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ABSTRACT

We present a sub-picosecond resolved investigation of the structural solvent reorganization and geminate recombination dynamics following 400 nm two-photon excitation and photodetachment of a valence p electron from the aqueous atomic solute, $I^-(aq)$. The measurements utilized time-resolved X-ray Absorption Near Edge Structure (TR-XANES) spectroscopy and X-ray Solution Scattering (TR-XSS) at the Linac

Coherent Light Source x-ray free electron laser in a laser pump/x-ray probe experiment. The XANES measurements around the L_1 -edge of the generated nascent iodine atoms (I^0) yield an average electron ejection distance from the iodine parent of 7.4 ± 1.5 Å with an excitation yield of about 1/3 of the 0.1M NaI aqueous solution. The kinetic traces of the XANES measurement are in agreement with a purely diffusion-driven geminate iodine–electron recombination model without the need for a long-lived (I^0 :e⁻) contact pair. Nonequilibrium classical molecular dynamics simulations indicate a delayed response of the caging H₂O solvent shell and this is supported by the structural analysis of the XSS data: We identify a two-step process exhibiting a 0.1 ps delayed solvent shell reorganization time within the tight H-bond network and a 0.3 ps time constant for the mean iodine–oxygen distance changes. The results indicate that most of the reorganization can be explained classically by a transition from a hydrophilic cavity with a well-ordered first solvation shell (hydrogens pointing toward I⁻) to an expanded cavity around I⁰ with a more random orientation of the H₂O molecules in a broadened first solvation shell.

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I. INTRODUCTION

Solvation dynamics and recombination effects in aqueous solution play a crucial role in (photo)chemical reactions such as intermolecular electron transfer. These fundamental processes control the outcome and efficiency of chemical and biological reactions, as they can actively trigger, assist, or hinder key reaction steps.^{1–8} Examples are the transport of ions in water,^{9,10} transport through membranes—where the hydration shell reorganizes in the initial and final stages of the membrane-crossing mechanism¹¹—and the recombination dynamics of the solvated electron, $e^-(aq)$, which is an important species that highly influences chemical reactions^{7,8,12,13} and physical processes¹⁴ in aqueous environments through charge transfer. Consequently, those processes have attracted significant interest and many experimental and theoretical studies have been performed over the past decades.^{15–18}

Solvation and recombination dynamics have been studied with ultrafast optical ultraviolet-visible (UV-VIS) spectroscopy, including transient absorption (TA) and fluorescence techniques as well as infrared (IR) laser measurements since the 1980s.^{1,19-26} Concerning the former, studies have traditionally been carried out on laser-induced dipole moment changes of dye molecules in solution, electronic changes to which the surrounding solvent molecules react via structural reorganization to minimize the free energy. This rearrangement is then measured, down to femtosecond time resolution, by, e.g., fluorescence, excited state absorption, or stimulated emission of the molecule.^{21,27,29} However, in particular, on ultrashort time scales, the distinction between the response of the solvent and that of a molecular solute has been difficult due to the fastest response of the solvent taking place on the same sub-picosecond time scales as the internal relaxation of the solute.³⁰ This issue can be completely removed by studying atomic solutes that change their electronic structure upon photoexcitation. A challenge is that atoms often have high-energetic optical transitions.³⁰ However, a class of systems allowing such investigations are the aqueous halides.³¹ These systems are also considered to be model systems for the initial step of intermolecular charge (e.g., electron) transfer reactions between donor and acceptor systems in solution and can equally be used to study geminate recombination dynamics of the solute-electron system in the absence of any nuclear degree of freedom.

The aqueous halides exhibit intense broad absorption bands below <270 nm (4.6 eV) in the deep UV, not present in the gas phase.^{39,41,42} They correspond to the so-called charge-transfer-tosolvent (CTTS) states, which are bound metastable states of the

