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Static structure factor and electron correlation effects studied by inelastic x-ray scattering spectroscopy
X-ray emission spectroscopy of bulk liquid water in “no-man’s land”

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The structure of bulk liquid water was recently probed by x-ray scattering below the temperature limit of homogeneous nucleation ($T_H$) of ~232 K [J. A. Sellberg et al., Nature 510, 381-384 (2014)]. Here, we utilize a similar approach to study the structure of bulk liquid water below $T_H$ using oxygen K-edge x-ray emission spectroscopy (XES). Based on previous XES experiments [T. Tokushima et al., Chem. Phys. Lett. 460, 387-400 (2008)] at higher temperatures, we expected the ratio of the $1b_1''$ and $1b_1''$ peaks associated with the lone-pair orbital in water to change strongly upon deep supercooling as the coordination of the hydrogen (H-) bonds becomes tetrahedral. In contrast, we observed only minor changes in the lone-pair spectral region, challenging an interpretation in terms of two interconverting species. A number of alternative hypotheses to explain the results are put forward and discussed. Although the spectra can be explained by various contributions from these hypotheses, we here emphasize the interpretation that the line shape of each component changes dramatically when approaching lower temperatures, where, in particular, the peak assigned to the proposed disordered component would become more symmetrical as vibrational interference becomes more important. © 2015 AIP Publishing LLC. [http://dx.doi.org/10.1063/1.4905603]

I. INTRODUCTION

Water has been studied intensively for hundreds of years but is still far from fully understood. Knowledge about the H-bond network in water is essential for understanding its numerous unusual chemical and physical properties. For example, water can exist in the liquid state far below its melting temperature (supercooled water) where its thermodynamic response functions, such as coefficient of thermal expansion ($\alpha_T$), isothermal compressibility ($\kappa_T$), isobaric heat capacity ($C_P$), and correlation length ($\xi$), appear to diverge at a difficult to reach temperature of 228 K.¹–³ Water crystallization occurs very rapidly below the homogeneous nucleation temperature ($T_H$) of ~232 K,⁴ which has prevented measurements of the liquid phase and led to a “no-man’s land” devoid of experimental results. The anomalous behavior of water...
becomes important already at ambient conditions giving a maximum density at 277 K, and a minimum in $C_P$ and $\kappa_T$ at 308 K and 319 K, respectively.

The apparent divergence of $C_P$ and $\kappa_T$ upon supercooling has led to several proposed scenarios involving fluctuations between local high-density liquid (HDL) and low-density liquid (LDL) configurations. These fluctuations could, upon deeper supercooling, lead either to a liquid-liquid transition (LLT) with a liquid-liquid critical point (LLCP) at positive pressure\textsuperscript{3} or a continuous transformation without discontinuity, called the singularity-free (SF) model.\textsuperscript{6} In the critical point-free (CPF) model, the LLCP would instead occur at negative pressure\textsuperscript{7} (outside the physically relevant part of the phase diagram), but there is also a recent suggestion that only a liquid-solid transition exists,\textsuperscript{8} which is currently under debate.\textsuperscript{9–14} Alternatively, a thermodynamic stability limit (SL) of the liquid state has been suggested\textsuperscript{15} and that ice crystallization occurs on a timescale faster than liquid equilibration in “no-man’s land.”\textsuperscript{16} These scenarios do not agree, and to date convincing experimental data have been lacking. Recently, a new technique has been demonstrated where it is possible to enter “no-man’s land” through fast evaporative cooling of micron-sized water droplets and ultrafast probing using x-ray scattering from an x-ray laser.\textsuperscript{17} It was shown that water could be probed in a metastable liquid phase down to 227 K and that a continuous increase of structures with local tetrahedral coordination became more enhanced upon deep supercooling, which indicates an accelerated transition toward LDL.

