On the X-ray scattering pre-peak of linear mono-ols and the related micro-structure from computer simulations

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Abstract

The X-ray scattering intensities $I(k)$ of linear alkanols $\text{OH} (\text{CH}_2)_{n-1} \text{CH}_3$, obtained from experiments (methanol to 1-undecanol) and computer simulations (methanol to 1-nonanol) of different force field models, are comparatively studied, particularly in order to explain the origin and the properties of the scattering pre-peak in the $k$-vector range $0.3 \AA^{-1} - 1 \AA^{-1}$. The experimental $I(k)$ show two apparent features: the pre-peak position $k_P$ decreases with increasing $n$, and more intriguingly, the amplitude $A_P$ goes through a maximum at 1-butanol ($n = 4$). The first feature is well reproduced by all force field models, while the second shows a strong model dependence. The simulations reveal various shapes of clusters of the hydroxyl head-group, from $n > 2$. $k_P$ is directly related to the size of the meta-objects corresponding to such clusters surrounded by their alkyl tails. The explanation of the $A_P$ turnover at $n = 4$ is more involved. The analysis of the atom-atom structure factors indicates that the pre-peaks arise from an incomplete cancellation between two types of contributions from the various atom-atom structure factors: positive contributions from atoms part of the head group (essentially hydroxyl group), and a mostly...
negative contributions from the cross head/tail correlations. The former dominate for small alkanols \( n < 3 \), while the latter gain importance from \( n = 4 \), which explains the pre-peak amplitude turnover. The flexibility of the alkyl tails tend to reduce the cross contributions, thus revealing the crucial importance of this parameter in the models. Force fields with all-atom representation are less successful in reproducing the pre-peak features for smaller alkanols \( n < 6 \), possibly because they blur the charge ordering process since all atoms bear partial charges. The analysis clearly shows that it is not possible to obtain a model free explanation of the features of \( I(k) \)

1 Introduction

Radiation scattering is generally considered to provide the most direct insight into the spatial microscopic structure of liquids, complementary to thermodynamic or spectroscopy techniques. Linear mono-ols have been investigated by X-ray and neutron scattering techniques since the 1930s, and the most remarkable known feature is the presence of a pre-peak of the scattering intensity \( I(k) \), in the \( k \)-vector region \( 0.3-1 \text{Å}^{-1} \) [1, 2], in addition to the usual main peak around \( k \approx 1.4-2 \text{Å}^{-1} \). The main peak can be interpreted through \( \sigma = 2\pi/k \) as corresponding to the mean atomic size \( \sigma \approx 3-4 \text{Å} \), such as in the X-ray or neutron diffusion in simple atomic liquids [3]. There has been several investigations of the origin of the pre-peak, which is now generally interpreted as corresponding to the existence of short chain-like or ring-like clusters of hydrogen bonded hydroxyl groups. Interestingly, this conclusion was reached by several different routes: the direct reconstruction of the pre-peak as a diffraction pattern from assumed cluster shapes [4, 5], the thermodynamic route of matching the internal energy with H-bond associations [6], a hard sphere model for the spherical cluster aggregates [7], and more recently various computer simulation analyses [8, 9, 10, 11, 12, 13, 14, 15, 16]. All these descriptions emphasize a causal link between the existence of clusters formed by the hydroxyl groups and the pre-peak in \( I(k) \). This is directly inspired from the fact that radiation is scattered off objects, usually atoms or molecules, hence specific features in \( I(k) \), such as the pre-peak, should refer to the existence of corresponding meta-objects, such as clusters, aggregates or self-assembled structures [17]. These descriptions also pose the question as to whether or not it is possible to explain the features of \( I(k) \) in a model free approach.

In the present work, we wish to emphasize that it is not so much the meta-objects, but the density correlations associated with the atomic constituents of such objects, which help explain the details in \( I(k) \). In particular, the atom-atom correlation functions reveal differences in the atomic ordering, depending on the head group atoms (mostly hydroxyl atoms), and the alkyl tail atoms. These differences are explained in terms of force field representations by the fact that head group atoms are charged, while alkyl tail atoms are overall neutral (or very weakly charged). Hence, the head group atoms tend to cluster through the Coulomb interaction, mostly into short chain-like aggregates surrounded by the
tail atoms. The corresponding micro-structure affects the shape of the atom-atom distribution functions, which in turn allow to infer the local ordering of the various types of atom groups. Such correlation functions, which depend strongly in the choice of the force field models, can only be obtained from computer simulations, enforcing the idea of an unavoidable model based understanding of the features of $I(k)$.

This interpretation is directly supported by the very definition of the scattering intensity \cite{18, 19} as related to the statistical thermal or ensemble average

$$I(k) \propto \langle \sum_{ij} \exp(ik \cdot (\vec{r}_i - \vec{r}_j)) \rangle$$

where the sum runs over all pairs of atomic sites $i,j$. Under this form, it is not possible to recognize how atomic sites organise themselves into clusters or aggregates, both inside a molecule and across different molecules. However, the r.h.s. of the above equation can be written in terms of atom-atom structure factor, and the Debye formula \cite{18, 19} allows to rewrite Eq.(1) as

$$I(k) = r_0^2 \rho \sum_{ij} f_i(k)f_j(k)S^{(T)}_{ij}(k)$$

where $S^{(T)}_{ij}(k)$ contains the intra-molecular atom-atom structure factor $w_{ij}(k)$ (to be discussed in Section 2.2), and the Fourier transform of the atom-atom intermolecular pair correlation function $g_{ij}(r)$

$$S^{(T)}_{ij}(k) = w_{ij}(k) + \rho \int d\vec{r} [g_{ij}(r) - 1] \exp(ik \cdot \vec{r})$$

$\rho = N/V$ is the density (where $N$ is the number of molecules in the volume $V$), the $f_i(k)$ functions are the form factor of atom $i$, and $r_0 = 2.8179 \cdot 10^{-13}\text{cm}$ is the electronic radius. Eq.(2) shows that $I(k)$ is related to the density correlations of the liquid through the pair correlation functions $g_{ij}(r)$. Since these quantities describe local atomic and molecular ordering, any specific feature in $I(k)$ is necessarily related to this local order, as expressed through the atom-atom structure factors. Therefore, it is of primary interest to analyse first these structure factors, in order to better understand the origin of the pre-peak. This is the route we use herein.