solute-solvent system.^{30,39} Laser excitation at energies above 4.6 eV eventually leads to ejection of a valence electron from the halide into the solvent, leaving a neutral halogen atom behind. This happens for both single-photon^{25,26,32-40} and multiphoton^{23,31,38} excitation. The dramatic change in electronic structure of the solute triggers dynamic configurational changes of its surrounding solvent shell.^{30,31,44–46} The dynamic response after single-photon and multiphoton excitation of aqueous halides has been intensively investigated using TA and fluorescence spectroscopy in the visible to near-IR region, 23,25,26,31-40,43,44,47 next to ultrafast liquid phase photoelectron spectroscopy (PES).48,49 In particular, many studies have focused on aqueous iodide, I-(aq), which has the energetically lowest lying CTTS bands of all halides with maxima at 226 and 194 nm. 41,42 Many of these TA experiments, which detected the transient changes in the absorption band of the appearing solvated electron in the visible to near-IR range,⁵⁰ mainly concentrated on the iodine-electron geminate recombination dynamics^{23,25,26,31–34,36–40,44} as the observation of pure solvation dynamics around the nascent atom with this method is challenging: Optical methods have high sensitivity toward changes in both electronic and vibronic energy levels, but they are not directly sensitive to structural changes.4

The experiments performed after photoexcitation into the energetically lowest lying CTTS state of I⁻(aq) concluded that a localized bound state between the halogen atom and the electron, a so-called contact or caged pair, is formed on ultrafast time ^{6,32,33,37–39} This is supported by quantum molecular dynamscales.² ics (MD) simulation results reported by Sheu and Rossky⁵¹ and Staib and Borgis,^{54–56} who have shown that CTTS excitation produces a localized state. This is bound not by Coulomb attraction between the electron and halide nucleus but by the polarization of the solvent surrounding the anion.^{57,58} The ejected electron remains close to its geminate (original) partner in a stable contact or close pair that is bound by several k_BT . The model implies that for the halides, after formation of this contact or caged pair on a time scale of several hundred femtoseconds and thermalization within a few picoseconds,²⁶ the subsequent kinetics are dominated by two competing processes: The caged electron escapes into the solvent or nonadiabatically recombines with the parent atom to the ground state halide. Both processes occur on a time scale of tens of picoseconds.³² This is distinctly different from the geminate recombination dynamics observed, e.g., after low-energy photoexcitation of polyatomic anions, which is well described by a pure diffusionlimited encounter of the photoseparated species and without the need for a caged pair.^{37,39} Energy-dependent TA measurements for

higher photoexcitation energies [including 389 nm two-photon excitation (6.4 eV)] concluded that one main difference from the lowenergy excitation is an altered initial electron ejection distribution, i.e., where average ejection distances increase stepwise from <5 Å for 4.9 eV to ~15 Å for 7.6 eV, while even higher energies above 8.2 eV yield much larger average ejection distances >40 Å.^{38,39} This difference is explained by the transition from CTTS-driven ejection for the lower energies to direct ejection into the conduction band of water for higher energies.^{38,39} While these measurements extracted important kinetic information on the >5 ps time scales (up to hundreds of picoseconds), a robust description of the early recombination dynamics remained challenging as such TA experiments need to include a description of the observed dynamic Stokes shift of the absorption spectrum of the solvated electron within the first picosecond. 23,25,26,31-40 This obscures a clear identification of the earliest recombination processes. Similar difficulties arise for the PES signal of the solvated electron, which overlaps that of the departing electron, as well as the signal of the subsequent dynamics.³⁰ In addition, especially for high-energetic/multiphoton excitation conditions, undesired water solvent ionization can occur,⁵⁹ also generating solvated electrons, and disentangling these contributions from those of the halide detachment process is nontrivial.

The experimental challenges are often sought to be overcome by coupling the spectroscopic observables to modeling studies (both for dynamic solvation and recombination). These have included classical, semiclassical, and quantum/ab initio MD simulations^{3,51,53-56,60-64} on cluster models of solvent molecules, but they often lack a direct connection to (structural) observables, complicating direct validation of the model predictions. Regarding the dynamic recombination, such simulations have suggested the formation of an (I⁰:e⁻) contact or close pair⁵¹⁻⁵³ on an ultrafast time scale after initial photoexcitation into the lowest-lying CTTS state of aqueous I⁻, thus guiding the interpretation of the results from, e.g., TA experiments. Concerning investigations of the local solvent structure, MD simulations have extracted a large distribution of rather different solvent cage configurations.⁶⁵ However, for the water molecules in the first solvation shell around I⁻(aq), one of the hydrogens of these nearest water molecules has been concluded to point toward the I⁻ due to electrostatic interactions.⁶ Around $I^{0}(aq)$, the water molecules lose this well-defined orientation around the solute, a process driven by entropy cost and mutual repulsion of the closest hydrogens.^{65–67} As a result of the changed molecular orientations of the solvent cavity, it is also possible that I⁰ receives a partial charge back donation from the oxygen lone pairs of the polar H₂O molecules. Some of these simulation studies have also suggested the formation of a transient I⁰(OH₂) complex within the first picosecond after electron abstraction.