There are several spectroscopic techniques such as vibrational spectroscopy and core-level spectroscopy that are sensitive to the hydrogen-bonding (H-bonding) environment.\textsuperscript{18} In particular, recent x-ray emission (XE) spectra show a split of the lone-pair (1b\textsubscript{1}$^\prime$) peak in water that has been interpreted as arising from structures with distorted H-bonds (1b\textsubscript{1}$^\prime$) and from tetrahedrally H-bonded structures (1b\textsubscript{1}$^\prime$).\textsuperscript{18,20–27} As the temperature is lowered, intensity is transferred from the low-energy 1b\textsubscript{1}$^\prime$ peak to the high-energy 1b\textsubscript{1}$^\prime$ peak, but no broadening or additional features are observed.\textsuperscript{20,28} Also, selective excitations show a clear correlation between the two 1b\textsubscript{1}$^\prime$ peaks and the various resonances in the x-ray absorption spectrum of water consistent with the hypothesis of two structural motifs.\textsuperscript{20,28} An interpretation has been proposed in terms of these local structures being related to fluctuations in the liquid giving rise to local regions of HDL (distorted) and LDL (tetrahedral) character.\textsuperscript{18,22,28,29}

However, the strong isotope effect in the spectra has led to other suggestions of the origin of the two 1b\textsubscript{1}$^\prime$ peaks.\textsuperscript{19,20} The first alternative interpretation suggests that the two peaks are due to intermediate or final states of ultrafast dissociation during the core-hole lifetime.\textsuperscript{19,30} The peak at low emission energy was proposed to come from OH$^-$ species generated from water through core-hole-induced dissociation while the other was proposed to be due to intact water.\textsuperscript{19,30} The second alternative interpretation, based on theoretical simulations, proposed that the peak at high energy is of lone-pair 1b\textsubscript{1} symmetry, whereas the peak at low emission energy is of 3a\textsubscript{1} symmetry and originating from the strong dynamical response toward dissociation found in the simulations.\textsuperscript{31,32} Excitation-selective, polarization-dependent measurements indicated first that there could be a difference between the two peaks in terms of symmetry,\textsuperscript{33} but later more data and better statistics demonstrated that the two lone-pair peaks are both of 1b\textsubscript{1} symmetry ruling out that the 1b\textsubscript{1}$^\prime$ peak would be of a\textsubscript{1} symmetry.\textsuperscript{37} Furthermore, theoretical calculations of the XE spectrum of a model water dimer including vibrational-lifetime interference in a full Kramers-Heisenberg treatment show only the appearance of an asymmetrical line shape and not two distinct peaks.\textsuperscript{34,35} The asymmetry of the 1b\textsubscript{1} peak of the lighter isotope is much larger which could explain the isotope effect.

Based on the interpretation in terms of two structural motifs, we would expect a rapid rise in intensity for the 1b\textsubscript{1}$^\prime$ peak if the liquid is cooled down to temperatures within “no-man’s land.” Here, we show that this is not the case. We will review several possible explanations for this result. Although no definite conclusion can be drawn, it is essential that these data are made available, since it is the first x-ray emission spectroscopy (XES) study of liquid water venturing into “no-man’s land” and since it led to results in seeming contradiction to expectation.

