Camel back shaped Kirkwood-Buff Integrals

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Abstract

Some binary mixtures, such as specific alcohol-alkane mixtures, or even water-tbutanol, exhibit two humps “camel back” shaped KBI. This is in sharp contrast with usual KBI of binary mixtures having a single extremum. This extremum is interpreted as the region of maximum concentration fluctuations, and usually occurs in binary mixtures presenting appreciable micro-segregation, and corresponds to where the mixture exhibit a percolation of the two species domains. In this paper, it is shown that two extrema occur in binary mixtures when one species forms "meta-particle" aggregates, the latter which act as a meta-species, and have their own concentration fluctuations, hence their own KBI extremum. This "meta-extremum" occurs at low concentration of the aggregate-forming species (such as alcohol in alkane), and is independant of the other usual extremum observed at mid volume fraction occupancy. These systems are a good illustration of the concept of the duality between concentration fluctuations and micro-segregation.

1 Introduction

The so-called Kirkwood-Buff integrals (KBI) are defined as the integrals of the species-species pair correlation functions $G_{ab} = \frac{1}{\Omega} \int d\mathbf{X}_1 d\mathbf{X}_2 [g_{ab}(\mathbf{X}_1, \mathbf{X}_2) - 1]$ where $\mathbf{X}_i$ is the set of position, and if required, the orientations, of molecule $i$, where $a$ and $b$ designates the species indexes ($\Omega = V \omega^2$, where $\omega$ is the angular integral, equal to $4\pi$ or $8\pi^2$, depending on the symmetry of the molecules). It can be shown that this integral is identical to that involving the pair correlation between any two atoms belonging to each molecules. Following this, the KBI are more simply defined as

$$G_{ab} = 4\pi \int_0^\infty dr r^2 [g_{ia,jb}(r) - 1]$$

(1)
where \( i \) and \( j \) designate any two atoms \( i \) and \( j \) of, respectively, species \( a \) and \( b \), and \( g_{i,j}(r) \) the atom-atom pair correlation function. The Kirkwood-Buff theory relates specific thermodynamic properties, such as the compressibility, or the partial molar volumes, for example, to be related to these integrals, thus providing an appealing link between macroscopic measurable properties to the microscopic structure, albeit in an integrated form, where all microscopic details are lost. Ben-Naim [5], Matteoli and Lepori [6] and other authors [7, 8, 9, 10, 11] have managed to invert these relations, providing a way to calculate these integrals from the experimentally available data on compressibility, partial molar volume and vapour pressure or Gibbs free-energy. The corresponding expressions for a binary mixture \((a, b)\) are well known [12]:

\[
G_{aa} = G_{ab} + \frac{1}{x_b} \left[ \frac{\bar{V}_b}{D} - V \right] \tag{2}
\]

\[
G_{ab} = \frac{\chi_T}{k_BT} - \frac{\bar{V}_a \bar{V}_b}{VD} \tag{3}
\]

were \( x_b \) is the mole fraction of component \( b \), \( \bar{V}_c \) is the partial molar volume of component \( c \) \((c = a \) or \( b)\), \( V \) is the molar volume, \( \chi_T \) is the isothermal compressibility (with \( T \) the temperature and \( k_B \) the Boltzmann constant) and \( D \) is related to the concentration fluctuations and given by

\[
D = x_i \left( \frac{\partial \beta \mu_i}{\partial x_i} \right)_T \tag{4}
\]

where \( \mu_i \) is the chemical component of species \( i \) and \( \beta = 1/k_BT \) is the Boltzmann factor.

Fig.1 of the seminal Matteoli and Lepori paper [6] displays KBI for various types of aqueous mixtures, and it can be seen that most of them have a single extremum, which is positive for like KBI \( G_{aa} \) and negative for then unlike \( G_{ab} \) with \((a \neq b)\). This extremum has its origin in the shape of the coefficient \( D \) above, which is often a U-like shaped curve with a single minimum. An example of such typical KBI and \( D \) are given in Fig.1, and other similar examples can be found in various place in the literature such as [13, 14, 15, 16, 12].

