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Size-Resolved Photoelectron Anisotropy of Gas Phase Water Clusters and Predictions for Liquid Water

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ABSTRACT

We report the first measurements of size-resolved photoelectron angular distributions for the valence orbitals of neutral water clusters with up to 20 molecules. A systematic decrease of the photoelectron anisotropy is found for clusters with up to 5-6 molecules, and most remarkably, convergence of the anisotropy for larger clusters. We suggest the latter to be the result of a local short-range scattering potential that is fully described by a unit of 5-6 molecules. The cluster data and a detailed electron scattering model are used to predict the anisotropy of slow photoelectrons in liquid water. Reasonable agreement with experimental liquid jet data is found.

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A detailed understanding of elastic and inelastic scattering of electrons in liquid water is of fundamental importance for modelling radiation damage in biological systems, describing the behaviour of the solvated electron in chemistry, and for the quantitative interpretation of photoelectron spectra of liquid water and aqueous solutions [1-8]. For slow electrons (electron kinetic energy eKE \leq 50 eV), detailed experimental scattering parameters (differential cross sections, energy losses) were only reported for amorphous ice [9] - apart from *liquid water* data for eKE \leq 6 eV recently obtained from photoelectron velocity map imaging (VMI) of water droplets [10]. As the energetics of electronic scattering processes (dissociative electron attachment, electronic excitations, ionization) are hardly affected by the finer details of the bulk environment, there is little reason to expect substantial differences between amorphous ice and liquid water for eKE \geq 6 eV [9]. The amorphous ice and liquid droplet data [9,10] should thus provide a reasonable basis for scattering simulations of liquid water. In addition, electron attenuation lengths (EALs) for eKE \gtrsim 3 eV are available from various microjet studies [11-13]. The individual scattering contributions required for quantitative predictions can however not be extracted from EALs because EALs are broad averages over many different scattering processes.

As the photoelectron angular distribution (PAD) is particularly sensitive to electron scattering it has recently received increasing attention in this context [7,10,13-18]. It is often described by a single anisotropy parameter β , defined by

$$I(\theta) \propto 1 + \frac{\beta}{2} (3\cos^2 \theta - 1).$$
 Eq. (1)

 θ is the angle between the light polarization and the ejection direction of the photoelectrons. $I(\theta)$ is the electron signal detected at that angle. For the liquid microjet, where the spherical symmetry of the sample is broken, this is an approximation, which we also use in the present work. For ionization from the O1s orbital of water, Thürmer et al. observed a more isotropic PAD (a smaller β -value) for the liquid than for the gas phase in the eKE range $\sim 12-450$ eV [13]. For core-level ionization, this reduction should mainly arise from electron scattering within the liquid. For the ionization from the valence orbitals 1b₁, 3a₁, and 1b₂, additional changes in the initial state due to orbital mixing, partly mediated by hydrogen-bonding, are expected to contribute to differences in β between gas and liquid phase. While monomer gas phase β -parameters have been reported for the three valence orbitals at photon energies 18 eV $\leq hv < 139$ eV [15,16,19-21], corresponding liquid water values have only been reported at hv = 38.7 eV [16]. Zhang et al. [15] made a first attempt to distinguish between initial state and scattering effects on β , based on measurements at hv = 40 and 60 eV of (H₂O)_n clusters with broad size distributions and estimated average sizes of $\langle n \rangle \geq 58$. The results hinted at intrinsic differences between gas phase monomer and cluster PADs arising from alterations in

the initial states. Table T1 [22] summarizes the existing literature values for β -parameters of water clusters and liquid water.

