potential no parameters are available which are fitted to the orthobaric properties of the liquid, we have determined them on the basis of our perturbation theory with a hard dumbbell reference. These parameters are compared in Table 1 to the 2CLJ-parameters. It is seen that the elongations are systematically smaller, which is compensated by a bigger size parameter o. Figure 4 shows for ethane the potential behaviour for four significant orientations, with the parameters from Table 1 or from Boublik [13], resp.. Figure 5 compares the center-center pair distribution function for the three potentials.

Another significant difference is the ratio of characteristic energy to ionization potential, which is more or less constant for 2CLJ, increases with elongation for SSR-LJ, but decreases with elongation for the Kihara potential. These numbers are given in Table 3. The second virial coefficients, calculated with the potential parameters from the liquid phase properties, are added in Table 2. It seems premature to draw definite conclusions from these comparisons, but at present it can be said that the nCLJ-(or nCH-) potential offers a very reasonable model.







Fig. 4. The 2CLJ(a), SSR-LJ(b) and Kihara(c) potential with the parameters of ethane for the four significant orientations:

tetrahedral	•
parallel	ΠĹ.
T-shaped	1-,
and end to end	



Fig.5. The centercenter pair distribution function for ethane for the 2CLJ, SSR-LJ, and Kihara potential.

EXTENSION TO MIXTURES

The straightforward extension of eqn. (1), (3), (5) to mixtures yields

$$\left(\frac{A^{\star}}{NkT}\right)_{mi} = \left(\frac{A^{\star}}{NkT}\right)_{o,mi} + \frac{\rho}{2} \Sigma \quad x_{i}x_{j} \int \hat{y}_{ij} < (u_{ij}/kT) e^{-u_{ij}^{o}/kT} > d\underline{r}$$
(9)

$$\left(\frac{A^{\star}}{NkT}\right)_{0,mi} = \left(\frac{A^{\star}}{NkT}\right)_{H,mi} + \frac{\rho}{2}\sum_{i,j} x_{i} x_{j} B_{ij}$$
(10)

where the x_{\perp} are mole fractions of component i.

The evaluation of \hat{y}_{1} via Baxter's method has been done in an approximate way by Perram [28] and in a refined version by Fischer and Lago [7]. The ques-tion remains how to deal with the sum of the blip integrals in eqn. (10). Fischer and Lago retained the condition for the pure components $B_{ij} = 0$ and $B_{ij} = 0$, and evaluated the term with B_{ij} as a correction term, whereby ^вјј ر H <e^{-u}/kT

$$e^{-\alpha}$$
ij $^{\prime \times 1}$ > was given by the condition of additivity of sphere diameters

$$d_{ij} = (d_{ii} + d_{jj})/2$$
(11)

Later, following a suggestion by Perram, Bohn et al. [8] used

together with eqn. (11) for determining d_{11} , d_{12} , and d_{22} in the mixture. This way d_{11} became slightly dependent on composition. However, comparison with computer simulaiton showed the advantage of this procedure. Recently, Shukla [29] has used the same conditions.

several comparisons to computer simulation have been made. First, it Again. was possible [30] to reproduce the extensive Monte Carlo calculations of Singer and Singer [31] on model mixtures of spherical molecules. Then, Haile [32] has compared g^E for repulsive soft spheres and Coon et al. [33] have compared excess properties for 1CLJ+2CLJ mixtures. Finally, Lotfi and Fischer [34] have calculated Henry constants for mixtures of spherical molecules by the perturbation theory and by computer simulation. Even in these extreme cases the agreement was very satisfactory.

Two questions should be answered now on the basis of the perturbation theory. First, how sensitive are the excess properties of mixtures of spheri-cal molecules on the pair potential used? Second, how important are shape effects for excess properties?

In dealing with these questions, the model mixtures of Singer and Singer [31] are used as the start. That means that the calculations are for zero pressure, for T = 07 K, for $\epsilon_{12}/k=133.5$ K, and for σ_{12} = 3.596 A. Assuming the Lorentz-Berthelot conditions, i.e.

$$\varepsilon_{12} = \sqrt{\varepsilon_{11} \varepsilon_{22}}$$
(13a)
$$\sigma_{12} = (\sigma_{11} + \sigma_{22})/2 ,$$
(13b)

the excess properties will be calculated as a function of the energy ratio

$$\delta = (\varepsilon_{22}/\varepsilon_{12} - 1)\sqrt{\varepsilon_{12}/\varepsilon_{22}}$$
(14)

and size ratio

and

$$\mu = \sigma_{22} / \sigma_{12} - 1 \qquad (15)$$



$(p = 0, T = 97 \text{ K}, \epsilon_{12}/k = 133.5 \text{ K}, \sigma_{12} = 3.596 \text{ A}).$							
<u>ه</u> م ۲	0	0.06	0.12				
-0.211	-161	-45	+71				
0	-138	-28	+86				
+0.211	-115	- 6	+102				

Fig.6. The excess properties for the Singer and Singer model mixtures for 1CLJ and 1CH potentials.