II. METHODS

A. Experimental setup at the Soft X-ray Materials Science (SXR) instrument, Linac Coherent Light Source (LCLS)

Non-resonant O K-edge XES was performed at the SXR instrument\textsuperscript{36} at the LCLS using a nominal photon energy of 550 eV (well above the O K-edge absorption resonance) and a nominal pulse duration of 100 fs. These pulses, delivered at 120 Hz and containing between 6×10\textsuperscript{11} and 12×10\textsuperscript{11} photons/pulse (accounting for 10% beamline transmission from gas detectors to the sample\textsuperscript{37}), were focused to a focal spot size of 75×520 μm\textsuperscript{2} (horizontal × vertical) at the interaction point, corresponding to a peak fluence of 0.27 J/cm\textsuperscript{2}, which is comparable to the threshold of ∼0.2 J/cm\textsuperscript{2} where a non-linear fluence dependence of the XE yield starts to occur due to valence-hole reabsorption.\textsuperscript{37} Liquid D\textsubscript{2}O (Fischer Scientific #AC166311000, 99.95%); and Sigma-Aldrich #151882, 99.9 at. %) was delivered to the interaction region using the Liquid Jet End Station, which consists of a liquid microjet in a differentially pumped high vacuum (2×10\textsuperscript{-3} millibars at the jet, 4×10\textsuperscript{-7} millibars at the detector) and a Grazing spectrometer XES 350\textsuperscript{38} mounted at 90° with respect to the incident x-ray beam, as described in detail elsewhere.\textsuperscript{39} The liquid microjet was produced by a gas-dynamic virtual nozzle (GDVN) operated at 800 psi liquid pressure and 330 psi He gas pressure. The microjet underwent Rayleigh breakup\textsuperscript{41} after a few hundred microns, driven by a piezoelectric actuator at 900 kHz (sine wave with 50 V amplitude), producing a single-file train of uniform droplets with a diameter of 5 μm, droplet spacing of 16 μm, and droplet speed of 14 m/s. The droplet diameter $D_{\text{drop}}$, droplet spacing $l_{\text{drop-drop}}$, and droplet speed $v$ were estimated by mass conservation for a single-file droplet train of uniform diameter, namely, $v = Q/A_{\text{jet}}$, $l_{\text{drop-drop}} = v/f$, and $D_{\text{drop}} = 2(Q/4\pi f)^{1/3}$, where $f$ is the droplet frequency determined by the piezoelectric actuator, $Q$ is the in situ
measured volume flow rate of 3.6 $\mu$L/min, and $A_{\text{jet}}$ is the cross-sectional area of the initial liquid jet, determined to be 4.2 $\mu$m$^2$ at given flow rate using a scanning electron microscope. Due to the large vertical focal spot size compared to the droplet spacing, up to 34 droplets were probed simultaneously for each shot; each spectrum was averaged over up to $2 \times 10^6$ shots, resulting in at least 90 000 detected photons. The distance between the dispenser nozzle and the interaction point, which determines the temperature of water droplets, is controlled by a 3-axis motorized sample stage. The temperature was estimated by the center of mass of the fluorescent signal and translated by the monochromator, which only changes the fluence but a reabsorption. Due to its long optical path length as well as coma-free mode operation.\(^{47}\) The incident photon flux was varied by tuning the front end slit placed at the upper stream of the monochromator, which only changes the fluence but affects neither the energy resolution nor the spot size at the sample position. Using this setup, we can thus precisely study the fluence dependence of the O K-edge XE spectra of liquid H$_2$O. In contrast to the free-electron laser measurements described in Sec. II A, the synchrotron measurements at SPring-8 were performed with picosecond x-ray pulses ($\sim$50 ps) at much higher repetition rates\(^{48}\) ($\sim$500 MHz), resulting in that more than one pulse probe the same volume and that much lower fluorences are obtained than at the free-electron laser. Hence, the fluence measurements are very different from the previous study by Schreck \textit{et al.}\(^{37}\) using LCLS to probe valence-hole reabsorption.

D. Simulated XE spectra

The calculations were performed using density functional theory for cluster models containing 32 molecules. Ten structures were extracted as a sampling of the vdw-DF2 \textit{ab initio} molecular dynamics NVE simulation of 64 water molecules performed by Mogelhøj \textit{et al.}\(^{49}\) The central water was computed as either intact or as one of the fragments (O, O$^+$, OH, OH$^+$). Spectra were calculated using the ground state orbitals\(^{50}\) and shifted such that the 1b$''$ peak computed for H$_2$O coincides with the experimental position; spectra for the fragments were shifted by the same amount assuming that the neglected relaxation effects are similar in all cases. Emission energies and transition probabilities were computed with orbital-based density functional theory with the deMon2k code\(^{51}\) and the revised Perdew, Burke, and Ernzerhof\(^{52}\) (revPBE) gradient-corrected functional. The IGLO-III basis set of Kutzelnigg \textit{et al.}\(^{53}\) was used to represent the oxygen of the core-excited molecule. In the other oxygen, the 1s core electrons were replaced by an effective core potential\(^{54}\) and the remaining electrons were described using 4s4p basis functions contracted to 2s2p. All hydrogens were described using 5s2p functions contracted to 2s2p.