Experimentally, new information on the solvation and recombination changes following photoabstraction has been opened up by using x rays.^{31,44,45} These approaches (i) allow to probe the generation and survival of the nascent iodine atom itself, instead of focusing on the kinetic behavior of the solvated electrons alone, and (ii) enable structural sensitivity. Time-resolved X-ray Absorption Near Edge Structure (TR-XANES) spectroscopy experiments at the I L₁-and L₃-edges have demonstrated the potential of this approach.^{43–45} The 2s-5p L₁ pre-edge transition (at 5.184 keV) serves as a new observable for the electron detachment step while picosecond Extended X-ray Absorption Fine Structure (EXAFS)

spectroscopy at the I L₃ absorption edge identified a change in the solvent shell right after photoexcitation.⁴³⁻⁴⁵ However, these pioneering studies exploiting time-sliced femtosecond x rays from a synchrotron lacked the statistical significance and/or time resolution to reliably track the earliest dynamic changes at scales ranging from sub-picoseconds to a few picoseconds.³⁰ Expanding on this approach, here we present results from the much more intense Linac Coherent Light Source (LCLS) X-ray Free-Electron Laser (XFEL) using its hard x-ray beam as an element-specific and structuresensitive probe. For this purpose, we exploit a combination of femtosecond time-resolved X-ray Solution Scattering (TR-XSS)6and TR-XANES.7 ⁸⁰ The TR-XSS measurements are directly sensitive to changes in the solute-solvent distances⁸¹ and are analyzed using structural information from equilibrium and nonequilibrium classical MD simulations of aqueous I^- and I^0 to extract a detailed picture of the solvation dynamic structural changes, while TR-XANES is used to track the early geminate iodine-electron recombination dynamics on a <10 ps time scale by monitoring the 2s-5p pre-edge feature at the I L₁ absorption Combining these probes allows us for the first time to edge.4 elucidate both processes on these ultrafast time scales.

The 3.1 eV pump photon energy (400 nm) in this study is below the CTTS states of I⁻ (>4.6 eV or <270 nm) and necessitates a multiphoton (2- or 3-photon) excitation process.44 The required vertical energy to directly excite the valence iodide 5p electron into the water conduction band is 7.7 eV.^{39,82,83} In a 400 nm twophoton excitation process, an energy of 6.2 eV is absorbed, which lies well below this threshold but within the CTTS manifold. Thus, two-photon absorption will not result in direct promotion of the electron in the water conduction band. It is also unlikely that a two-photon absorption process from the 5p orbital populates the lowest-lying s-like CTTS state, from which the electron then separates adiabatically^{25,51-53} as for one-photon excitation at, e.g., 255 nm. Instead, it is assumed that two-photon absorption accesses higher-lying p-type CTTS states⁵⁸ and electron detachment becomes already possible through nonadiabatic coupling to the water conduction band.^{38,39} Three-photon absorption processes (9.3 eV) would allow direct ionization into the water conduction band with considerably larger ejection distances than for two-photon excitation,38,39 and we will use this information to discriminate between both processes when analyzing the iodine-electron recombination kinetics.

II. EXPERIMENTAL APPROACH

The dynamics following photo-oxidation of aqueous I⁻ was tracked with time-resolved pump-probe XANES and XSS experiments at the X-ray Pump-Probe (XPP) beamline of the XFEL facility LCLS. The general experimental setup has been reported elsewhere^{70,72-75} and details concerning the present setup and data processing schemes are described below and in the supplementary material. Briefly, the sample consists of a 0.1 mm thick flat sheet liquid jet containing 0.1M aqueous iodide (by dissolving NaI in water). The laser and x-ray beams were temporally and spatially overlapped on the sample, with an x-ray spot size of ~60 μ m (fwhm) at the sample position and a laser spot size of 160 × 220 μ m². The solvated iodide was excited by a ~50 fs (fwhm) laser (pump) pulse at 400 nm (316 μ J pulse energy) and probed at selected time delays with a ~50 fs (fwhm) x-ray (probe) pulse.