A single extremum for \( D \) is expected for a curve which reaches the same value at both end points. In fact, Eq.\((4)\) can be shown (see Appendix) to be related to the stability limit criteria for a binary mixture, that is precursor for demixing, namely

\[
D \propto C_{11} C_{22} - C_{12}^2 \tag{5}
\]

where \( C_{ij} = \delta_{ij} - \sqrt{\rho_i \rho_j} \tilde{c}_{ij}(k = 0) \), with the \( \tilde{c}_{ij}(k = 0) \) being the integrals of the direct correlation functions and \( i,j \) refer to species index. Indeed, when \( D \to 0 \) the mixture becomes mechanically unstable [3], such when approaching
Figure 1: Examples of single extremum KBI. Red curves experimental KBI curves for aqueous-acetone from Ref.[17]; blue curves for aqueous-2propanol from Ref.[12]. Circles for Water-water KBI, squares for cross water-solute KBI, and triangles for solute-solute KBI. The inset shows the corresponding D (Eq.1) with same color codes, both showing a single minimum.

a spinodal or a critical point. Therefore, a single extremum in $D$ can be physically interpreted as the point where concentration fluctuations are maximal, and with the eventuality that when these grow beyond the mechanical stability, the mixture undergoes demixing. Subsequently, the extremum in the KBI is a signature of maximal concentration fluctuations. Since many binary mixtures, such as aqueous mixtures, for example, show stable local nano-segregation of constituents [18, 19, 20, 21, 22], the extremum can be interpreted as maximal amplitude in local segregation.

In this context, it came as a surprise that Jan Zielkiewicz published experimental KBI for various alcohols-heptane mixtures [23], which clearly showed the existence of two extremum. In Fig.2 of his paper, he shows KBI written as $\rho_a G_{ab}$ where $\rho_a = N_a/V$ is the partial density of species $a$ ($N_a$ is the number of particle in volume $V$), which amplifies the second extremum, the resulting KBI looking like “camel back” shaped, as opposed to the usual single extremum KBI which are “dromadary camel back” shaped.

The purpose of this paper is to understand the origin of this dual extremum in terms of fluctuations. The driving idea is the following. It is well known that alcohols generally tend to cluster their hydroxyl groups into linear patterns, forming chains, loops, etc... [24, 25, 26, 27, 28, 29] When put in small concentration in an inert alkane solvent, such as heptane, for example, small “chain-micelles” of alcohols form and act as independant meta-particles. Consequently, they exhibit their own concentration fluctuations, which corresponds to one of the observed extrema in the corresponding KBI. With increasing concentration, the frontier between well separated micelles and alcohol micro-segregated domains become loose, and the mixture becomes a simple micro-segregated mix-
ture, which has its own independent maximum in concentration fluctuations. It is not obvious that such scenario should happen, and the purpose of this paper is to use computer simulation to provide evidence that this scenario is correct, principally through the analysis of pair correlation functions and the KBI.

2 Theoretical and simulation details

The study of equilibrium locally micro-segregated mixtures has proven to be very difficult in the past decades, particularly in the case of aqueous mixtures. Perhaps a canonical example is that of aqueous tert-butanol (TBA) mixtures, for which the force field induced slow demixing at very small TBA concentrations around $x_{\text{TBA}} \approx 0.01$ was shown \[30, 31, 32, 33, 34\] to rule out many classically robust force field models such as OPLS \[35, 36\] or TraPPE \[37, 38\]. One may even consider this problem to remain unsolved. Similarly, the water-tetrahydrofuran mixture remains difficult to simulate, in particular for putting into evidence the loop coexistence phase diagram \[39, 40, 41\]. One of the principal obstacle for successfully simulating such mixtures is the fact that both water and the alcohol tend to form hydrogen bonded clusters, and it would seem that it is precisely the competition between these cluster formation which tend to drive the simulated mixture into demixing, often after unusually very long times.

These problems might be avoided if water is replaced by a more inert solvent such as an alkane. In a previous work \[42\], we have studied ethanol-benzene mixtures and found that the description of the strong micro-heterogeneity (MH) in the low ethanol content region posed statistical issues which necessitated large size simulations. We expect to find similar issues in the present study.