III. RESULTS

Fig. 1 shows XE spectra measured at LCLS at non-resonant conditions (550 eV) on the water droplets at three different temperatures corresponding to $\sim$290 K, 226$^\pm_2$ K, and 222$^\pm_3$ K. We notice quite small changes in the relative peak heights of the low-energy 1b$'$ and high-energy 1b$''$ peaks, but the 1b$''$ peak width narrows with decreasing temperature, resulting in a shift of the center of mass of the peak from 526.74 eV at $\sim$290 K to 526.66 eV and 526.63 eV at 226$^\pm_2$ K and 222$^\pm_3$ K, respectively. Because the XE spectrum of crystalline ice\(^{55}\) only shows a prominent 1b$'$ peak with the high-energy 1b$''$ peak reduced to a weak shoulder, we may exclude that a significant volume fraction of the illuminated droplets would have crystallized prior to detection upon supercooling the liquid. We note that the two peaks are less well-resolved in comparison to previous measurements, which can be attributed to different energy resolutions. Fig. 2 shows that the LCLS data at $\sim$290 K can be fitted with relative intensities between the
Based on that the spectra at ~290 K measured at LCLS and at 300 K measured at SPring-8 can be fitted with similar populations of species related to distorted and tetrahedral H-bonds, we can decompose the spectra into relative populations as a function of temperature using the LCLS data at lower temperatures while using the SPring-8 data at higher temperatures. For this decomposition, we assume that the line shapes of the two components are independent of temperature. Fig. 3 shows the temperature dependence of the relative populations of the $1b'_1$ and $1b''_1$ components as obtained under this assumption. It is clear that the fraction of distorted H-bonds is decreasing and the fraction of tetrahedral H-bonds is increasing with decreasing temperature, consistent with previous findings.\textsuperscript{20,28} Furthermore, we also observe the change in the high-energy $1b''_1$ peak position toward lower emission energy (Fig. 1) to be consistent with the previous studies. However, there is a major discrepancy as to what we would expect in terms of intensity of the low-energy $1b''_1$ peak in “no-man’s land” where the liquid is expected to become LDL-like with tetrahedral H-bonds. Analysis of inherent structures from the TIP4P/2005 force field shows that at 230 K the LDL and HDL populations should reach a 1:1 ratio and at temperatures below 230 K the LDL population should dominate.\textsuperscript{24,56} This is supported by experiments in “no-man’s land” where it was found that the tetrahedrality in terms of the peak height of the 2nd shell in the O-O pair correlation function becomes exceptionally strong and mimics low-density amorphous (LDA) ice.\textsuperscript{17} The experimental temperature-dependent trend shows an even stronger increase in tetrahedrality than what is obtained from the TIP4P/2005 simulations.\textsuperscript{17} We would therefore expect that at 226 K a crossing between the $1b'_1$ and $1b''_1$ populations would have occurred with the low-energy $1b'_1$.
peak (interpreted as corresponding to local regions of LDL) becoming dominant. Clearly, this is not observed in Fig. 3.

IV. DISCUSSION

The lack of strong temperature dependence in the peak heights of the $1b_1''$ and $1b_1'''$ peaks opens questions regarding our detailed understanding of the XE process and its connection to the structure of the liquid. The XE energy corresponds to the difference in binding energy of the $1b_1$ and O1$s$ orbitals as could also be measured through photoelectron spectroscopy. We note first that the valence photoelectron spectrum of water only shows one peak in the lone-pair region and that this is too sharp to support two contributions at the separation observed in XES.57–59 Core-level photoelectron spectroscopy of liquid water shows only one O1$s$ peak as well, but in this case the O1$s$ peak is broad enough to support two unresolved features within the separation between the two $1b_1$ peaks in XES.60 Here, we will discuss various hypotheses that can explain the current data. We are thereby revisiting an old discussion,25 but with a focus on that the temperature dependence is somewhat unexpected.