The XSS measurements were performed using the unmonochromatized SASE beam (~ 10¹² photons per pulse) and were acquired in a sequence consisting of three laser-on shots (=laser-excited sample) followed by one laser-off shot (=static sample) to generate a transient signal $\Delta S = S_{on} - S_{off}$ with several thousand such sequences per time point. The XSS signal was recorded with the liquid jet perpendicular to the beam propagation direction and detected in the forward direction by the 2D CS-PAD detector.⁸⁴ Detector corrections were applied as described in Ref. 85. Following these corrections, the 2D difference images were azimuthally integrated to yield one-dimensional $\Delta S(Q, t)$ difference scattering signals, with Q the scattering vector defined by $Q = \frac{4\pi}{\lambda} \sin(2\theta/2)$, where 2θ is the scattering angle onto the detector and λ is the x-ray wavelength.

For the XANES measurements, the incident pink x-ray beam was monochromatized to $\delta E \sim 1$ eV around the I L₁ edge (at E = 5.188 keV) using the XPP fixed exit Si-111 double-crystal monochromator (DCM). The liquid jet was horizontally tilted to 45° with respect to the incoming x-ray beam. This allowed recording of the XANES spectra in total fluorescence yield (TFY) mode by an x-ray diode placed at 90° to the beam propagation direction and in the polarization plane to suppress contributions from elastically scattered x rays. The x-ray probe pulses used for the XANES spectra were detected in a sequence where three laser-on shots (=laser-excited sample) were followed by one laser-off shot (=static sample) to generate a transient signal $\Delta A = A_{on} - A_{off}$. The timedelay traces (measured at 5.184 keV) were generated by subtracting the signal at negative time delays from the measured transient signal with laser on (for details, see the supplementary material, Sec. III B). Using the timing tool at LCLS,⁸⁶ all the sets of measurements obtained were temporally re-binned into 20 fs bins (both for the XSS and XANES measurements).

III. COMPUTATIONAL APPROACH

As a starting point for the interpretation of the x-ray scattering data, the equilibrium structures of the solvent around the I⁻ and I⁰ species were modeled by two equilibrium MD simulations with I^- and I^0 each solvated in a cubic box (30 Å side length) of water molecules. The TIP4P-Ew potential was used for the water molecules. For I⁻, the OPLS-AA-consistent Lennard-Jones parameters used were $\sigma = 4.81$ Å, $\epsilon = 0.71$ kcal/mol with a partial charge of $-1.^{87}$ There are no atomic I⁰ Lennard-Jones parameters within the OPLS-AA force field, but since its iodobenzene-I parameterization has a partial charge of only 0.1 e, the iodobenzene-I Lennard-Jones parameters ($\sigma = 3.75$ Å, $\epsilon = 0.60$ kcal/mol) were chosen as an approximation of atomic I⁰, using a partial charge of $0.^{88}$ The I^{-}/I^{0} atoms were restrained at the center of the box and a 24 ps T = 300 K equilibration run (Berendsen thermostat⁸⁹) was performed. For the production run, MD trajectories were calculated with a multistep integrator⁹⁰ with the nonbonded-near and nonbonded-far (long-range electrostatic) interactions updated every 1 and 3 fs, respectively, and using a Nose-Hoover thermostat at 300 K. Radial Distribution Functions (RDFs) g(r) were sampled in 0.1 Å radial bins and over 2000 individual simulation time steps from the total trajectory length of 2 ns. To model the excitation, we follow the approach of previous studies^{44,46} and instantaneously remove the charge from iodide and run nonequilibrium MD simulations. We want to point out that in contrast to some mixed quantum–classical work, our MD simulations do not include the solvated electron^{91–93} as the difference scattering signal is dominated by the change in the electron-rich solute and its nearest surroundings. The nonequilibrium MD simulations were initiated from 200 starting configurations selected at 10 ps intervals from an I⁻ equilibrium trajectory. For these, the charge and the Lennard-Jones parameters were changed from the I⁻ to the I⁰ values at t = 0 and the 200 trajectories were propagated for 5 ps each.