In the present report, we compare the experimental X-ray scattering intensities $I(k)$ for several mono-ols, with computer simulation results for several force fields. Similar comparison have been reported by several authors \cite{20, 21, 22, 23, 12, 16}, with the generic aim of relating the pre-peak to the clustering of the hydroxyl groups. In the present work, we show that the alkyl tails also play a crucial role in the interpretation of the pre-peak. This is due to the fact that the charge ordering of the head groups are influenced by the ordering of the alkyl tail groups, as witnessed by the micro-segregation of the charged and uncharged groups observed in computer simulations.

Charge ordering has been recently studied in the context of room temperature ionic liquids \cite{24, 25, 26}, where the polar/apolar segregation of charged and
neutral atomic groups [27] provides a contextual link with the present developments. In such works, the scattering pre-peak is related to the nano-segregation of the charged and neutral groups, described as polar/apolar in the corresponding literature. The principal difference with the present system is the fact that in ionic systems the charges are free to move, whereas in alcohol molecules they are constrained both to molecular neutrality and to be attached to the alkyl tails. We come back to this difference in the Discussion section.

From the remarks above, it becomes very clear that the existence of a pre-peak in X-ray scattering $I(k)$ cannot be analysed without detailed molecular simulations, involving in particular approximate force field models. Therefore, it would be crucial to have a reliable model capable of reproducing the details of the shape of $I(k)$ for various mono-ols. The present study reveals that not all models are able to reproduce the pre-peak behaviour correctly. Hence this comparison provides a severe selection principle. In particular, the analysis reveals that the ability of a force field model to reproduce (or not) the pre-peak in $I(k)$, is less related to a proper account of the charge and domain ordering, than the ability to properly describe the incomplete canceling of these two forms of local order.

The remainder of this paper is as follows. In the next section we describe the X-ray experiments, simulation models and protocols, as well as the theoretical developments required to analyse the atom-atom correlation contributions to $I(k)$. The Results section shows detailed analysis of the X-ray intensities obtained from scattering experiments and molecular dynamics calculations. A discussion and conclusion sections close the presentation of this work.

2 Experimental and model simulation details

2.1 Experimental

The X-ray diffraction (XRD) experiments were performed at the beamlines BL2 and BL9 of the DELTA synchrotron radiation source using the setup for wide angle X-ray scattering [28] with an incident X-ray energy of 11keV and 13keV, respectively. The X-rays were monochromatized using a multi-layer monochromator at BL2 and a Si(311) double crystal monochromator at BL9 with a beamsize of 0.5x0.5 mm$^2$ and 1x1 mm$^2$ (vxh) at the sample position, respectively. The scattered photons were detected by a MAR345 image plate detector. The calibration of the two-dimensional diffraction patterns was performed with silicon and lanthanum hexaboride references. We measured the linear mono-ols methanol (purity $\geq 99.9\%$), ethanol ($\geq 99.9\%$), 1-propanol ($\geq 99.9\%$), 1-butanol ($\geq 99.8\%$), 1-pentanol ($\geq 99.8\%$), 1-hexanol ($\geq 99\%$), 1-heptanol ($\geq 99.9\%$), 1-octanol ($\geq 99\%$), 1-nonanol ($\geq 98\%$), 1-decanol ($\geq 99\%$) and 1-undecanol ($\geq 97.5\%$). All samples were purchased by Sigma Aldrich, except 1-octanol that was bought from Alfa Aesar, and used without further treatment. The linear mono-ols were filled into borosilicate capillaries with 3.5 mm (BL2) and 2 mm (BL9) diameter and measured at a temperature of about 293 K.
The XRD images were integrated with the program package Fit2D \cite{29} and converted to wave-vector transfer $k$ scale. The diffraction pattern were corrected for the scattering contributions of the capillaries as well as of air and were normalized to the mean integral of the calculated diffraction patterns in the $k$-range of 0.2 to 2.3 Å$^{-1}$.

### 2.2 Models

We have used several models for the linear mono-ols, previously studied in the literature. Namely, we considered the OPLS (Optimized Potentials for Liquid Simulations) \cite{30, 31}, TraPPE (Transferable Potentials for Phase Equilibria) \cite{32}, CHARMM (Chemistry at Harvard Macromolecular Mechanics) \cite{33, 34, 35}, and to some extent the GROMOS \cite{36, 37} force field models. All models, except for CHARMM are used in their united atom (UA) versions for the methyl/methylene groups. Indeed, CHARMM is an all-atom (AA) model by construction. The OPLS AA model for methanol and ethanol have been tested, in order to have an equivalent comparison with CHARMM. All force field models are based on the atom-atom interactions modeled as a Lennard-Jones centre with a partial charge. Since our interpretation of the pre-peak feature is based on local charge order, we have listed in Tables 1-5 in the ESI, the typical Lennard-Jones energy $\epsilon_i$ and diameter $\sigma_i$, and Coulomb charge $e_i$ for each of the atoms $i$, namely the hydrogen $H$, oxygen $O$ and first carbon group $C_1$, which are charged, and the remaining carbons numbered as $C_i$, for $i = 2, n$ where $n$ is the terminal carbon group. For the CHARMM model, the partial charges of the hydrogen and the carbon of the carbon groups are explicited. We note that all models are flexible, hence bond length and dihedral dynamics are considered consistently with the corresponding parameters in the various force fields. We will report elsewhere detailed analysis of the differences between the models concerning the contribution of the local structure on $I(k)$.

We retain from this section that all UA models attribute a negative charge for the oxygen atom and positive one for the hydrogen atom, followed by another positively charged first methylene atom, and remaining methyl groups are uncharged. From this observation, we can already predict that the 3 head group atoms will tend to cluster, leaving the uncharged tail atoms to randomly position around and constrained by the sole packing effects. If these two groups were separated, the resulting mixture will readily phase separate into a polar part and an apolar part. Since these two groups are bound into a single alcohol molecule, they produce a micro-structure which brings the full demixing to a local micro-segregation of the polar and apolar group. This is similar to what is observed ionic-liquids \cite{27}. The problem posed by this work is to figure out the link between this micro-structure and the pre-peak in $I(k)$.