Figure 6 shows this behaviour for mixtures of 1CLJ molecules (dashed) and 1CH molecules (full). The differences are small, except for extreme cases of δ . The H-potential is much more softly repulsive, which results in slightly more negative v^E values, and in turn in slightly less positive g^E and h^E values.

order to answer the second question, we first model molecule In and molecule 2 as 1CLJ (with the parameters fitted to vapour pressure and orthobaric density). Then we elongate molecule 2 (again with 2CLJ-parameters which satisfy vapour pressure and density). The difference in g^E , which results from this elongation, is given in Table 4 as a function of the 1CLJ energy and size ratios. It is seen that the elongation leads to a negative increment in g^E ,

when we start from equally sized spheres, i.e. when σ_2 becomes smaller than σ_{11} in the elongation process. But when we start from 2 small sphere 1 and a 1 big sphere 2 (μ big), then the elongation process leads to more equally sized σ_{11}

and σ_{22} , and the increment in g^E becomes positive. The dashed lines of Figure 6 and Table 4 are part of our general correlation formulae for the excess properties of 1CLJ+2CLJ mixtures [9].

COMBINING RULES

It is a crucial problem in the treatment of mixtures to assign the unlike interaction parameters ϵ_{12} and σ_{12} , in other words, to determine the small deviations from the Lorentz-Berthelot rule, characterized by the parameters ξ and η :

$$\epsilon_{12} = \xi \sqrt{\epsilon_{11}} \frac{\epsilon_{22}}{\epsilon_{22}}$$
(16a)

$$\sigma_{12} = \eta(\sigma_{11} + \sigma_{22})/2 \tag{16b}$$

For the long range tail of the potentials, the use of the simplified London formulae for the dispersion energy [35] is now well established, with the characteristic energies calculated via eqn. (7). This leads to

$$\epsilon_{12} \sigma_{12}^{6} = (3/8) \alpha_1 \alpha_2 h \nu_1 h \nu_2 / (h \nu_1 + h \nu_2)$$
(17)

For $n_1CLJ + n_2CLJ$ mixtures the factor n_1n_2 has to be added on the left hand side. If σ_1 is assumed to be the arithmetic mean (i.e. $n_1 = 1$), this leads to

$$\xi = \begin{bmatrix} \sqrt{\sigma_{11} \sigma_{22}} & 6 & \sqrt{h\nu_1 h\nu_2} \\ (\sigma_{11} + \sigma_{22})/2 & (18) \end{bmatrix}$$

That means, ξ is given by the ratio of the geometric to arithmetic mean of the σ 's to the sixth power times this ratio for the hv's to the first power. It is, therefore, very important to have reliable σ and hv values for making predictions of the unlike interaction. As it has been shown, this depends very much on choosing the correct molecular model.

The assumption n = 1, on which eqn. (18) is based, has been termed later [8] the old method of Kohler (KO). It is quite a difficult problem to arrive at better values. Kohler et al. [36] thought that it is the effective hard diameter, which is responsible for the repulsive forces, and assumed additivity for it. In order to have a simpler relation than the blip, they used the BH1 prescription [37], which can be formulated generally for nCLJ as

$$\int (\langle e^{-u} H^{/kT} \rangle - \langle e^{-u} o^{/kT} \rangle) dr = 0 , \qquad (19)$$

where $\langle e^{-u} H^{/kT} \rangle$ depends on the hard diameter d. The new relation

$$d_{12} = (d_{11} + d_{22})/2$$
⁽²⁰⁾

requires an iterative solution, which is easily obtained. Note that the integrand in eqn. (19) differs from the integrand in the blip (eqn. 5a) only by the factor $\hat{y}r^2$, so that the new procedure comes very near to choosing a σ_{12} , which leads together with conditions (12) to zero blips $B_{12} = B_{11} = B_{22} = 0.$ It gives mostly n values slightly larger than unity. It has been termed [8] the new method of Kohler (KN), or the new extended method (KNE), when applied to multicenter Lennard-Jones mixtures.

Recently [27], another method for determining η has been recommended, which goes back to the work of Smith [38] and Kong [39]. Their principal assumption can be rewritten as

$$\left(\frac{du_{12}^{rep}}{dr}\right)_{r_{1}+r_{2}} = \left(\frac{du_{1}^{rep}}{dr}\right)_{2r_{1}} = \left(\frac{du_{22}^{rep}}{dr}\right)_{2r_{2}}, \qquad (21)$$

which means that the repulsive force exerted by one molecule for a certain deformation is independent of the nature of the colliding molecule. Though one might question this assumption, the main drawback in the further treat-

ment was the insertion of the r^{-12} part of the LJ-potential in u^{rep} . This leads immediately to

$$\epsilon_{12}\sigma_{12}^{12} = \{ \left[\left(\epsilon_{11}\sigma_{11}^{12} \right)^{1/13} + \left(\epsilon_{22}\sigma_{22}^{12} \right)^{1/13} \right]/2 \}^{13} , \qquad (22)$$

which together with eqn. (17) gives ϵ_{12} and σ_{12} , or ξ and η (this combination might be termed Kohler-Smith-Kong (KSK).