The present work reports double imaging photoelectron photoion coincidence ($i^{2}PEPICO$) measurements of small $(H_2O)_n$ clusters (n ≤ 20) recorded at the synchrotron radiation facility SOLEIL [22-27]. As a unique feature, this technique allows us to record photoelectron VMIs for a particular cluster size n. Avoiding averaging over different cluster sizes and circumventing the issue of the overlap with the strong water monomer signal we can extract cluster size-resolved β -parameters. Size selectivity is particularly important for small clusters, where pronounced changes in β are expected for size changes by just one water molecule. To clarify the evolution of PADs with increasing cluster size is our main goal. Clusters are not expected to be good models for the PAD of the liquid since most of the effects of elastic and inelastic scattering during electron transport in the bulk (referred to as contribution (iv) below) would be missing. Still, many local effects (referred to as contributions (i)-(iii)) can already be observed in clusters. As a link between the monomer and the condensed phase clusters contribute to a better understanding of the complex electron scattering in liquid water. With this in mind, we predict β -parameters for typical liquid water microjet experiments using a detailed scattering model [7,9,10,22]. We focus on slow electrons with eKEs \leq 65 eV where PADs sensitively depend on electron scattering.

Fig.1 shows β for the gas phase monomer (n = 1) in the range 13.0 eV $\leq hv \leq$ 35.0 eV with corresponding values tabulated in Table T2 [22]. Fig.S2 [22] provides a comparison with published data for $hv \geq 18$ eV [15,16,19,20]. Our monomer data agree well with literature values and provide the first values below ~18 eV, which clearly confirm the trend towards low anisotropies at very low eKEs (~0.4 - 5.4 eV) predicted by calculations [21]. Fig.1 also includes a summary of our experimental β -parameters for (H₂O)_n with $2 \leq n \leq 20$, recorded with i²PEPICO (see Table T2 [22] for corresponding values with respective uncertainties). For larger clusters at higher hv, some data points are missing because the signal to noise ratio was insufficient to determine reliable β -parameters. Photoionization of a neutral water cluster (H₂O)_n is accompanied by a fast intracluster proton transfer with subsequent loss of an OH radical [28-33]:

$$(H_2O)_n + h\nu = (H_2O)_{n-1}H^+ + OH + e^-$$
 Eq. (2)



Figure 1: Lines labelled n=1 to n=20: Experimental β -parameters for H₂O monomer and (H₂O)_n clusters with $2 \le n \le 20$ recorded at 13.0 eV $\le hv \le 35.0$ eV. (a), (b), and (c) correspond to ionization from the 1b₁, 3a₁, and 1b₂ orbitals of water, respectively. Monomer values for eKEs < 25 eV from this work and for eKEs > 25 eV from ref. [20]. For (H₂O)₂, separate traces are shown for H₃O⁺ (full brown line; n=2 Eq. (2)) and for the intact dimer (H₂O)₂⁺ (brown crosses) (see Fig.2b). **Green triangles:** monomer (open symbols) and a cluster ensemble (full symbols) with an average cluster size $\langle n \rangle \square$ 58 from ref. [15]. **Blue squares:** monomer (open symbols) and liquid water (full symbols) from the microjet study by Faubel *et al.* [16]. **Calculated anisotropy parameters** for liquid water microjets: Open black circles: $\beta_{n=1}^{liquid}$ calculated with monomer values (n=1) as input for the local anisotropy in the liquid. Open orange diamonds: β_{liquid}^{liquid} calculated with gas phase elastic scattering cross sections alone [34].

Accordingly we assign clusters with n molecules to VMIs recorded in coincidence with mass $m = (n \cdot 18) - 17$. For small clusters the subsequent slow loss of water molecules from the initially formed protonated cluster is dominated by monomer loss with total decay fractions < 0.3 [28,29,35]. Fig.2 shows exemplary photoelectron spectra for n = 1, 2, and 6. In larger clusters polarization effects shift the vertical electron binding energy (VBE), i.e. the most probable electron binding energy (eBE), towards lower values (Fig. S4 [22]), although not yet reaching the liquid bulk value [36] or the values for large clusters [15,37]. The downward trend in cluster VBEs is consistent with the evolution of the cluster ion appearance energy from ref. [28]. The dimer spectrum in Fig. 2b consists of the two contributions from the intact dimer (H₂O)₂⁺ to ionization from the lone pair of the hydrogen-bond donor (referred to as (b₁)_D), while H₃O⁺ results from the ionization of an orbital delocalized over both hydrogen-bond donor and -acceptor (referred to as (a₁/b₁)) with a clearly different β -parameter (Fig.1a, brown crosses and full line, respectively). Table T3 [22] compares the corresponding dimer VBEs with literature data.

