Microscopic structure of liquid 1-1-1-2-tetrafluoroethane (R134a) from Monte Carlo simulation†

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1-1-1-2-tetrafluoroethane (R134a) is one of the most commonly used refrigerants. Its thermophysical properties are important for evaluating the performance of refrigeration cycles. These can be obtained via computer simulation, with an insight into the microscopic structure of the liquid, which is not accessible to experiment. In this paper, vapour–liquid equilibrium properties of R134a and its liquid microscopic structure are investigated using coupled–decoupled configurational-bias Monte Carlo simulation in the Gibbs ensemble, with a recent potential [J. Phys. Chem. B 2009, 113, 178]. We find that the simulations agree well with the experimental data, except at the vicinity of the critical region. Liquid R134a packs like liquid argon, with a coordination number in the first solvation shell of 12 at 260 K. The nearest neighbours prefer to be localized in three different spaces around the central molecule, in such a manner that the dipole moments are in a parallel alignment. Analysis of the pair interaction energy shows clear association of R134a molecules, but no evidence for C–H · · · F type hydrogen bonding is found. The above findings should be of relevance to a broad range of fluoroalkanes.

1. Introduction

Hydrofluorocarbons (HFCs) are now widely accepted alternative refrigerants to ozone-depleting chlorofluorocarbons (CFCs). Partially fluorinated alkanes have the same desirable physical properties as CFCs (i.e. low boiling point and high latent heat of vaporization), but without the harmful environmental side effects. In addition, hydrofluoroalkanes are also good lubricants, water- and stain-repellant products, chemical agents, and industrial solvents. The HFCs of particular industrial interest are fluorinated methanes and ethanes. 1-1-1-2-Tetrafluoroethane (R134a) is one of the most commonly used, primarily in domestic refrigeration and automobile heating, ventilation and air conditioning systems. Therefore, its thermodynamic properties and microscopic structure attract much attention.1–3

In evaluating the performance of refrigeration cycles, knowledge of the vapour–liquid equilibrium is important. Traditionally, such data are usually obtained from experiment or from empirical equations of state. These conventional methods can give accurate vapour–liquid equilibrium properties of pure substances or mixtures of components, but they cannot provide insight into the microscopic nature of the systems.

Computer simulation based on molecular modelling can predict vapour–liquid equilibria of fluids under a wide range of conditions, and also provides insight into their microscopic structure. Simulation can also model conditions under which experimental data are difficult or impossible to obtain. Several techniques have been proposed, such as the NPT + test particle method,4 Gibbs ensemble Monte Carlo (GEMC),5 Gibbs–Duhem integration,6 and histogram reweighting grand canonical ensemble Monte Carlo.7 Intermolecular potentials play a central role in obtaining accurate predictions of properties and microscopic structures. Molecular potential parameters are often optimized using experimental data for pure systems. The potential parameters can also be determined purely from ab initio calculations, or from a combination of ab initio calculations and experimental data.

There are several potentials for R134a. Lisal et al.8,9 proposed a set of effective pair potentials for R134a and pentafluoroethane using the Halgren’s Buff 14-7 and coulombic functions. In these potentials, only the internal rotation about the C–C bond is taken into account; bond lengths and bond angles are kept rigid. The potential functions are adjusted to reproduce the thermodynamic properties of the saturated liquid. Based on ab initio calculations (only for geometry optimization and atomic partial charges), Fermeglia et al.10 developed an all-atom potential model for R134a, which uses the 9-6 Lennard-Jones potential to represent the dispersion and induction energy. A dipolar two-centre Lennard-Jones model was also used to describe the vapour–liquid equilibrium of pure R134a and pentafluoroethane and their mixture.11–13 Peguin et al.14 developed an all-atom force field using a Lennard-Jones 12-6 function to describe the repulsion and dispersion energy and point charges to describe the coulombic energy. This new force field accurately predicts the bulk and interfacial properties of R134a. It reproduces the experimental vapour–liquid equilibrium properties especially well. Since the 12-6 Lennard-Jones potential is the most commonly used potential in computer simulations, this force field is straightforward to combine with established potentials, such as SPC/E for water15 and EPM for carbon dioxide (CO₂),16–17 to study these interesting mixtures.