We emphasize again that, unlike the OPLS and TraPPE UA models, the CHARMM model is an AA model, hence the alkyl tail carbons contain partial charges over the central atom and the surrounding hydrogen atoms. Hence, these alkyl tail groups will likewise tend to exhibit local charge ordering. Although the total charge of the carbon groups is zero for the remote alkyl groups,
there is nevertheless a Coulomb charge influence in the way these groups will tend to position with respect to each other and the hydroxyl groups. We will show below that, although appearing at first more realistic than the AA models, this model poses some problems such as the absence or suppression of scattering pre-peak for methanol and ethanol.

2.3 Simulations

All simulations were performed with the program package Gromacs [38]. We systematically used $N = 2048$ number of molecules for all systems, which corresponds to box sizes ranging from approximately $52 \, \text{Å}$, for methanol to $82 \, \text{Å}$ for 1-nonanol. For 1-octanol, simulations of $N = 8000$ molecules were performed for testing purpose, which produced correlation functions not distinguishable from $N = 2048$, indicating that this system size was sufficient even for longer alkanols.

The systems were simulated in the isobaric-isothermal (constant $NpT$) ensemble, at the temperature of $T = 300 \, \text{K}$ and pressure $p = 1 \, \text{bar}$. Those conditions were achieved with the Nose-Hoover thermostat [39] [40] and Parrinello-Rahman barostat [41] [42]. It is worth emphasizing this procedure allows to test the models under the same conditions as the experimental ones. As a consequence, the calculated densities may differ from the experimental ones. We do not enforce the experimental densities using the isochoric constant $NVT$ ensemble, since it would create a bias in the model analysis. In particular, the correlation functions and the structure factors differ according to whether these are calculated in the $NpT$ or $NVT$ ensemble, because of the density differences. This is important to consider when comparing calculated intensities $I(k)$ obtained from these different methods.

We followed the same procedure for every simulation. Packmol [43] was used to obtain the initial configurations of the systems from the pdb files of each molecule. After energy minimization, the systems were equilibrated in the $NVT$ and then $NpT$ ensemble, for a total of 1 ns each. The following production runs lasted 5 ns, in order to sample at least 2000 configurations for calculating the site-site correlation functions $g_{ij}(r)$. In many cases, several independent series of such 5 ns runs were conducted, in order to ensure convergence of the $g_{ij}(r)$.

2.4 Theoretical details

2.4.1 The intra-molecular correlations

The evaluation of the total structure factor in Eq. (3) requires that of the intra-molecular term $w_{ij}(k)$. In previous works, we had approximated this term by its rigid molecule form [44] $w_{ij}(k) = j_0(kd_{ij})$, where $j_0(x)$ is the zeroth order spherical Bessel function, and $d_{ij} = |\vec{r}_i - \vec{r}_j|$ is the inter-atomic distance between atoms $i$ and $j$ on the same molecule. While this approximation might be acceptable for smaller molecules, such as ethanol for example (see Fig.1a below), this
is no longer true for longer molecules such as 1-octanol, for example, for which the flexibility leads to mean inter-atomic distance to be very different from the rigid molecule values. To account for this, we have evaluated the intra-molecular correlation functions \( w_{ij}(r) \) directly from the configurations obtained from Gromacs, by using the same standard neighbour histogram method employed for the evaluation of the inter-molecular correlation \( g_{ij}(r) \). The \( w_{ij}(k) \) were then evaluated by the same Fast Fourier Technique (FFT) technique used to obtain the \( S_{ij}(k) \) from the \( g_{ij}(r) \) \[9\].

In Fig.1 we show for the OPLS force field model the differences between the rigid molecule version of the \( w_{ij}(k) \) (in dashed lines) and the flexible version (in full lines), for specific atom pairs. Namely, we focus on 3 alcohols, ethanol, 1-pentanol and 1-octanol, for which we show \( w_{OC_i}(k) \) between the oxygen atom \( O \) and all the carbon atoms \( C_i \) (\( i \) ranging from 1 to the terminal one, which is 2 for ethanol, 5 for 1-pentanol and 8 for 1-octanol), as well as \( w_{C_1C_j}(k) \), for the correlations between the first carbon atom and all remaining others. The choice of these atoms allows to test the flexibility as seen from the oxygen atom to all carbon groups, as well as that of the carbon atoms between themselves. In all 3 plots, the gray vertical line marks the pre-peak position in the corresponding OPLS \( I(k) \) (see Fig.4 in the next section).

![Figure 1: \( w_{XC_1}(k) \) for ethanol, 1-pentanol and 1-octanol, where \( X=O \) (upper panels) or \( X=C_1 \) (lower panels), and \( i = 1, 2, \ldots n \). The value of \( i \) is shown for 1-pentanol and 1-octanol, next to the corresponding curves. Full curve is with flexibility of alkyl tail and dashed curve without. See text for more details.](image-url)

Fig.1 a) shows that, for the ethanol molecule, the flexibility does not affect much the intra-molecular correlations, since the full and dashed lines are mostly superposed, particularly around the pre-peak region \( k_P \approx 0.7\,\text{Å}^{-1} \). For the case of 1-pentanol (middle panel), this is no longer true for the oxygen atom and the alkyl tail end carbon atoms, starting from \( C_3 \) (upper panel). This difference is particularly important around the pre-peak position \( k_P \approx 0.51\,\text{Å}^{-1} \). However, the carbon atoms are not much affected by the flexibility issue. In particular, we observe that the mean distance between \( C_3 \) and the neighbouring carbons...
is nearly the same: \(d_{C_3C_1} \approx d_{C_3C_5}\) and \(d_{C_3C_2} \approx d_{C_3C_4}\). These relations remain true for the case of 1-octanol (right panel). But we also see that the flexibility issue is aggravated when considering atoms that are much further apart, as one would indeed expect from a fully flexible tail. An important feature seen in both panel b) and c) is that the intra-molecular correlations for the flexible case are above that of the rigid case. This feature is crucial in determining the sign of the pre-peak in \(I(k)\), as we will see in the results section 3.6.