Herein, we focus on the ethanol-heptane mixtures, simulated in the following range of ethanol mole fractions: 0.02, 0.05, 0.1, 0.2, 0.3, 0.4, 0.5, 0.6, 0.7, 0.8 and 0.9. All of the mixtures contain N=16000 particles, in order to properly describe extended MH. The program package Gromacs was used to perform molecular dynamics simulations \[43, 44\]. The simulation protocol has been the same for all mole fractions of ethanol. The initial random configurations of 16000 molecules were created with the program Packmol \[45\]. These initial configurations were first energy minimized and then equilibrated in the NpT ensemble 5 ns. The length of the production runs varied for each mole fraction of ethanol and can be found in Table 1. The shortest production runs of 10 ns sampled on average 1000 configurations, whereas the longest runs of 35 ns sampled more than 3500 configurations.

The TraPPE forcefield for heptane \[37\] and ethanol \[38\] was used throughout the range of ethanol mole fractions. The mixtures were simulated at $T = 300$ K and $p = 1$ bar. Temperature was maintained constant using mostly the v-rescale \[46\] or Nose–Hoover \[47, 48\] thermostat, while pressure was maintained with the Parrinello–Rahman barostat \[49, 50\]. The temperature algorithms had a time constant of 0.2 ps, while the pressure algorithm was set at 2 ps. The integration algorithm leap-frog \[51\] was used at every time-step of 2 fs. The short-range
interactions were calculated within the 1.5 nm cut-off radius. The electrostatics were handled with the PME method [52], and the constraints with the LINCS algorithm [53].

The snapshots were made with VMD [54].

### 3 Results

#### 3.1 Experiments

The experimental KBI data for the ethanol-heptane mixtures scanned from Zielkiewicz paper are reported as lines in the main panel of Fig.2.

![Figure 2: Experimental “camel back shaped” KBI from Ref.[23] for the ethanol-heptane mixtures, reproduced as lines, with symbols for recalculated values (see text) for consistency check.](image)

In order to make sure that these results are fully consistent with one another, we have inverted Eqs(2,3) in order to extract $D$, using the following assumptions. The partial molar volumes have been replaced by the molar volume of each species, thus making these quantities independent of the concentrations.

| Mole fractions $x_{ETH}$ | Production run times | Mole fractions $x_{ETH}$ | Production run times |
|--------------------------|----------------------|--------------------------|----------------------|
| 0.02                     | 10 ns                | 0.5                      | 25 ns                |
| 0.05                     | 35 ns                | 0.6                      | 20 ns                |
| 0.1                      | 35 ns                | 0.7                      | 15 ns                |
| 0.2                      | 30 ns                | 0.8                      | 10 ns                |
| 0.3                      | 35 ns                | 0.9                      | 10 ns                |
| 0.4                      | 20 ns                |                          |                      |

Table 1: Production run times for the simulated ethanol-heptane mixtures.
The isothermal compressibility term has been neglected (assuming the incompressibility of the liquid mixtures). Finally, the excess volume of the mixture has been set to be a linear function of the pure liquid volumes, hence neglecting the excess volumes. These are usually found to be less that 0.3 cm⁻³/mol. All 3 assumptions can be justified only by the resulting KBI (as was proven by us for aqueous alcohol mixtures in Ref. [12, 21]). From Eqs (2,3) we have extracted the following equivalent expressions for $D$

\begin{align*}
D_{ab} &= -\frac{V_a B_b}{V G_{ab}} \\
D_{aa} &= \frac{x_b V_b - x_a V_a}{f_1 - f_2}
\end{align*}

with $f_1 = x_a x_b (G_{aa} - G_{bb})$ and $f_2 = V (x_a - x_b)$. The resulting values are reported in Fig.3.

Figure 3: Dual minima D for the ethanol-heptane mixtures, as calculated from the experimental results of Ref.[23] (see Eqs. (6,7) in the text).