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† Electronic supplementary information (ESI) available: (A) the potential parameters for R134a, (B) a description of the growing procedure for R134a, (C) the vapour-liquid equilibrium properties of R134a, and (D) an analysis of the orientation of pair interactions in the first solvation shell. See DOI: 10.1039/c0cp00620c
Besides the thermodynamic properties, the liquid structure of real fluids in general and of R134a in particular is also important from a theoretical point of view. There has been much effort devoted to characterizing the microscopic structure of many substances, including CO2,23–25 methylene chloride,26 and difluoromethane,27 and mixtures, e.g., CO2 + methane22,23 and CO2 + difluoromethane.23 However, there has not yet been a similar analysis of R134a, partly due to the lack of accurate potentials. The recent R134a potential of Peguin et al. gives us the opportunity to investigate the microscopic structure of this substance. Due to the large electronegativity of the fluorine atom, it is interesting to investigate the possible formation of C–H···F type hydrogen bonding in liquid R134a. In fact, this has been studied for difluoromethane, but no evidence for such hydrogen bonding was found.23 Our study of a larger molecule (R134a) will augment the previous findings in this regard. Beside the possibility of weak C–H···F hydrogen bonding, other factors can also play an important role in determining the liquid structure of R134a. R134a has a relatively high dipole moment of 2.06 D. Thus, strong hydrogen bonding was found.25 Our study of a larger molecule (R134a) will augment the previous findings in this regard. Beside the possibility of weak C–H···F hydrogen bonding, other factors can also play an important role in determining the liquid structure of R134a. R134a has a relatively high dipole moment of 2.06 D. Thus, strong hydrogen bonding was found.25 Our study of a larger molecule (R134a) will augment the previous findings in this regard. Beside the possibility of weak C–H···F hydrogen bonding, other factors can also play an important role in determining the liquid structure of R134a. R134a has a relatively high dipole moment of 2.06 D. 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simulation results. Due to the finite size effect, GEMC cannot simulate the phase equilibrium properties of any system at the vicinity of the critical point ($T_c = 373.9$ K). This is a well-known problem and has been addressed elsewhere. The simulations slightly overestimate the saturated vapour pressure of R134a (Fig. 1b). However, the deviations are quite small and still within the range of the statistical uncertainties of the simulations (shown in parentheses in Table S3, ESI†).

The latent heat of vaporization (Fig. 1c) is satisfactorily predicted at high temperatures, but slightly overestimated at low temperatures.

3.2 Microscopic structure of R134a

Our main goal here is to investigate the liquid structure of R134a. We take the simulated data of saturated liquid R134a at 260 K as an example to probe its intermolecular structure. Two established methods, which are often used to analyze the microscopic structure of substances, are employed. The radial distribution functions (RDFs) probe the occurrence of an atom at a distance from another atom. Peaks in an RDF are associated with solvation shells or specific neighbours. The orientational distribution function (ODF) gives information about the distribution of average angles between a pair of molecules at a given separation. In addition, we also analyze pair interaction energies.

To begin, we establish the distribution of the conformation of R134a molecules under these conditions, as the CF$_3$ and the CH$_2$F groups can freely rotate about the CC bond. The ensemble of configurations collected in the course of the simulations is analyzed. Fig. 2a shows the distribution of the average FCCF dihedral angles at 260 K. There is a sharp peak at 180° ($\cos(\phi) = -1$) and a lower one at about 60°, i.e., the FCCF dihedral angles of R134a have a preference for both 60° and 180°. Snapshots of the conformation of R134a are shown in Fig. 2b. This conformation is similar to ethane and is the most stable, as all six CF and CH bonds are furthest from each other, minimizing steric interactions.