2.4.2 Correlation functions and the computation of \(I(k)\)

The correlation functions obtained in our simulations are quite smooth, even at the long range part. In some cases, it was necessary to do additional production runs, in order to ensure that the \(g_{ij}(r)\) were well defined in the long range part. The inter-molecular structure factors were evaluated by FFT techniques from the \(g_{ij}(r)\) (the second term at the right-hand side of Eq.(3)), as in all our previous works \[45, 46, 47\]. Since the box size \(L\) is finite, all our \(S_{ij}(k)\) are not reliable for \(k_{\text{min}} < 2\pi/L\). This is typically \(k_{\text{min}} \approx 0.115 \text{Å}^{-1}\) for methanol, and \(k_{\text{min}} \approx 0.072 \text{Å}^{-1}\) for 1-nonanol. These values are much smaller than the respective pre-peak positions \(k \approx 1 \text{Å}^{-1}\) and \(k_p \approx 0.4 \text{Å}^{-1}\), ensuring that the box size does not affect the physical features we discuss herein.

The numerical evaluation of \(I(k)\) in experimental units of \(\text{cm}^{-1}\) has been described in \[48\]. We remind that this requires to replace the frontal \(r_0^2\rho\) term of \(I(k)\) in Eq.(1) by the following pre-factor \(r_0^2(N/L^3)/100\), \(N\) is the number of molecules in the simulation box of size \(L\) (expressed in meters).

3 Results

3.1 Experiments

The experimental \(I(k)\) are shown in Fig.2 for the measurements performed at BL2 of DELTA and are in reasonable agreement with the data published by Vahvaselkä et al. \[10\] and Tomšič et al. \[23\] [16] (see Fig.1 in SI).

Despite the different experimental setup used at BL9 of DELTA, the data taken at both experiments resemble each other. Complementary measurements were performed on a D8 laboratory diffractometer with less statistical accuracy but are also consistent with the ones presented here. The patterns exhibit two distinct peaks, the main diffraction peak and the well known pre-peak feature in mono-ols \[50, 51, 49, 52, 23, 53, 54, 12, 55\]. The latter can be observed in the \(k\)-range \(0.3 \text{Å}^{-1} < k < 1 \text{Å}^{-1}\) and is well separated from the main diffraction maximum (\(1.4 \text{Å}^{-1} < k < 1.75 \text{Å}^{-1}\)). Interestingly, the pre-peak is a simple shoulder for methanol. The main peak position and amplitude varies considerably from methanol to 1-propanol, above which it stabilises in position while increasing in amplitude. As for the pre-peak, two important features are observed. The first concerns the systematic decrease of the pre-peak position with increasing alcohol length. A second feature is the fact that
Figure 2: Experimental X-ray scattered intensities for alcohols from methanol to 1-nonanol. The inset shows the pre-peaks, as extracted from a procedure explained in Section 3.3.

The pre-peak amplitude increases with \( n \) for small alkanols, until a maximum for 1-butanol, after which it decreases systematically. This behaviour suggests a crossover phenomena, which we will study in the following sections. Both observations are line with a previous similar report of these trends [56], and require an explanation.

3.2 Simulations

The results for \( I(k) \) from computer simulations, and for various models are shown in Fig.3, and are compared with the experimental ones as shown in Fig.2. The results are presented for the OPLS model in blue, for the TraPPE model in red, for the CHARMM model in green and the GROMOS model in gold, while the experiment is plotted as black dashed lines. The gray area represents the small-\( k \) region for which the finite size errors from the simulations affect the estimations of the calculated \( I(k) \).

It is immediately seen that, for all models, the overall shapes of the calculated \( I(k) \) for various mono-ols are in agreement with the experimental ones, including in particular the main peak in the range 1.3 \( \text{Å}^{-1} < k_M < 1.7 \text{Å}^{-1} \), and the pre-peak in the range 0.3 \( \text{Å}^{-1} < k_P < 1.0 \text{Å}^{-1} \). The OPLS model seems more consistent than the two others, in the sense that it reproduces both main peak and pre-peak features better than the other models. It generally overestimates main peak height and underestimates pre-peak height. The TraPPE model does not provide a good description of the main peak for longer mono-ols (amplitude and position), but produces a pre-peak consistent with experiments. The position of the pre-peak is in remarkable agreement with the experiment up to 1-propanol, then deviates significantly but approaches again the experimen-
Figure 3: $I(k)$ calculated with the OPLS (blue), TraPPE (red), CHARMM (green), GROMOS (gold) model from methanol to 1-nonanol compared to the experimental diffraction patterns (black dashed lines).

tal pre-peak position for the largest mono-ols. CHARMM model is the most inconsistent one, since it is unable to produce the pre-peak for lower alkanol (except for 1-propanol - but see below). Interestingly, this model predicts an unphysical negative pre-peak (anti-peak) for 1-pentanol. It predicts consistent pre-peaks for longer chains.

The good agreement at the main peak indicates that the force field model is able to describe short range spatial correlations relatively well. Most force fields are designed to achieve this to some extent. Indeed, most force fields give a good account of thermodynamic properties such as the enthalpy. This quantity is related to the elementary integrals $\int d\vec{r}g_{ij}(r)v_{ij}(r)$, where $v_{ij}(r)$ is the pair interaction modeled by the force field, for a given pair of atomic sites $i$ and $j$. Such interactions have a spatial range limited to first or second neighbours, even in presence of long range interactions, because of the Coulomb screening phenomena in dense liquids. Therefore, a good approximation of the enthalpy indicates that both the interaction $v_{ij}(r)$ and the resulting pair correlation $g_{ij}(r)$ must be well reproduced by the force field model.

A good agreement for the pre-peak is a more demanding feature, indicating that the force field is able to capture cluster/domain formation accurately. This
information is hidden in the long range part of the correlations, and is more sensitive to the microscopic details.

A systematic analysis of both main peak and pre-peak features is provided in the next sub-section.

### 3.3 Main peak and pre-peak analysis

For a detailed analysis we determined the main-peak using a combination of a Pearson VII function and a linear slope. Then in a second step we fitted the pre-peak likewise in the corresponding pre-peak region after subtraction of the main-peak fit and revealed their peak intensity, peak position and FWHM (Full Width at Half Maximum). The corresponding error bars were estimated analyzing the measurements carried out at the various experimental end-stations and are dominated by the background treatment. This procedure was likewise performed for the computed $I(k)$ for proper comparison between experiment and theory. However, using this procedure it was not possibly to extract a pre-peak for all the computed $I(k)$, i.e. for methanol, ethanol, 1-butanol and 1-pentanol obtained from CHARMM force field. The error bars of this analysis were obtained by variation of the corresponding fitting ranges. The results of the evaluation are shown in Fig. 4 for the main peak and Fig. 5 for the pre-peak.