Fig.1 provides the first quantitative β -values for the initial condensation steps (n = 2-20). The largest absolute decrease of the anisotropy with increasing cluster size is observed for the 1b₁ orbital (out-of-plane lone pair), followed by the 3a₁ orbital (in-plane lone pair). The 1b₂ orbital (σ_{OH} bond orbital) shows the smallest variations in β , but within the estimated uncertainty no systematic trend with cluster size. The larger sensitivity of β for ionization from 1b₁ compared with 1b₂ seems reasonable because the 1b₁ orbital acts as an acceptor in hydrogen bonds, while 1b₂ (σ_{OH}) is not directly involved in hydrogen bonding. Similar trends compared with the monomer were observed for the cluster ensemble at 40 eV [15] and the liquid microjet at 38.7 eV [16]. The most striking result in Fig.1 is the convergence of β for the two outermost valence orbitals for cluster sizes n \gtrsim 5-6. A simple estimate based on the maximum number of monomers that can evaporate from a cluster after proton transfer and the reported cluster decay rates shows that the slow cluster evaporation does not significantly affect the β -value derived for a given cluster size [28,29,35] and thus cannot explain the observed convergence.

We suggest the following qualitative explanation for the systematic decrease of $|\beta|$ with increasing n and its convergence for $n \ge 5-6$. (Note that for larger clusters the observed β is the average over several conformers). The difference between molecular and cluster PADs arises from several contributions: (i) A change in the initial molecular electron wavefunction, i.e. *a change in the local orbital character* (polarization, orbital mixing) due to condensation. In (H₂O)_n clusters, electron delocalization over hydrogen-bonds is likely a major factor determining the change of orbital character. For increasing cluster size, changes in the orbital character typically reduce $|\beta|$. (ii) The second contribution is attributed to *multicentre* *ionization* – again a change in the initial state: The larger the cluster the more equivalent units are taking part in a single ionization event (quasi-degeneracy). Interference of partial waves from many centres tends to reduce $|\beta|$, likely the more pronounced the larger the cluster. (iii) The third contribution comes from a *change in the ion core potential*, by which the outgoing electron wave is scattered. This is the result of the delocalization of the electron hole over nearest neighbours connected through hydrogen bonds, as discussed in detail for the case of the dimer in ref. [30]. Again, this tends to reduce $|\beta|$. Qualitatively, all three contributions favour more isotropic PADs, i.e. a decrease of $|\beta|$, with increasing cluster size. This expectation agrees with the experimental observation for cluster sizes up to n = 5-6 (Fig.1). The convergence of β observed for $n \gtrsim 5-6$ implies that the range of contributions (i)-(iii) only extends over a few molecules. $n \approx 5-6$ coincides with the smallest cluster sizes for which three-dimensional hydrogen-bond networks with more than two hydrogen bonds per water molecule become more stable than ring-topology structures ([38-41] and references therein). It is plausible that the typical range for changes in orbital character and in the ion core potential is comparable to the range of local hydrogen-bridges. Similarly interference effects from multicentre ionization should be most pronounced just in a local environment. The convergence of β for $6 \leq n \leq 20$ agrees with an intrinsic, short-range scattering potential that is described by a cluster with $n \approx 6$. Since the spatial extent of clusters with $n \leq 20$ is very small (~7-10 Å) the long-range scattering potential is essentially an unshielded Coulombpotential. Even semi-quantitative descriptions of the cluster PADs would require very highlevel quantum chemical calculations [17,42-44] - still a big challenge for such complex systems.