It is seen that they are quite similar, with the denominator in Eq.(7) producing a singularity at $x_{\text{Eth}} \approx 0.7$. The dots represent the values which have been selected to best represent a good compromise between the two sets. The most prominent feature is the very apparent existence of the 2 extrema, one at $x_{\text{Eth}} \approx 0.2$ and the other at $x_{\text{Eth}} \approx 0.5$. Interestingly, both evaluations of $D$ coincide almost perfectly in the entire range of the minimum part of $D$. Using these values of the extracted D function, we return to evaluate back the 3 KBI integrals, by still preserving the 3 approximations mentioned above. The resulting values are reported in Fig.2 as dots, and are seen to nicely superpose to the original data. This simple exercise proves that a single form of $D$, which must be quite close to the one reported in the inset, must have served in Ref.[23] to calculate the KBI, and moreover, that the use of true partial molar volumes, compressibilities and excess volume did not affect much the final shapes. We
also prove that the 2 extrema of the KBI originate from the 2 extrema observed in $D$.

### 3.2 Computer simulations

#### 3.2.1 Snapshots

When mixtures present strong local heterogeneity, it is often instructive to look at snapshots in order to visually appreciate the nature and the extent of the concentration dependence of the spatial segregation. Fig.4 shows snapshots of 4 typical concentrations of ethanol, with different styles highlighting the morphological changes in the ethanol clustering in heptane solvent.

![Snapshots of ethanol-heptane mixtures](image)

**Figure 4:** Snapshots of the ethanol-heptane mixtures for 4 typical ethanol concentrations. Ethanol oxygen, hydrogen and carbon groups, in red, white and cyan, respectively, and heptane carbon groups in blue. Heptane is shown as ghost pale blue for $x = 0.1$ and 0.2. Ethanol carbon groups are also shown as ghost pale blue for $x = 0.2$. (see text for details).

For ethanol concentration $x = 0.1$, full ethanol molecules are shown (oxygen atom in red, hydrogen in white and carbon groups in cyan) while heptane molecules are shown in transparent mode. It can be seen that small droplets of ethanol float in the midst of heptane solvent. For $x = 0.3$, only the hydroxyl groups are fully shown, with all carbon groups of both species shown in transparent mode. This way, hydroxyl clusters and their distribution are highlighted. For $x = 0.5$ and $x = 0.8$, ethanol molecules are fully shown like for $x = 0.1$, and heptane molecules are shown in blue. These last 2 snapshots highlight the clear micro-segregation of both species.
Fig. 5 shows a zoom on the hydroxyl group clustering for the same 4 ethanol concentrations highlighted in Fig. 4. For $x = 0.1$ one sees that the small ethanol droplets of Fig. 4 consist in fact in pentameric rings of hydroxyl groups, forming ethanol pentameric “micelles”. This structure vanishes for the other higher ethanol concentrations shown in Fig. 5, replaced by more complex cluster structures, such as globules and various type of chains conformations.

Detailed studies of snapshots for the entire concentration range shows that ethanol molecules start to spontaneously form these pentameric micelles at the smallest ethanol concentrations studied herein, which is $x = 0.02$. These pentameric micelle structures persist until $x = 0.3$, after which larger aggregated structures take over. The visual inspection of formed structure tends to confirm that there are 2 regime of clustering, pentameric ethanol micelles for $x < 0.3$ and larger clusters for higher concentrations. We now confirm this through cluster and pair correlation function studies.

### 3.2.2 Cluster structure

The cluster size probability distributions were calculated as:

$$P_n = \frac{\sum_k s(n,k)}{\sum_k \sum_n s(n,k)}$$

where $P_n$ is the probability for the cluster formed of $n$ sites, $s(n,k)$ represents the number of clusters of the size $n$ in the configuration $k$. $P_n$ is obtained by averaging the number $s(n,k)$ of clusters of size $n$ over several such configurations. The cut-off distance ($r_c$) for a calculation depends on the first minimum.
in the pair correlation function of a particular site. For the oxygens in ethanol, the $r_c = 3.7$ Å.

Fig. 6 shows the size $s$ dependence of the cluster distributions $P(s)$ in 2 different ways. Left panel (a) shows the distribution as function of the cluster size, and for different ethanol concentrations. What is apparent is the pentamer peak, which exists for all concentrations. In addition, we observe the appearance of a shoulder-like feature for large clusters, showing that the probability of larger clusters is not negligible for ethanol concentrations above $x > 0.3$. Moreover, while the probability of monomers is higher than that of pentamers for $x < 0.3$, and becomes comparable or lower for $x > 0.3$.