![Figure 4: Main peak analysis of $I(k)$ from the experiment (black triangles), OPLS model (blue dots), CHARMM model (green squares) and TraPPE model (red diamonds). Peak position $k_M$ (left), peak amplitude $A_M$ (center) and FWHM (right).](image)

Fig.4 shows that the main peak amplitude $A_M$ increases with alcohol chain length showing an almost linear behavior for longer mono-ols starting with 1-propanol, while the position $k_M$ strongly decreases and then changes almost linearly with chain length. Surprisingly, the FWHM exhibits a clear non-linear dependency. This can be assigned to the proximity of the position of both pre-peak and main peak up to 1-butanol and the corresponding crosstalk in the fit. For larger chain lengths the difference in peak positions strongly increases.
providing a more independent fit of both features. However, this behavior can
be observed consistently in simulations (OPLS and CHARMM) and experiment.

These trends in the main peak can be rationalised as follows. The position
$k_M$ of the peak is related to the mean atomic diameter $<\sigma>$ for a given alcohol,
according to $k_M \approx \frac{2\pi}{<\sigma>}$. For a dense liquid, this is equivalent to $<\sigma>$
being related to the average particle-particle contact. For small alcohols, this
is dominated by the small diameter of hydrogen, hence a larger $k_M$. But for
the longer ones it would be dominated by the diameter of the methylene group,
hence a smaller $k_M$. Since, for longer alcohols the methylene groups dominates
the averaging, it explains why $k_M$ tends to saturate at the lower value of $1.4\text{Å}^{-1}$,
which corresponds to $\sigma \approx 4.5\text{Å}^{-1}$, which is close to $\sigma_{\text{CH}_2} \approx 4.3\text{Å}$. This is con-
sistent for OPLS and CHARMM, but not TraPPE, which indicates too a large
methylene diameter about 4.65Å. The amplitude $A_M$ is governed, according
to Eq.(2), by the main peak amplitude of the various structure factors $S_{ij}(k)$. Although the number of methylene-methylene correlations increases with $n^2$,
the density term $\rho = N/V$ in Eq.(2) moderates this trend, since simulations
indicate that, for a fixed number of molecules $N = 2048$, the box size $V = L^3$
increases with $n$.

Overall, the OPLS model appears to follow the overall experimental trends
better than the CHARMM and TraPPE models.

Fig.5 shows a similar analysis for the pre-peak.

Figure 5: Pre-peak analysis of $I(k)$ from the experiment (black triangles), OPLS
model (blue dots), CHARMM model (green squares) and TraPPE model (red
diamonds). Peak position $k_P$ (left), peak amplitude $A_P$ (center) and FWHM
(right).

The pre-peak position $k_M$ is seen to decrease monotonously with $n$. This
trend can be understood the following way. $k_M$ is related to the size of the
meta-object, but which? Is it the mean length of the hydroxyl head group linear
aggregate? The cluster analysis (see Fig.6 below) shows that the cluster size
is either dominated by pentamers for all mono-ols (OPLS, TraPPE) or changes
very little (CHARMM), hence in contradiction with the trend observed in Fig.
4. It seems more reasonable to associate the meta-object to the ensemble of the chain aggregate surrounded by the alkyl chain, whose mean size \(< d >\) would increase with \(n\), in agreement with the observed behaviour for \(k_P \approx 2\pi/\langle d \rangle\). The FWHM of the pre-peak decreases continuously from methanol to 1-undecanol. As for the amplitude \(A_P\), the experimental data show that it increases up to 1-butanol, saturates, and then decreases starting with 1-octanol. A similar observation for the increase of pre-peak was made by Tomšič et al. [16], in very good agreement with our findings. The decrease of the amplitude \(A_P\) is more complicated to rationalize. It is related to the importance of the alkyl tail contributions with increasing \(n\), and will be discussed later in the light of the simulations.

Overall by comparison between experiment and theory, the OPLS-UA model provides the most prominent agreement with the experimental findings.

3.4 Snapshots

Snapshots of typical alcohols are shown in Fig. 6, and for various models. Visual inspection confirms mostly the existence of short chain-like aggregates for all models and alcohols.

These snapshots are designed to highlight the hydroxyl groups, with oxygen in red and hydrogen in white, while the alkyl tails are shown as semi-transparent cyan. While the density of the H-bonded chains decreases with increasing alkyl chain length, which is obvious since the number of methyl groups increases, it clearly appears that the H-bonded chains seem longer for longer mono-ols. This feature can be rationalized by the fact that charged groups can cluster more easily in large groups when the alkyl chain is longer. The visual inspection of the CHARMM model is complicated by the fact that this is an AA model, hence the alkyl tails are fatter because of the prominent hydrogen atoms. This makes the cyan background somewhat denser and hinders the clear vision of hydroxyl chains, and gives a false impression of a lesser density of the hydroxyl groups. Despite this drawback, the somewhat lesser clustering tendencies of this model are visually apparent. A more detailed investigation reveals that all sorts of geometry and size of clusters exists, such as linear chains, branched chains, loops, branched loops, lassos. In that, clustering shape in all mono-ols is very similar to what we reported earlier for methanol [9], with the exception that larger clusters more prominently appear for longer mono-ols.

3.5 Cluster analysis

Fig. 7 shows the clustering probabilities of the oxygen atoms for selected mono-ols and models. The methodology used is the same as in our previous works [9, 57].

Both OPLS and TraPPE model show a systematic trend for higher clustering probability with increasing alcohol length, both with an average chain length around 5. This is somewhat consistent with what is observed in the snapshots of Fig. 6. In particular, for longer alcohols, the probability of observing pentamers
of hydroxyl groups exceeds that of free monomers. But, the most striking and unexpected feature is the near absence of a cluster peak for the CHARMM model. For this model, the monomer probability is the highest, suggesting that the number of pentamer hydroxyl groups is small. However, the total number of hydrogen-bonded molecules is still much larger than the number of monomers. In that, the CHARMM model is not different than other model, as far as the global clustering property is concerned. It confirms what the visual inspection of the snapshots in Fig.6 suggests. This problem may be related to the absence of a pre-peak for this model for many alcohols, as shown in Fig.3. This marked difference with the other models requires some clarifications, which the analysis of the correlation functions is likely to provide.
3.6 Correlation function analysis

Since atom-atom structure factors $S_{ij}(k)$ appear in the definition of $I(k)$ through Eq. (2), in this entire section we will analyse how the specific micro-structure of various types of the alcohols we have studied herein, contribute to the $S_{ij}(k)$. In the remainder, we choose to graphically represent the inter-molecular part of the structure factor with a convention similar to a single atom liquid [44], namely:

$$S_{ij}(k) = 1 + \rho \int d\vec{r} [g_{ij}(r) - 1] \exp(i\vec{k} \cdot \vec{r})$$ (4)

This way, aside the common additive term 1, we will be essentially comparing the Fourier transforms of the atom-atom inter-molecular correlation functions $g_{ij}(r)$.