First, could the dissociative mechanism better explain the appearance of the two peaks?19,31,32 This has recently been extensively reviewed25 and excluded based on both high-level theoretical simulations of a model water dimer34,35 and polarization-dependent measurements.27 It was shown that, although the potential energy surface in the core-excited state has its minimum at the accepting oxygen, the proton becomes delocalized between the two water molecules and the resulting O–H stretch has to be described as a superposition of several excited vibrational states, giving rise to important interference effects when the appropriate Kramers-Heisenberg formalism is applied. Indeed, the computed XE spectrum of the water dimer does not exhibit two distinct peaks but rather an asymmetrical $1b_1$ line shape that can also explain the isotope effect.25,34,35 Furthermore, the difference in binding energy between $1b_1$ and O1$s$ in photoelectron spectroscopy is nearly identical for OH and water,26 thus not giving an extra feature in the $1b_1$ region and, in addition, we would expect to see a distinct feature at 523 eV characteristic of a well-defined OH species.25 A dissociative mechanism involving OH and giving a sharp spectral feature in the lone-pair spectrum, but not at 523 eV, thus cannot account for the dependence of the two peaks on temperature,26 polarization,27 and excitation energy.20,25,27,28

A second possibility is that the two peaks observed in XES reflect an additional and different bimodal distribution than that given by distorted and tetrahedral H-bonds which then could have temperature dependence different from that expected for HDL and LDL from molecular dynamics simulations. This seems highly unlikely, however, and there is currently no suggestion from simulations or theoretical considerations of any additional division of structures into classes with the observed weak temperature dependence and which would give rise to the two peaks in XES but be distinct from the suggested species.

A third explanation could be that the split is due to a ground state electronic structure effect in the liquid which depends on the local environment where the $1b_1$ peak is split due to band structure effects as for the 3a$_1$ in ice.57,61 The argument against such an interpretation is that the valence photoelectron spectrum of liquid water should then also show two such peaks in the lone-pair region.57,61 In particular, measurements at relatively low photon energy, where the photoionization cross section is dominated by the O2$p$ contribution,57,61 should give spectra similar to XES which is also completely derived from the O2$p$ character through the dipole selection rule for O K-edge emission. This, however, is not observed.

A 4th hypothesis is that there could be radical species building up in the liquid, not from ultrafast dissociation, but from the Auger decay cascade.62 These could include various fragmented water species, such as the OH radical, that could contribute to the $1b_1''$ peak. This is contradicted by theoretical calculations of the $1b_1$ to O1$s$ emission energy for OH, OH$^+$, O, and O$^+$ as fragments, which all lead to a contribution at the $1b_1''$ position of intact water or at higher energy (Fig. 4). The theoretical results are further supported by experiments shown in Fig. 5 where high-resolution spectra of room temperature water, in this case H$_2$O, were measured at different incident fluence at BL07LSU at SPring-8 to see if the $1b_1'$ could be related to some fragmentation processes; since more fragments would be expected at higher fluence, the $1b_1'$ peak would be expected to be enhanced at higher fluence. In agreement with the calculations, no specific additional intensity grows in the $1b_1'$ region with increasing fluence, indicating that this feature is not related to OH, OH$^+$, O, or O$^+$ fragments. However, there is a small and almost negligible conversion of some $1b_1'$ to $1b_1''$ intensity with increasing fluence that will be discussed below. Although the currently tested radical species are the most probable62 we cannot definitely rule out that there could exist other species not tested here that could give rise to spectral intensity at lower energies. We note that non-linear effects in terms of reabsorption can affect the spectra but only

FIG. 4. Computed XE spectra for H$_2$O (red solid curve) and neutral (solid curves) and positive (dashed curves) fragments of OH and atomic O in a sampling of 32 molecules water clusters (Methods). The same shift has been applied to all spectra such that the $1b_1$ peak position of H$_2$O coincides with the experimental $1b_1''$ peak position. Contributions from the fragments appear in the vicinity of the high-energy $1b_1''$ peak but do not contribute to the $1b_1'$ peak at lower emission energy.
The lower blue curve is the difference spectrum between the normalized 200 nA and 20 nA spectra.