Figure 6: Cluster distribution probability function $P(s)$ as function of (a) cluster size $s$ and (b) ethanol concentration. (See text for details).

The right panel (b) shows $P(s)$ as function of ethanol concentration, and for different types of $n$-mers. What is strikingly apparent here is that, for the entire concentration range, there is a sharp fall of $P(s)$ from monomer to dimers and 3-mers, followed by a second dramatic increase for 4-mers and 5-mers, followed by 6-mers, while higher $n$-mers are the same level as monomers and dimers. An important feature is that the maximum of the curves seems to saturate around $x = 0.3$, indicating a turnover of cluster structures above this concentration.

The cluster study complements the visual inspection of Fig. 5 by showing more details, such as for example the existence of 4-mers at small concentrations, and not only pentamers, as the visual inspection may suggest.

3.2.3 Atom-atom pair correlation functions

In order to further confirm the sharp separation of cluster structure highlighted in the previous sections, we show in Fig. 7 the evolution of 3 typical pair correlation functions as a function of ethanol concentration. The right panel shows the ethanol oxygen-oxygen correlation function $g_{OO}(r)$ for all the ethanol concentrations calculated. Focusing on the second peak, which represent second neighbour correlations, hence the influence of clustering beyond the first neigh-
bour peak, we clearly a difference between two successive concentrations: the
gap is wider for $x < 0.3$. In order to facilitate this observation, the curve for
$x = 0.3$ is marked in thicker orange color.

The narrower gap for $x > 0.3$ can be interpreted as lesser difference in the
second neighbour correlations as a mark of insensitivity for the concentration
dependence. This is consistent with hydroxyl clusters being of more varied
shapes. In contrast, the almost similar gap for $x < 0.3$ shows a linear dependence
in concentration, which would be expected if clustering was the same and would
only depend on the concentration of the pentamers.

In order to further confirm this trend, we examine in the middle panel of
Fig.7 the cross species correlations $g_{OM_1}(r)$ between the ethanol oxygen and the
heptane first/last carbon group termed M1. These correlations are seen to be
less concentration dependent than the previously examined $g_{OO}(r)$ correlations.
This is expected, since heptane site are not charged, hence only Lennard-Jones
like correlations exist, which are less prominent then for those between charged
groups [42]. Nevertheless, we observe that for $x < 0.3$ all correlation functions
are nearly superposed (below the thick orange curve), and start to show appreciable differences only when $x > 0.3$. Again, this is fully consistent with the
existence of same type of clusters for $x < 0.3$. Indeed, is clustered objects are
the same, the correlations between the carbon sites of heptane and the meta
objects would be nearly similar, and very weakly dependent on $x$. In contrast,
if the clustered objects are very different in shape, and if this depends strongly
on ethanol concentration, we would indeed expect a larger concentration dependence of $g_{OM_1}(r)$.

The last right panel Fig.7 shows a weak concentration correlations between
the heptane last/first carbon site $g_{M_1M_1}(r)$, very similar to that observed for
$g_{OM_1}(r)$. First of all these are Lennard-Jones like correlations, and correlations
below $x<0.3$ are also nearly superimposed, witnessing the same insensitivity to concentration in presence of same type of aggregates of the ethanol.

If we reconsider now the experimental KBI in the light of this sharp separation of the behaviour of the correlations for $x<0.3$ and for $x>0.3$, we obtain the result shown in Fig.8, where the thick orange vertical line at $x=0.3$ perfectly separates the concentrations under the first KBI extremum for $x<0.3$, where mostly pentameric ethanol micelles are observed, from the concentrations under the second weaker peak at right for $x>0.3$, for which usual micro-segregation of the 2 species is observed. In order to fully confirm this picture of sharp separation between a meta-object mixture and a usual mixture, we need to calculate the KBI from the simulations and reproduce the same KBI shapes as observed in experiments.