3.6.1 Charge ordering and domain ordering

From our previous work [58] [59] [47], the key feature in the hydroxyl group chaining is the charge ordering between the negatively charged oxygen and positively charged hydrogen, leading to a characteristic linear chaining of the type $+-+-+-+---...$. As a consequence, the associated atoms having many of their like neighbours aligned, they have less second and third neighbours of their kind. This leads to two specific features [47] in the corresponding inter-molecular $g_{ii}(r)$ for the correlations between like atoms $i$: a high first peak, witnessing the strong association, followed by depleted correlations, due to lesser neighbours of like atoms. This feature is illustrated in Fig. 8 for the oxygen-oxygen correlation.
functions \( g_{OO}(r) \) of the OPLS model for selected 3 mono-ols (ethanol(green), 1-pentanol (blue) and 1-octanol (black)).

Figure 8: Oxygen-oxygen correlation functions \( g_{OO}(r) \) of the OPLS model for selected mono-ols (ethanol (green), 1-pentanol (blue) and 1-octanol (black)), illustrating the features which give rise to the structure factor pre-peak (see text). The inset shows the corresponding pre-peak in \( S_{OO}(k) \).

It is this double feature which leads to the pre-peak feature in the corresponding \( S_{OO}(k) \), as was demonstrated in Ref. [47]. The essence of the demonstration is the following and holds for any pair of associated atoms \( a \): the high first peak of \( g_{aa}(r) \) contributes to a wide positive peak at \( k = 0 \) for \( S_{aa}(k) \), while the depletion part of \( g_{aa}(r) \) contributes to a narrow negative peak at \( k = 0 \). The total contribution is a positive pre-peak in \( S_{aa}(k) \). This is illustrated in the inset of Fig.8 for \( S_{OO}(k) \).

Another feature which is fundamental to understand the pre-peak in \( I(k) \), is the existence of anti-correlations between the charged and uncharged groups [60, 48]. Indeed, since charged groups prefer to associate together, the uncharged groups tend to occupy the remaining empty space, with the permanent constraint that these groups are attached to the full molecules. The spatial alternation of charged and uncharged atomic groups is illustrated in Fig.9 for the OPLS model and 3 selected alkanols.

The charged groups (oxygen, hydrogen and first methylene atom) are shown in blue, while the remaining methylene/methyl atoms are shown in semi-transparent cyan. The charged groups form compact semi-linear blue clusters, separated from the surrounding cyan neutral atoms.

This alternation of groups/domains, leads to long range anti-correlations between them. This is illustrated in Fig.10 for oxygen-oxygen \( g_{OO}(r) \) and oxygen-terminal methyl group \( g_{OC_n}(r) \) correlation functions, for the OPLS models of the 3 alkanols selected in the previous snapshots in Fig.9.
Figure 9: Snapshots for 3 alcohols (ethanol, 1-pentanol and 1-octanol) for the OPLS model, illustrating the nano-segregation between the charged atoms (dark blue) and the uncharged ones (pale blue, shown in semi-transparent).

The long range phase opposition domain oscillations can be observed in all 3 cases, but these are broader for the longer mono-ols. The width of the oscillations period increases with the alkanol chain length. The inset shows the first peak details. The influence of the Coulomb induced large first peak is clearly seen in the $g_{OO}(r)$, which is a direct consequence of the H-bonding correlations between the oxygen atoms.

Figure 10: Correlation functions for OPLS model $g_{OO}(r)$ (blue) and $g_{OC_n}(r)$ (black) illustrating the domain ordering between charged head group atoms (here O) and tail atoms (here the last carbon Cₙ), and for 3 different alcohols (ethanol, 1-pentanol and 1-octanol).

These anti-correlations give raise to negative pre-peaks (or anti-peaks) in the structure factors, as illustrated in Fig.11. The anti-peaks are easy to understand:
since the correlations $g_{OC_n}(r)$ are in opposing phase with those of the $g_{OO}(r)$, their contribution to the Fourier transform in Eq. (4) has necessarily the opposite sign, hence contribute in the opposite direction than that of the $S_{OO}(k)$ pre-peak.

![Structure factors](image)

Figure 11: Structure factors $S_{OO}(k)$ and $S_{OC_n}(k)$ corresponding to the correlation functions shown in Fig. 10.

While the positive pre-peaks are quite prominent, and greater than the main peaks, the anti-peaks are often negative. It is interesting to note that for the smaller mono-ols, the anti-peak is merely a small dip (top panel in Fig. 11). This is a direct consequence of the alkyl tails being very short, and being only weakly correlated with the charged head groups. However, for longer alcohols, these tails are strongly anti-correlated with the head groups. This anti-correlation reflects the nano-segregation observed in the snapshots Fig. 6, between the aggregated hydroxyl groups and the alkyl tails. Indeed, if the two types of groups, charged hydroxyl and neutral tail, were detached, one would observe a macroscopic phase transition between them. Since they are tied into the same molecule, one observes a depleted correlation between the two, as that seen in $g_{OC_n}(r)$ in Fig. 10, which is at the origin of the anti-correlation.

Now, we are in position to understand how these correlations and anti-correlations play a capital role in the pre-peak of $I(k)$, as a result of cancellations in Eq. (1). The lower panels of Fig. 12 show all the atom-atom structure factors $S_{ij}(k)$ (as defined in Eq. (2), that is with the extra 1 added for presentation purpose) for the same 3 OPLS-UA alkanols, highlighting the like and cross correlations in the pre-peak region. The upper panels show the $I(k)$ (in black), together with the like correlation contributions (in blue) and cross correlation contributions (in red).
Figure 12: Scattered intensities $I(k)$ (upper panels) and all corresponding atom-atom structure factors $S_{ij}(k)$ for the OPLS model of 3 alcohols (ethanol, 1-pentanol and 1-octanol). The upper panels equally show the like atom contributions to $I(k)$ in blue, and cross atom contributions in red. The lower panels show hydroxyl atom contributions in blue, alkyl tail carbon atom contributions in gray, and cross contributions in green (see text).