A 5th hypothesis is that there is beam-induced transformation, not in terms of dissociation but in terms of local structural environment. This has been suggested in a recent study of the x-ray absorption spectra of cubic and hexagonal ice in order to explain the variation of the pre-edge intensity in various samples and using different measurement techniques.\(^{63}\) It was proposed that even if the ice is prepared as a pure hexagonal ice phase, the interaction with the beam may transform portions of the ice from hexagonal to high-density amorphous (HDA) ice. It is known that the enthalpy difference between ice I\(_h\) and HDA ice is 0.58 ± 0.1 kcal/mol\(^{64}\) at 1 atm while the absorption of a photon at 550 eV corresponds to 12 680 kcal/mol, which is 4 orders of magnitude higher and thus more than sufficient to convert ice I\(_h\) into HDA ice if the energy is not efficiently transferred away from the excited molecule. We postulate that the absorbed energy will dissipate slowly through the solid in the case of ice, creating many sites where significant energy is dumped and causing a local pressure rise that converts a fraction of the volume into HDA ice. Since the low temperature hinders thermal motions, these molecules are kinetically hindered to convert back to low-density hexagonal ice.

In the liquid state, at ambient temperatures, the large thermal motion and ultrafast dynamics of breaking and forming H-bonds on a picosecond timescale\(^{35}\) allow for a beam-induced structural defect to convert back nearly instantaneously and no beam-induced structural transformation will be seen. This can account for the almost negligible intensity redistribution at room temperature in connection to Fig. 5. However, as water is supercooled below \(T_H\), the relaxation time in the liquid will increase significantly. It has been shown that the structural relaxation time at 220 K in simulations of liquid water using the TIP4P/2005 water model is \(~10\) ns.\(^{66}\) Under a certain flux that transforms the liquid and if the rate to transform back to the structures at equilibrium is slow due to long relaxation time there could build up a steady-state concentration in deeply supercooled water of distorted H-bonds through this deposited energy resulting in converting some of the \(1b_1′\) to \(1b_1′′\) intensity. The effect would become more prominent the colder the liquid becomes through the increasing relaxation time. The challenge to this hypothesis is that the x-ray pulse from LCLS is only around 100 fs in duration and the question is if the liquid would have time to undergo any transformation within this time period. A vibrational period in the H-bond translational mode is around 200 fs,\(^{67,68}\) which is relatively slow in comparison to the pulse length. However, regarding the deposited energy as equivalent to a local heating of the liquid it is most likely the librational mode that will mediate a structural transformation, where the water molecule will rotate out to avoid directional H-bonds. When comparing HDA ice to LDA ice, it is mostly bending of H-bonds that occurs\(^{69}\) and we anticipate similar changes in a conversion of tetrahedrally H-bonded structures to local structures with distorted H-bonds at low temperatures. The vibrational period of the librational mode is of the order of 50 fs and a fractional period of motion could occur on a few femtoseconds to tens of femtoseconds timescale. This could result in a high rate of structural transformation, since the intensity during the LCLS pulse is very high, allowing for many excited molecular centers. Note however, that even if such a conversion would occur in soft x-ray experiments using free-electron lasers, we do not expect this effect to be significant for the recently performed hard x-ray scattering experiment at LCLS\(^{17}\) since at 9.4 keV the absorption cross section is lowered by several orders of magnitude in comparison to just above the O K edge.

Finally, a 6th hypothesis is that a modified fitting scheme of the two components is required, where also the line shape depends on temperature. It is most likely that the local structures with distorted and tetrahedral H-bonds will undergo changes with temperature. This is seen through the high emission energy \(1b_1′′\) peak position changing with temperature\(^{20,28}\) similarly to what is also seen in the XAS temperature-dependent spectra.\(^{28,70}\) We could view the two structural motifs as two different ensembles of molecules so that the line shape of the two peaks could depend on the temperature and not only on the distribution of species within each subensemble. This could occur through the local environment affecting the vibrational lifetime interference causing a change in the degree of asymmetry in the line shape. Let us first demonstrate that this is possible by simply changing the line width of the \(1b_1′\) and \(1b_1′′\) components in the 222 K spectrum as shown in Fig. 6. Since the spectral resolution in the LCLS spectra is lower than in the SPring-8 spectra, the minimum between the \(1b_1′\) and \(1b_1′′\) peaks is not well resolved in the LCLS spectra. This, together with the lower signal-to-noise ratio, allows for a wide range of spectral component widths to be consistent with the data and opens up a possibility that these can indeed undergo a large change with temperature, in particular in the deeply supercooled region; note that the higher resolution and better statistics of the SPring-8 spectra puts stricter limits on this range. Fig. 6 demonstrates that nearly any ratio between the two components can be achieved.
and if the line width would be allowed to change with temperature we can obtain an equal ratio between the two components as we enter into “no-man’s land.” Fig. 7 shows how we can fit the spectra with temperature-dependent line shapes to reconcile the expected nearly equal ratio between structures with tetrahedral and distorted H-bonds in terms of the area of the 1b′/1b′′ intensity ratio as well as the 1b′/1b′′ line width ratio increases monotonically with increasing 1b′ line width.