In order to investigate the relative importance of the different types of interactions, we calculated separate contributions of...
the Lennard-Jones and electrostatic terms ($U_{LJ}$, $U_C$) to the total intermolecular interaction energy at different temperatures (Table 1). The electrostatic energy accounts for about 30% of the total, which suggests that dipole interaction is not dominant. Instead, liquid R134a seems to be dominated by dispersion energy, as the Lennard-Jones interaction accounts for about 70% of the total intermolecular interaction energy.

The pair interaction energy distribution is computed from the ensemble of individual molecule–molecule interactions. We have determined the distribution of the $U_{ij}$ pair interaction energies for the saturated liquid at three different temperatures, 220 K, 260 K and 340 K (Fig. 3). All curves have a shoulder at about –2 kJ mol$^{-1}$, on the attractive side of the main peak. This shoulder is a sign of the association of the molecules, which could arise from weak C–H···F hydrogen bonding or simply dipolar association.21 However, the RDFs of the H–F interactions show no evidence of specific interaction between hydrogen and fluorine, which suggests that C–H···F hydrogen bonding cannot be the dominant form of the molecular association. Similar shoulders also appear on the probability distribution of pair energies of difluoromethane, but no dipolar association.21 However, the RDFs of the H–F distribution of pair energies of difluoromethane, but no association. Similar shoulders also appear on the probability bonding cannot be the dominant form of the molecular about $1.4$, which is similar to that of liquid argon. These ratios also are found for other small molecules, such as methane, CO$_2$, and difluoromethane.21,22 The coordination number is about $12$, which is similar to that of liquid argon. These ratios up to their first minimum also yields a coordination number of about $12$. To have a clearer picture of the arrangement of pairs of atoms in the system, we calculate the RDFs, considering all interaction sites (Fig. 6). In the first solvation shell, the CC interactions are the most structured, as indicated by the

### Table 1 Intermolecular energy of liquid R134a

<table>
<thead>
<tr>
<th>$T$/K</th>
<th>$U_L$/kJ mol$^{-1}$</th>
<th>$U_{LJ}$/kJ mol$^{-1}$</th>
<th>$U_C$/kJ mol$^{-1}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>220</td>
<td>$-22.99$ (0.11)</td>
<td>$-16.13$ (0.10)</td>
<td>$-6.86$ (0.09)</td>
</tr>
<tr>
<td>240</td>
<td>$-21.66$ (0.13)</td>
<td>$-14.90$ (0.10)</td>
<td>$-6.76$ (0.09)</td>
</tr>
<tr>
<td>260</td>
<td>$-20.33$ (0.17)</td>
<td>$-14.61$ (0.11)</td>
<td>$-5.72$ (0.08)</td>
</tr>
<tr>
<td>280</td>
<td>$-18.96$ (0.27)</td>
<td>$-14.06$ (0.09)</td>
<td>$-4.90$ (0.08)</td>
</tr>
<tr>
<td>300</td>
<td>$-17.37$ (0.13)</td>
<td>$-12.54$ (0.13)</td>
<td>$-4.83$ (0.10)</td>
</tr>
<tr>
<td>320</td>
<td>$-15.97$ (0.19)</td>
<td>$-11.96$ (0.11)</td>
<td>$-4.01$ (0.17)</td>
</tr>
<tr>
<td>340</td>
<td>$-14.94$ (0.18)</td>
<td>$-10.46$ (0.20)</td>
<td>$-4.48$ (0.19)</td>
</tr>
</tbody>
</table>

$T$: temperature, $U_L$: configurational intermolecular energy in the liquid phase, $U_{LJ}$: configurational intermolecular energy due to Lennard-Jones interaction in the liquid phase, $U_C$: configurational intermolecular energy due to electrostatic interaction in the liquid phase. The numbers in parentheses indicate the statistical uncertainties.