The repulsive potential which is thought to be responsible for the structure is not the r^{-12} part of the LJ-potential, but that part originating from the WCA-division of the potential, i.e.

$$f^{ep} = 4\varepsilon \left[\left(\frac{\sigma}{r}\right)^{12} - \left(\frac{\sigma}{r}\right)^{6} \right] + \varepsilon \qquad r < 2^{1/6} \sigma \tag{23}$$

However, for this potential the condition (21) does not lead to a simple formula like eqn. (22), but gives n values depending on the assumed colliding distance r_1+r_2 . For colliding distances slightly smaller than σ_{12} the n values are of the same order as with the KN rule.

The inconsistency in the repulsive potential used in the KSK rule with that used in the statistical mechanical concept would eliminate the KSK rule if it would not give sometimes relatively good results. For this reason we will discuss it further. It seems to be a fact that the KO or KN rule has a tendency to produce somewhat high ξ values. The use of the very softly repulsive r⁻¹ term of the LJ-potential in the Smith-Kong treatment leads to a relatively big η value, which in turn brings the ξ value down. However, when an independent check on ξ and η is possible, the general finding is that the KSK ξ is quite good but the KSK η is too large. This is illustrated in Figure 7 for the example of the Kr+Xe mixture, which shows what pairs of ξ and η would be demanded to reproduce the excess properties v, g, and h. It is seen that h and g determine essentially ξ , but v is also very strongly

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 $\eta-dependent.$ In that example, the KSK-rule gives a slightly too small $\xi,$ whereas the KN-rule is on the high side. But the η value produced by KSK is definitely too large.

There is another difficulty with the KSK rule for $n_{c}CLJ + n_{c}CLJ$ mixtures. For very small elongations, the rule should go continuously over to the 1LCJ+1CLJ mixture, eqn. (22). Then it should be written

$$n_{1}n_{2}\epsilon_{12}\sigma_{12}^{12} = \{ [(n_{1}^{2}\epsilon_{11}\sigma_{11}^{12})^{1/13} + (n_{2}^{2}\epsilon_{22}\sigma_{22}^{12})^{1/13}]/2 \}^{13} .$$
 (24)

This we will call KSKI. For large elongations, where only different sites "see" each other, formula (22) should be retained, but with the ε_{1} and σ_{1} being the site-site potential parameters. This version we call KSKII. Table 15 gives a summary of various systems calculated by KNE, KSKI, and KSKII.

TABLE 5. Comparison of experimental and predicted excess properties $g^{E}/J \text{ mol}^{-1}$, $h^{E}/J \text{ mol}^{-1}$, and $v^{E}/\text{cm}^{3}\text{mol}^{-1}$ of 101 H201 J and 201 H201 J mixtures

		exp	KNE	KSKI	KSKII
Ar + 0 ₂ 84 K	gE hE v	37 60 0.14	10 16 0.01	-1.6 -11 -0.03	82 121 0.30
Ar + N ₂ 84 K	gE hE v	34 51 -0.18	35 34 -0.15	30 27 -0.18	82 112 0.11
Ar + CO 84 K	gE hE v	57 96 0.09	43 53 -0.02	46 59 -0.01	99 146 0.28
^{CH} 4 + ^N 2 91 K	gE hE v	170 138 -0.54	106 57 -0,57	97 44 -0.63	253 297 0.20
Kr + C ₂ H ₄ 116 K	gE hE v	240 315 0.21	136 154 -0.13	146 144 -0.17	212 239 0.01
Xe + C ₂ H ₄ 161 K	gE hE v	145 185 0.35	108 131 0.07	14 -14 -0.27	275 399 0.63
Ar + C ₂ H ₆ 91 K	ge hE v	317 239 -0.38	191 127 _0.75	422 439 -0.17	272 232 -0.54
Кг + С ₂ Н ₆ 116 К	gE hE v	80 49 -0.22	26 -14 -0.34	79 65 -0.20	56 33 -0.26
^{СН} 4 ^{+С} 2 ^Н 6 104 К	gE hE v	121 74 -0.45	109 59 -0.59	125 85 -0.54	141 106 _0.50
Xe + C ₂ H ₆ 161 K	ge he v	-29 -52 -0.12	3 -19 -0.10	-91 -172 -0.41	92 109 0.18
⁰ 2 ^{+ N} 2 78 К	ge he v	40 60 -0.25	55 59 -0.21	57 62 -0.20	81 100 -0.09
N ₂ + CO 84 K	gE hE v	23 - 0.13	8 11 0.01	14 23 0.05	53 91 0.32
^С 2 ^Н 4 ^{+С} 2 ^Н 6 161 к	ge hE v	99 193 0.16	69 100 0.10	109 162 0.23	113 169 0.25



Fig. 7. The values of ξ and η needed to reproduce the excess properties of Kr + Xe, and the prediction by various combining rules.

We believe that inspite of some shortcomings the KNE rule should be preferred, and that the inconsistency inherent in the KSK rule masks some approximations of the models, which could be easier seen and eventually remedied otherwise.

Acknowledgement

The authors acknowledge gratefully support by the Deutsche Forschungsgemeinschaft.

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