The structure factors in the lower panel are divided into those of the hydroxyl groups (in blue), the methyl groups (in gray) and the cross correlation between the two (in green). The specificity of the various atoms are intentionally ignored, since we want to highlight the influence of head groups versus the alkyl tail, which is at the origin of the local segregation in Fig.6 and Fig.9. The $I(k)$ in Eq.(2) can be rewritten as

$$I(k) = I_{\text{like}}(k) + I_{\text{cross}}(k)$$

with

$$I_{\text{like}}(k) = r_0^2 \rho \sum_i f_i^2(k) S_{ii}^{(T)}(k)$$

$$I_{\text{cross}}(k) = r_0^2 \rho \sum_{i \neq j} f_i(k)f_j(k)S_{ij}^{(T)}(k)$$

where we have separated the like atom contributions $I_{\text{like}}(k)$ from those from the different atoms $I_{\text{cross}}(k)$. From the lower panels of Fig.12, it is clear that $I_{\text{cross}}(k)$ is likely to contain negative anti-peak contributions in the case of some alcohols, such as 1-pentanol and 1-octanol, for example. The upper panels of Fig.12 show in black lines the total $I(k)$ built from all total structure factors from Eqs.2,3, as well as the contributions to $I(k)$ coming from like atoms and cross atomic contributions. Comparing the structure factors in the lower panel, to the $I(k)$ in the upper one, it is clear how the like contributions tend
to contribute positively to the pre-peak, winning over the negative ones from the cross contributions. It is important to note that in the alkyl tail carbon atom contributions, those carbons close to the hydroxyl head contribute to a positive pre-peak (since they are strongly attached to the head group), while the cross correlation of these atoms with the trailing carbons contribute to a negative anti-peak. This explains why there are negative contributions in the gray curves as one goes to longer alcohols. It is also noteworthy that structure factors between trailing carbon atoms do not have any pre-peak, and look just standard Lennard-Jones structure factors. In other words, the tail atoms by themselves do not contribute to the pre-peak. This remark shows the importance of interpreting $I(k)$ as resulting from statistical correlations between different parts of the meta-objects, and not only to some of them, such as the hydroxyl group chain.

Looking closer at the lower plots, we notice that the structure factors of ethanol have almost no anti-peaks. The reason for this is simply because there is only one neutral atom: the tailing methyl group. This is not the case any more for 1-pentanol and 1-octanol, and we clearly see the importance of the anti-peaks. The positive pre-peak of the hydroxyl correlations also grows in intensity when going from ethanol to 1-octanol, reflecting the tightness of the hydrogen bonding for larger mono-ols. Another feature is the apparent inversion of peak heights between the structure factors and $I(k)$. Indeed, the structure factors show a large pre-peak and a smaller main peak, while this is inverted in $I(k)$. The reason is that it is the number of contributions in Eq.(2) which matters, and the smaller main peaks of the structure factor end up overwhelming that of the pre-peak, resulting in the actual shape of $I(k)$. Details of these correlations, intentionally undifferentiated here, will be discussed in a subsequent work.

3.6.2 The important role of the intra-molecular correlations

In the previous part we have seen that the cross domain correlations involve large negative contributions in $I(k)$. Such contributions could potentially cancel the pre-peak contributions in some cases, and they indeed do, as we have demonstrated for the case of aqueous 1-propanol mixture [60]. This observation poses the problem of how to interpret the appearance of a pre-peak in $I(k)$ as residual part of canceling effects, and more importantly how to relate this to an underlying microscopic physics such as self-assembly. For example, in another work [48], we interpreted the experimental evidence for the appearance of a scattering pre-peak in aqueous 1-propylamine mixtures, as opposed to its absence in aqueous 1-propanol mixtures, as the signature of water hydrogen bonding to the amine and hydroxyl groups is different for each type of mixtures. In the present case, we observe that, if instead of the correct $w_{ij}(k)$ involving the molecular flexibility, we use the approximation of the rigid molecule (see Section 2.4.1), then the pre-peak of longer alcohols vanishes, or even becomes a negative anti-peak, which is an unphysical feature since $I(k)$ should be positive for all $k$. This is illustrated in Fig.12, where we show, for the case of the 3 alcohols selected for Fig.10, the like $I_{\text{like}}(k)$ and cross contributions $I_{\text{cross}}(k)$ together.
with the total $I(k)$, for the cases of the rigid and flexible $w_{ij}(k)$, respectively in dashed and full lines.

Figure 13: Illustration of the crucial influence of flexibility in the pre-peak description, for the OPLS model of 3 alcohols (ethanol, 1-pentanol and 1-octanol). The full curves are for when flexibility is properly included in the $w_{ij}(k)$ functions (see Fig.1), and thin dashed curves when $w_{ij}(k)$ for rigid molecules is used. Black curves are for total $I(k)$, blue curve for like atom-contributions of $I(k)$ and red curves for cross atom contributions.

It is quite clear that, in the case of the rigid intra-molecular approximation with dashed lines, the pre-peak feature is considerably modified and does not match the respective experimental data. Also, this effect is more pronounced for longer alkanols. Indeed, for 1-octanol, $I(k)$ is seen to develop an unphysical negative anti-peak (black dashed curve in the left panel) when flexibility is not properly described, and the dashed red curve shows that the negative contribution comes from the overwhelming cross atoms structure factor, involving in particular correlations between the hydroxyl $h$

4 Discussion

Radiation scattering in liquids is a statistical probe of the microscopic features in the disorder, which is why it is related to purely statistical observable such as correlation functions and structure factors, as in the Debye formula. It is the analysis of the structure factors which would uncover which microscopic features contribute to specific features of the scattered intensity, such as pre-peaks. Many types of investigations of alcohols point towards the existence of hydroxyl group aggregates. But the present work shows clearly that it is not only these aggregates which contribute to the features of $I(k)$, but the entire meta-object formed by the surrounding alkyl tails as well, which is seen by the scattered radiation. The alkyl tails are shown to play an increasingly important role with $n$, particularly through their flexibility. More importantly, it appears necessary to include the anti-correlations between the hydroxyl head group and the alkyl tail, in order to interpret the decrease of the pre-peak when these alkyl
tails grow in size beyond \( n = 4 \). The study clearly suggests that the pre-peak cannot be fully understood with a model free investigation, hence necessarily involving the penalties associated with modeling shortcomings.