### 3.3 Kirkwood-Buff integrals from simulations

The evaluation of the KBI through computer simulation requires that the asymptotes of the various site-site correlations are well defined and converging to 1 as expected. Recent investigations have shown that there are 2 major obstacles. The first obstacle is that computer simulations conducted in the Canonical or Isobaric ensemble cannot lead to the proper asymptote 1, this value being reached only in the Grand Canonical ensemble \[55, 56, 57\]. The second obstacle concerns the existence of micro-segregation, whose spatial extent and slow kinetics alters the statistics of the correlations at large atom-atom separations. Both cases are illustrated below, where we examine both the tail of the correlation
functions $g_{ab}(r)$, but also the so-called running KBI defined as (1):

$$G_{ab}(r) = 4\pi \int_0^r ds s^2 [g_{ab}(s) - 1]$$

which, when $r$ is large enough, is expected to converge to the KBI defined in Eq. (1).

In a first example, we examine the case of ethanol concentration $x = 0.7$, for which we have shown above that micro-segregation is dominant. The typical order parameter for micro-segregation is $g_{OM_1}(r)$, the cross species pair correlation between the ethanol oxygen atom and the first/last carbon group atom of heptanol. Fig.9 shown this function in the main panel for 3 different runs of 5ns each. The strong depletion between adverse ethanol and heptane nano-domains is clearly visible, as all 3 curves stay below 1 until 20 Å or so.

Figure 9: Long range tail contribution for ethanol-heptane cross correlation function $g_{OM_1}(r)$ for ethanol concentration $x = 0.7$ shown in the main panel for 3 different runs of 5ns each. The upper inset is a close-up of the tail part. The lower inset shows the respective RKBI $G_{OM_1}(r)$ functions, along with the experimental KBI (in blue).

On this scale, all 3 curves seem nearly similar, which is what is expected from statistics. However, the zoom on tail region in the upper inset shows that there are visible differences between the 3 runs, as the segregation of domain is not the same in each of the 3 runs. What is observed is the typical domains oscillatory correlations, with half-period about 20 Å, which is the depletion extent. All 3 curves oscillated about the asymptote 1, hence the first obstacle mentioned is not really apparent. The lower inset shows the corresponding running KBI $G_{OM_1}(r)$, as well as the expected experimental value. It is seen that the various runs have an oscillatory asymptotic feature, witnessing the domains alternation, but
Figure 10: Long range tail contribution for ethanol-heptane cross correlation function $g_{OM_1}(r)$ for ethanol concentration $x = 0.2$ shown in the main panel for 5 different runs of 5ns each. The upper inset is a close-up of the tail part. The lower inset shows the respective RKBI $G_{OM_1}(r)$ functions, along with the experimental KBI (in purple).

oscillate around the experimental value. This result shows the rather excellent agreement between the experimental KBI and the calculated one.

In the second example, we examine the case of ethanol concentration $x = 0.2$, for which we expect an homogeneous distribution of ethanol micelles in the midst of heptane solvent. In this case, micro-segregation is very different in that it generates these micelles. Fig. 10 shows the same order parameter $g_{OM_1}(r)$ in the main panel, and for 5 different runs of 5ns each.

This function again shows appreciable depletion, but of the size of the ethanol micelles, which is more about 12Å. But now, the zoom of the tail in the upper inset shows that there are no domain oscillations, which is indeed expected if the ethanol droplets are homogeneous distributed. However, we observe that all the asymptote are clearly shifted upwards from 1. This is a direct manifestation of the first obstacle mentioned above, which occurs here since our calculations are in the isobaric ensemble instead of the Grand canonical ensemble. It can be shown that the asymptote of the correlation functions has the following form

$$\lim_{r \to \infty} g_{ab}(r) = 1 - \frac{\epsilon_{ab}}{N}$$

where $N$ is the number of particles in the simulation box, and the value of $\epsilon_{ab}$ depend both on species $a,b$ and the statistical ensemble [55, 56, 57]. For like correlations the shift is downwards from 1, and for unlike it is upwards [42]. The RKBI are shown in the lower inset, and are seen to lie above the experimental value, and not to have the expected flat asymptote for a proper definition of the KBI value for $G_{OM_1}$. In order to correct for this “spurious” behaviour, we
simply multiply the pair correlation function with the appropriate coefficient 
\( \gamma_{ab} = 1/(1 - \frac{\epsilon_{ab}}{N}) \) before applying Eq.(10) to 
\( \bar{g}_{OM_1}(r) = \gamma_{OM_1} g_{OM_1}(r) \), which 
now has the correct asymptote 1. The corresponding results are shown in Fig.11.