IV. RESULTS

A. Molecular dynamics (MD) simulations

Figures 1(a) and 1(b) show the I–O and I–H RDFs of I⁻ and I⁰ solvated in water as obtained from the equilibrium MD simulations. A significant difference is observed in the solvation shells around the ions vs the atoms, with the negatively charged I⁻ exhibiting a more well-defined solvation shell configuration (narrower peaks, three distinguishable shells with the first and second shells containing ~8 and ~22 oxygens, respectively). For I⁻, the hydrogen atoms of the water molecules in the nearest solvation shell are oriented



FIG. 1. Top: Schematic model showing the structural dynamics of the H₂O solvent shell during the I⁻ \rightarrow 1⁰ photoreaction after pumping with an ultrashort 400 nm laser pulse. Upon photodetachment of the electron from I⁻, the first solvation shell of the water molecules undergoes a reorganization. Bottom: Results from equilibrium MD simulations of aqueous I⁻ and I⁰. The I–O (a) and I–H (b) radial distribution functions (RDFs) show how the solvation shell around I⁻ (blue) is more well-ordered (narrower peaks) with the hydrogens being significantly closer to I.

toward the solute as evidenced by the first peak in the I–H RDFs being 1.0 Å closer to I⁻ than the one for I–O. The solvation shells of the neutral I⁰ are less ordered (broader peaks, two distinguishable shells with the first shell containing 22 oxygens) with more similar I–O and I–H RDFs. Compared to the case of I⁻, the nearest water molecules no longer have a well-defined orientation with respect to I⁰. From the viewpoint of the solute, an expansion of the solvation cage is observed as the first peak in the I–H and I–O RDFs moves outward with ~0.9 Å (from 2.65 to 3.55 Å) and 0.1 Å (from 3.65 to 3.75 Å), respectively.

Figure 2 shows the results obtained from the nonequilibrium MD simulations. Figure 2(a) shows the $g_{IO}(r, t)$ and $g_{IH}(r, t)$ RDFs as a function of time for the first 3 ps of the 5 ps nonequilibrium trajectory. For both RDFs, the dynamics take place on a <0.5 ps time scale, and for $g_{10}(r)$, a lowering and slight symmetric broadening followed by a slower broadening toward longer distances is observed, whereas for $g_{\rm IH}(r)$, the first peak exhibits an almost 1 Å shift to longer distances, again combined with a lowering of the peak height. Showing this in more detail, Fig. 2(b) shows $g_{10}(r)$ for four selected time delays. The peak height is observed to have reached its new equilibrium value after <100 fs with some symmetric broadening on the same time scale, whereas the shift toward longer distances and the formation of a new distinct solvation structure (characterized by broadening of the first peak and a peak turning into a dip at r = 5.2 Å) develops on a time scale of hundreds of femtoseconds. Figure 2(c) highlights these dynamics by showing the magnitude of $g_{10}(r, t)$ at the peak and on the long-*r* shoulder of the

peak, indicating time scales of 50–75 fs for the peak lowering and ~300 fs for the peak shift. Figure 2(d) shows the short-time part of $g_{IO}(r, t)$, with gray lines indicating the traces in Figs. 2(b) and 2(c). Similar plots are shown for $g_{IH}(r, t)$ in the supplementary material, Fig. S1.

To further investigate the structural dynamics indicated by the two simulated g(r,t) shown in Fig. 2(a), Fig. 3 shows the results of a Singular Value Decomposition (SVD) of $g_{IO}(r,t)$. For this, the $g_{IO}(r,t)$ matrix was decomposed as the matrix product $g = USV^T$, where the left-singular vectors (columns of *U*) represent the typical "shapes" describing how g(r) evolves and where the right-singular vectors (columns of *V*) represent the time evolution of each component in this time evolution. The matrix *S* is diagonal, with the elements sorted in descending order and describing the relative magnitude of the paired components in *U* and *V* in terms of the contribution to the total signal.