Fig. 3 Probability distribution of the pair interaction energy of R134a molecules at 220 K (solid line), 260 K (dashed line) and 340 K (dotted line).

C–H···F hydrogen bonding is found either.21 Analogous observations have been made for formic acid,32 as a result of C–H···O hydrogen bonding, and for liquid acetone33 and acetonitrile,34 due to the dipolar association of the molecules. Fig. 4 shows the lowest pair energy encountered in the simulation. The angle between the dipole moment of the two molecules is $104^\circ$, indicating that the dipole–dipole interaction does not play an important role in the determination of this dimer arrangement.21

Fig. 5 shows the RDF of R134a molecules, considering one site per molecule at 260 K. For convenience, we identify the centre of the molecule, which we refer to as CoCC, as the midpoint of the CC bond rather than the molecular centre of mass. The RDF exhibits all the features of a closely packed system, given the size of the molecule. There are two peaks, at CoCC–CoCC distances of about 5 Å and about 9.5 Å. The ratio of the positions of the second and first peak is about 1.9, and the ratio of the first minimum and the first peak position is about 1.4, which is similar to that of liquid argon. These ratios are also found for other small molecules, such as methane, CO$_2$, and difluoromethane.21,22 The coordination number is about $12$, which is similar to that of liquid argon. These ratios up to their first minimum also yields a coordination number of about $12$. To have a clearer picture of the arrangement of pairs of atoms in the system, we calculate the RDFs, considering all interaction sites (Fig. 6). In the first solvation shell, the CC interactions are the most structured, as indicated by the

Fig. 4 Configuration of the R134a dimer with the lowest interaction energy ($-6.518$ kJ mol$^{-1}$) taken from simulation at 260 K.

Fig. 5 Radial distribution function of saturated liquid R134a, considering one site (centre of the CC bond) per molecule at 260 K.
narrow and sharp peaks ($g_{C2C2}$, $g_{C2C3}$, $g_{C3C3}$). This shows that carbon atoms aggregate together at about 5 Å. In contrast, the other interactions are not very structured; their peaks are low and broad.

Augmenting the RDFs, we use the ODFs to characterise the environments around each molecule in the liquid. Fig. 7a shows the distribution of the angle between $C2C3$ vectors and the intermolecular vectors $CoCC$–$CoCC$ of pairs of R134a molecules at 260 K. This distribution tells us how R134a molecules are positioned around a reference molecule in the liquid. In this figure, the vertical axis represents the distance between the centres ($CoCC$s) of pairs of R134a molecules. There are three maxima, which correspond to the first solvation shell. Two are located at a $CoCC$–$CoCC$ distance of about 5.2 Å and angles of 0° and 180°, with a coordination number of two for each peak. A larger peak appears at about 5.0 Å and an angle of 90°, with a coordination number of six. Thus, R134a molecules preferentially populate a circle of radius 5.0 Å around the midpoint of the CC bond of the central molecule. They also prefer to sit above and below the central molecule (Fig. 7b).

In order to investigate further the orientations of the neighbouring molecules, we divided the space around a reference molecule into four spaces: $p_F$, $p_H$, $p_T$ and $p_B$ (Fig. 8). A plane that cuts through $C2C3$ and is perpendicular to the $C2C3F6$ plane is used to separate the spaces $p_F$ and $p_H$. One would expect that the orientations of the molecules in the four spaces would be different. Therefore, we analyze the preferred orientations of the neighbouring molecules in all spaces separately. In the space $p_T$, the $C2C3$ vectors of the nearest neighbours in the first solvation shell have a preference for a perpendicular alignment (Fig. S2a, ESI†), while dipole vectors in this space have a preference for a parallel alignment, with some variation up to 30° with each other (Fig. S2b, ESI†).