In this picture, water behaves as a simple liquid at temperatures close to the boiling point, where most interactions are isotropic and dominated by non-directional van der Waals interactions.\textsuperscript{71} As the liquid cools down, the water molecules start to stick to each other through directional H-bonds. This will appear through two classes of configurations,\textsuperscript{18,24,28,29} tetrahedral structures with four strong H-bonds (LDL-like regions) and asymmetric structures with distorted H-bonds (HDL-like regions), with fluctuations between them on some timescale. The first class of structures is enthalpically favored due to efficient H-bonding. Since cooperativity effects make the bonds stronger if water is H-bonded to other waters that are also in tetrahedral structures, fluctuations into tetrahedral structures expand into small local regions that become LDL-like with strong H-bonds, which grow in size since the balance between enthalpy and entropy in the free energy shifts more and more to favor enthalpy with decreasing temperature. There are fluctuations between the LDL-like and HDL-like structures. Although the timescale of these fluctuations slows down as we cool the liquid, the molecules in the asymmetric structures with distorted H-bonds fluctuate more and more frequently into tetrahedral structures with four strong H-bonds, which grow in size since the balance between enthalpy and entropy in the free energy shifts more and more to favor enthalpy with decreasing temperature. There is also a continuous change in the structure of the HDL-like species with temperature. The switching time of OH groups participating in strong or weak/broken H-bonding is furthermore expected to slow down and the amplitude in the motion to decrease with temperature. The HDL-like regions will approach more and more tetrahedral character with strong H-bonds as in high-density ices with many interstitials rather than as in hexagonal ice. Here, we propose that as the HDL-like structures become more symmetrical, similar to HDA ice, the two OH groups in the water molecule encounter more similar environments. This would lead to increased vibrational lifetime interference due to both OH groups contributing in the core-excited electronic state, rather than only one. This
would make the line shape of the high-energy $1b'\prime$ peak more symmetrical, similar to the $1b'\prime$ peak, and also narrower from the smaller thermal motion at lower temperature. As water is supercooled, the correlation length increases\(^3\) indicating that the LDL-like regions become significantly larger. We would therefore assume that there would be different subclasses of tetrahedral molecules depending on the long-range ordering, where some are in the center of the region and others at the boundary. This could lead to variations in the position of the low-energy $1b'\prime$ peak, which would be fully consistent with the fitting shown in Fig. 7.

V. SUMMARY

Though all of the six hypotheses have different strengths and weaknesses, we regard the most likely explanation for the subtle changes in the lone-pair spectral region to be provided by the 6th hypothesis with minor contributions from the other mechanisms. If we assume a plausible line-shape temperature dependence, the ratio between $1b'\prime$ and $1b''$ becomes closer to what we expect, where the now smaller deviation from the expected behavior could come from beam-induced local structural transformations of tetrahedral to distorted H-bonds. Clearly, this requires further investigation in the future utilizing both experiments and theory. Here, we would like to remark that the interpretation of XES of liquid water is indeed less straightforward than previously anticipated and other assumptions, such as temperature-dependent line shapes and potential beam-induced structural transformations of water, may be necessary to explain XE spectra of water upon deep supercooling. To further determine the line shape in the supercooled regime, it would be essential to measure water with superior energy resolution.

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