In principle, the $|\beta|$ -value of larger clusters could also be reduced by non-local *elastic and inelastic scattering of the electrons during electron transport through the cluster*, referred to as contribution (iv). We determine its influence by simulating cluster VMIs for different cluster sizes with our scattering model described in section S3 [22] and [10]. It is based on a Monte-Carlo solution of the transport problem, equivalent to a random succession of suitably parameterised isolated scattering events. Our simulations explicitly take into account the cross sections, energetics, and angular-dependences of all relevant scattering processes (inelastic electron-phonon, electron-vibron, electron-electron scattering and elastic scattering). For $n \leq 50$ the influence of contribution (iv) is negligible (Fig. S7 [22]). A significant change of β (on the order of 0.1) is only found for clusters with more than $n \approx 100$ molecules i.e. beyond cluster sizes studied here. This also shows that previously used simple modelling approaches, such as gas phase scattering between the monomers in a cluster, are not suitable to describe the cluster behavior.



Figure 2: Experimental photoelectron spectra of (a) water monomer and $(H_2O)_n$ clusters for (b) n=2 and (c) n=6. Selected VMI images are shown in Fig. S3 [22]. The dimer spectrum has contributions from the intact dimer $(H_2O)_2^+$ (red line) and from H_3O^+ (black line) formed after fast proton transfer. (d) Calculated photoelectron spectra for the liquid water microjet for two polarization directions $\theta = 0^\circ$ (blue line) and 90° (black line) of the light (Fig.S5 [22]) for monomer input $\beta_{n=1}$ (Fig.1). The photon energies *hv* are indicated in the figure.

The water dimer is a special case because ionization from the lone pair of the hydrogenbond donor $(b_1)_D$ is distinguishable from ionization of the mixed (a_1/b_1) orbital, which is delocalized over donor and acceptor (Fig.2b). The β -parameters for $(b_1)_D$ (brown crosses in Fig.1a) are slightly lower than the monomer value (n=1, full black circles). The $(b_1)_D$ orbital can be considered as a monomer orbital disturbed by the presence of the second H₂O molecule. As it is not directly involved in the hydrogen-bond the decrease in β relative to the monomer mainly arises from contribution (i). The larger decrease of β for the (a_1/b_1) orbital (n=2, full brown line) results from the simultaneous action of all three contributions (i)-(iii). It is unfortunately not possible to estimate the relative magnitude of (i) compared with (ii-iii) from the absolute changes of the two different dimer β -parameters, but the general trend is consistent with our expectation that contributions (i)-(iii) tend to reduce $|\beta|$. When comparing absolute changes one must account for the non-linearity of the relation between β and the observed signal $I(\theta)$ (Eq.(1)).

In contrast to the clusters studied here, the liquid water PAD is not only determined by the three local contributions (i)-(iii), but also strongly by contribution (iv), i.e. elastic and inelastic scattering during electron transport in the liquid. We treat this effect within the detailed scattering model mentioned above (section S3 in [22] and [7,9,10]) to simulate a typical liquid microjet experiment [12-14,16-18], illustrated in Fig. S5 [22], where β^{liquid} is determined from polarization-dependent (θ -dependent) measurements. We assume the local contributions (i)-(iii) in liquid water to be either the same as in the monomer or the same as in a cluster with n=6 (converged cluster value). Correspondingly we use either the experimental monomer ($\beta_{n=1}$; full black circles) or the experimental cluster ($\beta_{n=6}$; full orange diamonds) β -parameters from Fig.1 to describe the local anisotropies in the liquid. With the chosen local input anisotropy $(\beta_{n=1} \text{ or } \beta_{n=6})$ and our scattering model to account for contribution (iv) we calculate the liquid anisotropy parameters, $\beta_{n=1}^{liquid}$ (open black circles) and $\beta_{n=6}^{liquid}$ (open orange diamonds), respectively, shown in Fig.1. The comparison with $\beta_{n=1}$ and $\beta_{n=6}$ illustrates the pronounced effect of contribution (iv) on the PADs of the liquid, viz. a reduction of the anisotropy. Fig.2d shows examples of photoelectron spectra of the liquid calculated for 0° and 90° laser polarization (see also Fig. S6 [22]), which agree well with experimental liquid-jet spectra [16,36] (note the large gas phase fractions in the spectra of ref. [16]). We finally note that polarization-dependent liquid jet measurements in principle yield marginally higher β -values than other methods, such as VMI, because the coupling of the electromagnetic radiation into the jet depends on the laser polarization (section S3, p. S9 [22]).