Like the dipoles, the $C3F6$ bonds also prefer a parallel alignment, but with greater variation (Fig. S2c, ESI†). Combining the information from these three figures, we can deduce the preferred orientations of the neighbouring molecules in the $p_T$ space of the first solvation shell, as shown in Fig. S2d ESI†. In the space $p_H$ (Fig. S3, ESI†), the distribution of the angle between $C2C3$ vectors is the same as that in the $p_T$ space. However, at short distances, the $C2C3$ vectors have a preference for antiparallel alignment, but this alignment disappears quickly as the distance between molecules increases. We do not observe this alignment of the angle between $C2C3$ vectors in the $p_F$ space. Both the dipole vectors and the $C3F6$ bonds in this space prefer a parallel alignment. Unlike in the spaces $p_T$ and $p_H$, the $C2C3$ vectors of the nearest neighbouring molecules prefer a parallel alignment in the $p_F$ space (Fig. S4a, ESI†). The distribution of the average angle between dipoles (Fig. S4b, ESI†) also reveals that the dipoles prefer to be parallel to each other and so do the $C3F6$ bonds (Fig. S4c, ESI†). A similar orientation is preferred by the molecule that sits at the bottom of the central molecule in the first solvation shell (Fig. S5, ESI†). A snapshot of the first solvation shell of saturated liquid R134a at 260 K is shown in Fig. 9.

4. Conclusion

We have used coupled-decoupled CBMC simulations in the Gibbs ensemble to study the microscopic structure of saturated liquid R134a. We also calculated the vapour–liquid equilibrium properties for R134a. The potential employed has been recently published, where it was used in grand canonical MC (GCMC) simulations to determine vapour–liquid equilibrium properties. Both GCMC and GEMC methods yield good agreement with experiment, which implies that the methods...
are effective and the intermolecular potential is relatively accurate. Knowledge of vapour–liquid equilibrium is important in evaluating the performance of refrigeration cycles, because the two main processes in refrigeration cycles are the condensation and the evaporation of the refrigerants. The simulations of this system will augment empirical and experimental data in this regard.

Data for saturated liquid R134a at 260 K are taken from the GEMC simulations to analyze its microscopic structure. The RDFs reveal that the ratio of the positions of the second and first peaks, and the ratio of the first minimum and the first peak position are similar to that of liquid argon. The coordination number is about 12, which is similar to that of liquid methane, CO₂, and difluoromethane. The CC interactions are the most structured, whereas the other interactions are not very ordered. The ODFs reveal that the dipole moments of liquid R134a of the nearest neighbours in the first solvation shell have a preference for a parallel alignment. This is also characteristic of liquid CH₂F₂. The alignments of the neighbouring molecules in the π₅ and π₄ spaces around the central molecule are different. Particularly, in the π₅ space, the C₂C₃ bonds have a preference for anti-parallel alignment at short distance (4.7 Å), which is not observed in the π₅ space. The alignments of the neighbouring molecules above and below the central molecule are more or less the same. In liquid R134a, the electrostatic energy only accounts for about 30% of the total intermolecular interaction energy, whereas the Lennard-Jones interaction accounts for about 70%. The association of the molecules is found from the pair interaction energy analysis, but there is no evidence of weak C–H–F hydrogen bonding.

As computational power continues to improve, we expect to see more *ab initio* force fields in the future. This will allow us to obtain the thermodynamic properties from pure theory, and also provide a more comprehensive understanding of the molecular interactions at a microscopic scale. The mixing rule is a very convenient approximation for unlike interactions, but there is no guarantee that it works for every unlike interaction. Strictly speaking, we should calculate the potential parameters for every single interaction in the system, ideally from first
principles and use non-additive potentials. Work along these
lines is in progress. Nevertheless, the semi-empirical force
field employed in our work gives satisfactory agreement with
experiment for vapour–liquid equilibrium properties, and also
gives us the opportunity to analyze for the first time the micro-
scopic structure of R134a—an important industrial substance.

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