![Figure 11: Tail corrected version of pair correlation functions shown Fig.10 (see text)](image)

Due to the extreme smallness of \( \frac{\epsilon_{ab}}{N} \), \( \gamma_{ab} \) is nearly 1 and the main panel 
shows no difference with respect for Fig.10. However, while the upper inset 
confirms that the correct asymptote 1 has been set by the operation above, the 
lower inset shows that all the RKBI have now flat asymptotes, even though it is 
not quite the value set by the experiments. This discrepancy is expected, since 
the ethanol aggregation in the real mixtures may not be well represented by the 
model simulations.

For the calculations of the KBI, from the cross KBI \( G_{OM_1} \) we extract D 
from Eq.(6) and calculate the like KBI \( G_{OO} \) and \( G_{M_1M_1} \) using Eq.(7). Indeed, 
the simulations results for these KBI do not often converge to the consistency 
for D in Eqs.(7,6), and the above trick is one way to get around this problem. 
The origin of the inconsistency is due to system size problems, since the like 
domain statistics require larger simulations, with \( N=32000 \) particles or more 
[32, 33, 34]. When these operations are conducted for all ethanol concentrations, 
we obtain the result shown in Fig.12, which shows a remarkable agreement with 
the experimental results.

The very low ethanol concentrations for \( x = 0.02 \) and \( x = 0.05 \) pose particular 
problems for the proper evaluation of the asymptotes. Indeed, the correspon-
ding RKBI have very distorted shapes, which cannot be arranged to look 
flat by simple shifting of the asymptotes of the pair correlation functions. These 
distortions, which vary considerably from one run the other, witness the strong 
kinetics of the ethanol micelle formation. Since these micelles are about 15Å in 
diameter, while the half box length extends to 70Å, if the micelles were replaced
by soft spheres of same diameter, the statistics on the tail of the soft sphere correlations would be much better. Therefore, it is really the micelle-monomer exchange which affect those statistics at very small ethanol concentrations.

4 Discussion and Conclusion

It is not a priori obvious that the segregation of ethanol in small concentration in oil-like solvent should evolve in a discontinuous manner into the micro-segregation as this concentration is increased, leading to the separation illustrated in Fig.8. For instance, in a previous study of ethanol-benzene mixtures [42], we have found a single extremum shaped KBI. Yet, this study has shown that small micelle-like aggregates at low ethanol mole fraction mixtures. This difference suggests that closed carbon group molecules, such as benzene, affect differently the cluster structural changes from micelle to domain, than chain-like shaped alkanes. Indeed, these latter molecules can more easily merge with the micellar alkane corona than the disc-shaped benzene molecules. In order to explain this apparent discrepancy, we hypothetize here that it is solvent shape induced depletion entropic effects [58, 59, 60] which affect the structural transition between the ethanol micelles regime and the micro-segregated domain regime. Following this hypothesis, chain-like solvent would allow a smoother transition between micelles and domains, hence clearly separating the two manifestations in terms of fluctuations, thus leading to separate fluctuation regimes.

In the abstract, we mention that water tert-butanol mixtures equally show two extrema in KBI. This was indeed reported our previous work[12], where the KBI calculated by us (see Fig.6b in Ref.[12]) were obtained by Y. Koga from the vapour pressure measurements of the chemical potentials (see the plot of $D$ in Fig.6a of Ref.[12]). The double extrema is absent from the KBI data from
other authors (see Fig.6b in Ref.[12]), but present in the SANS data obtained in the same work. Interestingly, in both cases, the double extrema occurs clearly only for the water-water KBI. In addition, the equivalent of the micelle extrema - in the present case it would be tbutanol micellar aggregates, occurs for very small tbutanol concentrations in the interval \( 0.1 < x < 0.2 \), which is similar to that observed in the present work for an entirely different system. Interestingly, the aggregation in aqueous tbutanol is not clearly understood to date, despite several experimental [61, 62] and simulation [31, 32, 34, 63] investigations.

From these two points, it appears that the topic of the existence of camel back shaped KBI deserves further investigations as to the conditions where it might occur.