The comparison of the experimental and simulated \( I(k) \) shows the strong model dependence, and in particular the erratic behaviour of the CHARMM model deserves some investigation. This model differs from the OPLS and TraPPE models by the all-atom representation of the methylene and methyl groups, but more importantly by the fact that the corresponding hydrogen and central carbon bear relatively large partial charges. In addition, unlike the two other models which assign the same partial charges to the same atoms throughout the various alcohols, the CHARMM model has different adjustments depending on alcohols (see Tables). In view of the argumentation in the previous section, where we have shown the importance of charge ordering in the microscopic structure, we believe that this model blurs the charge ordering and diminish strongly the hydroxyl group association.

It is an experimental observation that the pre-peak in \( I(k) \) is always positive. In view of Eq. (2) and the existence of pre-peak and anti-peaks in the \( S_{ij}(k) \), one may ask why their summed contribution in Eq. (2) should always lead to a positive contribution in the experiments. The example of the CHARMM model shows that this is not always the case, as in the case of 1-pentanol, hence indicating that this model has something unphysical about it. The CHARMM model, in a sense, demonstrates that it is possible to obtain a micro-structure obeying charge ordering and domain ordering criteria, and yet give unphysical quantities, such as no-prepeak or negative anti-peaks.

However, one should not conclude that all all-atom models are bound to such fate. Indeed, we have tested the OPLS-AA for methanol and ethanol, and it gives a pre-peak for \( I(k) \), although somewhat lower than expected, when compared to the OPLS-UA model. Looking at the partial charges of this model in the Tables, we notice that those on the methyl/methylene groups are somewhat smaller than those of the CHARMM model. In a way, these groups will have less tendency to blur the charge ordering.

In order to support these arguments, we show in Fig.13 a comparison of the OPLS-AA model for methanol and ethanol with OPLS-UA and CHARMM.

It is seen that, for methanol, the OPLS-AA model misses the pre-peak shoulder feature, just like the CHARMM model. For ethanol however, OPLS-AA shows a pre-peak, although weaker than the OPLS-UA model. When looking at the charges of the terminal C group in Table 5, we find that OPLS-AA has 0.06 for the hydrogen, while CHARMM has 0.09, a higher value. In other words, the Coulomb interaction of the carbon group hydrogen would be more strongly felt for CHARMM than for OPLS-AA, leading to a stronger charge order homogeneity breaking for this latter model. This explains why OPLS-AA has stronger pre-peak than CHARMM.

Interestingly, OPLS-AA has a cluster distribution similar to that of CHARMM in Fig.7, suggesting that all AA models tend to favor higher monomer probability than their UA counterpart. We note that this is not in contradiction with the similarity of the correlation functions and structure factors in what concerns
Figure 14: Calculated $I(k)$ for methanol (top panel) and ethanol (bottom panel) illustrating the ability of all-atom and united-atom models to reproduce the pre-peak. CHARMM-AA in green, OPLS-AA in magenta and OPLS-UA in green.

the clustering features, since AA models have also a large number of clusters, as indicated by the large cluster distribution tail in Fig.6 for CHARMM.

5 Conclusion

In this work, we have compared the calculated X-ray scattering intensities $I(k)$ for a variety of mono-ols, ranging from methanol to 1-nonanol, to the corresponding experimental data (methanol to 1-undecanol), and for several of known force field models, such as OPLS, TraPPE, CHARMM and GROMOS. The principal focus of this calculation was twofold: on one hand to test the ability of various types of force field models to reproduce the overall shape of the experimental $I(k)$, and particularly the well known scattering pre-peak feature of these alcohols, and on the other hand to provide an explanation for the generic features of $I(k)$ across different alcohols. Two notable such features are the decrease of $k_P$ with increasing values of $n$, and the amplitude $A_P$ turnover around 1-pentanol.

As far as the first point is concerned, we find in general that the OPLS-UA model provides better results. In particular it reproduces well the pre-peak’s position and relative intensity variations throughout the whole series of mono-ols studied. The TraPPE model is less performant, particularly in what concerns the main-peak. But this model predicts pre-peak for all alcohols. The CHARMM model is the most problematic, particularly for small alcohols, since it fails to predict the pre-peak for methanol and ethanol, and predicts an unphysical negative anti peak for 1-pentanol. However, it is good for longer
The problems found for TraPPE and CHARMM around 1-pentanol could well be related to the physical fact that there is a crossover of behaviour of the aggregated structures around that particular alcohol. As far as the second point is concerned, we have shown that the origin of the pre-peak in $I(k)$ is not only related to the hydroxyl head groups aggregates, but to the entire meta-object formed by this central chain and the surrounding alkyl tails, promoting the picture of such alcohols made not only of monomer alcohol molecules, but also meta-objects. This is illustrated in Fig.15. $I(k)$ would then detect both the atomic constituents through the main peak and the meta-object through the pre-peak. More specifically, the pre-peak is the result of correlations between the charged head groups part and cross-correlations between the head and tail parts. The duality of the meta-object and the correlations between its part appears as a whole. Earlier studies of methanol reported the large variety of cluster shapes, such as chains, simple and branched, loops, lassos, etc... \[50, 51, 4, 5, 51, 52, 53, 21, 12, 9, 10\]. The present work indicates that this is also the case for all mono-ols.

Pre-peaks are found in many soft-matter systems \[62, 63, 64, 65, 66\], particularly in micro-emulsion systems, and geometric shapes of the aggregates are often deduced from the shape of $I(k)$. In view of Eq.(1) and the fact that aggregates are better defined from the atomic structure factors themselves, it is quite possible that a more detailed analysis and interpretation in terms of charge and domain order via atom-atom correlations may provide deeper insight into self-assembly. In any case, the need for computer simulations appears as necessary, in order to calculate the atom-atom structure factors. This could pose quite
formidable modeling problems, specially when considering typical soft-matter systems. We hope that the present investigation has provided elements for reconsidering usual interpretations of $I(k)$ in terms of microscopic properties, by establishing the need for the dual aggregate/correlation picture.

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