To the best of our knowledge experimental values for β^{liquid} in the valence region were only reported at hv = 38.7 eV in a microjet study by Faubel *et al.* [16] (Fig.1, full blue squares at eKEs ~21-28 eV). They lie reasonably close to our calculated β^{liquid} -values in Fig.1. The agreement is best for b₁, where monomer and liquid bands are well separated in the photoelectron spectrum (Figs. 2a and d). The fact that for a₁ and b₂ the values of Faubel *et al.* lie above our calculations might be attributed to overlapping monomer bands not fully accounted for in the analysis of their experiment. Note the high β values of the monomer in the corresponding eKE range. In principle, our model should underestimate the true β^{liquid} because it does not account for the strong shielding of the ion core potential in the liquid, which is mainly dielectric screening. It reduces the range of the ion core potential by roughly an order of magnitude compared with the isolated monomer or cluster. Our simulations with input values $\beta_{n=1}$ and $\beta_{n=6}$ do not include this effect. A shorter range of the ion core potential reduces the influence of scattering by the ion core on the local anisotropy of the electron. This effect could be described by correspondingly higher input values for $\beta_{n=1}$ and $\beta_{n=6}$. This in turn would yield higher calculated β^{liquid} -values. The apparent better agreement of the experimental data by Faubel *et al.* with $\beta_{n=1}^{liquid}$ than with $\beta_{n=6}^{liquid}$ appears to be fortuitous. Once dielectric screening is included, $\beta_{n=6}^{liquid}$ should actually agree better with experimental liquid bulk data, simply because the cluster input $\beta_{n=6}$ better represents the contributions (i)-(iii) as discussed above. A simple estimate of the influence of shielding is unfortunately not possible. Such estimates would require high-level ab initio calculations. In Fig.1 we also add a calculation for the liquid anisotropy $\beta_{elastic}^{liquid}$ (open black stars), for which we used just *elastic gas phase* monomer scattering cross sections [34] instead of the proper condensed phase values as for the other simulations [9,10]. The resulting $\beta_{elastic}^{liquid}$ are almost isotropic and clearly disagree with the experimental values at hv = 38.7 eV. This demonstrates that gas phase scattering parameters are not suitable to describe the liquid.

In summary, photoelectron photoion concidence imaging provides size-dependent photoelectron anisotropy parameters of $(H_2O)_n$ clusters for $n \leq 20$. The experimental data suggest that intracluster electron scattering in clusters containing between ~ 6 and 20 molecules is mainly determined by the short range potential of a unit consisting of 5-6 molecules, coinciding with the smallest cluster sizes for which three-dimensional hydrogenbond networks become the most stable structures. It seems reasonable that the short range scattering potential in liquid water is largely determined by this smallest unit; i.e. approximately by the first solvation shell. In contrast to clusters, however, the ion core potential is strongly shielded in the liquid. While a quantitative estimate of this effect can at present not be provided, it appears plausible that shielding will increase the anisotropy compared with the experimental cluster data. We suspect that the major difference between small clusters (n \leq 100) and the liquid arises from the additional elastic and inelastic electron scattering in the liquid. A detailed scattering simulation for the liquid starting from cluster anisotropies of the smallest unit confirms this presumption. Even with the shielding effect neglected, this model provides reasonable agreement with experimental liquid jet data. Our simulations demonstrate that gas phase scattering parameters are generally not appropriate for electron scattering in the liquid. Further validation of the role of the smallest cluster unit and the shielding in the liquid awaits more experimental data from liquid jets and larger water clusters as well as in-depth theoretical studies.

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