To conclude, the present study illustrates the difference between concentration fluctuations and micro-segregation both in the microscopic and macroscopic thermodynamic level. It is generally believed that micro-segregation would correspond to a more microscopic \( k \neq 0 \) part of the concentration fluctuations, related to the finite extent of segregated domains, while concentration fluctuations would be their thermodynamic limit, which correspond to \( k = 0 \). The present study shown that the KBI, which are related to the macroscopic \( k = 0 \) limit of the structure factors, themselves contain the difference in both manifestations. This was illustrated in Fig.8, through the sharp separation between concentration fluctuations of meta-objects (the ethanol micelles) for \( x < 0.3 \) and the micro-segregation regime for \( x > 0.3 \). To be more specific, while micro-segregation concerns the initial microscopic objects, namely ethanol and heptane molecules, the concentration fluctuations described here concern the meta-objects formed by the ethanol micelles, themselves floating in molecular heptane solvent. This study has been possible, precisely because heptane is an inert solvent, due to its uncharged carbon group sites, which cannot form associated entities. In this context, heptane concentration fluctuations are very small, and the neutrality of this oil-solvent enhances the charge association of ethanol molecules. This is not possible to observe with water-solvent and small alcohol molecules, because both species tend to associate. In order to observe a similar phenomena in water, one requires much larger solutes molecules, such as surfactants, which can form micelles and other self-assembled objects.

The present study also illustrates the importance of the pair correlation functions and their asymptotes, and through these quantities, the issues related to domain segregation and concentration fluctuations in finite size simulations, in particular through the Lebowitz-Percus shift of the asymptotes, illustrate here for the fluctuations between meta-objects, as opposed to the usual illustration for molecular objects.

Appendix

Eq.(2) in the Kirkwood-Buff paper [1] expresses the derivative of the chemical potential \( \mu_i \) of species \( i \) with respect to its mole fraction \( x_i \) in terms of the KBI, which we rewrite below using a trivial rearrangement of the original equation.
as:
\[
\left( \frac{\partial \beta \mu_i}{\partial x_i} \right)_{T,P} = \frac{1}{x_i [1 + \rho_j x_i (G_{11} + G_{22} - 2G_{12})]}
\]

The Ornstein-Zernike equation for a binary mixture can be written in a matrix equation as
\[
\begin{pmatrix}
S_{11} & S_{12} \\
S_{12} & S_{22}
\end{pmatrix}
\begin{pmatrix}
C_{11} & C_{12} \\
C_{12} & C_{22}
\end{pmatrix}
= I
\]
where \(I\) is the identity matrix, and where the structure matrix elements are \(S_{ij}(k) = \delta_{ij} + \sqrt{x_i x_j} \tilde{h}_{ij}(k)\) and the C-matrix elements are defined in terms of the direct correlation functions as \(C_{ij}(k) = \delta_{ij} - \sqrt{x_i x_j} \tilde{c}_{ij}(k)\), with the Fourier transform notation and definition \(\tilde{f}(k) = \int d\vec{r} f(\vec{r}) \exp(i\vec{r} \cdot \vec{k})\). Since the KBI are defined as in Eq.(1), one has \(G_{ij} = \tilde{h}_{ij}(k = 0)\), and \(S_{ij} = \delta_{ij} + \sqrt{x_i x_j} G_{ij}\).

Therefore, one can rewrite the equation above as (using \(\rho_i = x_i \rho\))
\[
\left( \frac{\partial \beta \mu_i}{\partial x_i} \right)_{T,P} = \frac{1}{x_i \sqrt{x_1 x_2} [S_{11} + S_{22} - 2S_{12}]}
\]

which can be rewritten by using the OZ equation as
\[
\left( \frac{\partial \beta \mu_i}{\partial x_i} \right)_{T,P} = \frac{\det C}{x_i \sqrt{x_1 x_2} [C_{11} + C_{22} + 2C_{12}]}
\]

Using Eq.(4) which defines the KBI coefficient \(D\), one has the relation relating \(D\) to the determinant of \(C\), which controls the stability limit of a mixture [3], as written in Eq.(5)
\[
D = \alpha \left[ C_{11}C_{22} - C_{12}^2 \right]
\]
with \(\alpha = 1/(\sqrt{x_1 x_2} [C_{11} + C_{22} + 2C_{12}])\).

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