In addition to the main component of $g_{IO}(r, t)$ [dark blue traces in Figs. 3(a) and 3(c), representing the steady-state g(r) for I^0], Fig. 3(b) shows the magnitudes [$S_{i,i}$, normalized to S(1, 1), which is omitted for clarity] of the individual components of the SVD of g(r, t) and indicates how only two components with magnitudes above the background serve to completely describe the evolution of g(r, t). Inspecting Fig. 3(b) [and noting that the sign of the traces in Fig. 3(a) should be inverted to show the evolution from g_{I^-} to g_{I^0}], the first of these components (light blue) is observed to describe how the initial g(r, t = 0) evolves toward the steady-state g(r) via a peak shift+broadening toward longer distances and the evolution



FIG. 2. (a) Time-resolved radial distribution functions (RDFs) $g_{|0}(r,t)$ and $g_{|H}(r,t)$ from the nonequilibrium MD simulations of the solvent dynamics following the |- photoabstraction process. (b) $g_{|0}(r)$ for four time delays, showing the details of the I–O structural dynamics with peak lowering and symmetric broadening followed by a shift toward longer distances. (c) Time evolution of $g_{|0}(r,t)$ magnitude at the first peak and long-*r* shoulder. (d) Short-time part of $g_{|0}(r,t)$ with horizontal gray lines indicating the data traces in panel (b) and vertical lines the traces in (c).



FIG. 3. Summarized results of a Singular Value Decomposition (SVD) of $g_{|0}(r, t)$. (a) The three first left-singular vectors (columns of U) describing the shape (dark blue) and change of shape of $g_{|0}(r, t)$ due to the electron abstraction. (b) Normalized magnitude of the diagonal entries in S with S(1, 1) omitted for clarity, showing that the evolution of $g_{|0}(r, t)$ is well described by three components only. (c) Right-singular vectors (columns of V) showing the time evolution of the signal components shown in (a).

of a pronounced dip in g(r) at $r \approx 5.5$ Å. Figure 3(a) also shows the second significant contribution (teal) to the changes in g(r), highlighting the pronounced lowering and slight symmetric broadening of the first-shell peak in g(r) discussed above. Figure 3(c) shows the time evolution for these two components, with the peak height decrease and slight symmetric broadening described by U^3 dominated by prompt grow-in (<100 fs), followed by further increase on a 200–300 fs time scale as shown by the time evolution described by V^3 . The overall shift toward longer distances of the first solvation shell (described by U^2 , light blue) exhibits a short wait time, <100 fs, followed by evolution on a 200–300 fs time scale as seen by the V^2 graph in Fig. 3(c) (light blue). These observations are further discussed in relation to the experimental results described below, and the corresponding results for $g_{IH}(r, t)$ are shown in the supplementary material, Fig. S2.

B. Femtosecond x-ray solution scattering (XSS)

Figure 4(a) shows the time-resolved XSS difference signals re-binned in time intervals of 50 fs from -1 to 7 ps after laser excitation as radially integrated $\Delta S(Q,t)$ curves. The difference signals arise mainly from two contributions, the local changes in the solvent molecules in the proximity of the solute (ΔS_{cage}) and structural changes in the bulk solvent structure due to heating ($\Delta S_{solvent}$),

$$\Delta S_{total}(Q, t) = \Delta S_{cage}(Q, t) + \Delta S_{solvent}(Q, t).$$
(1)

Qualitatively, $\Delta S_{solvent}(Q, t)$ arises from structural changes in the bulk solvent, which are due to changes in temperature and density of the bulk solvent following photoexcitation of the sample. In the case of bulk water on a sub-10 ps time scale (before any significant thermal expansion occurs), this solvent term has been shown to be generally well described by a difference signal $\Delta T(t) (\partial S(Q)/\partial T)|_{\rho}$, linear in temperature, which arises from the molecular rearrangement solely due to an increase in solvent temperature, ΔT .⁹⁴ This term was measured in a separate experiment on neat water, following the procedure outlined in the work of Kjær *et al.*⁶⁹ In line with previous analysis of LCLS experiments utilizing high excitation power,⁷² the nonlinear changes in scattering due to significant local heating are included in the model as the $\epsilon(t)S_{dT^2}(Q)$ term, where $\epsilon(t)$ is a free time-dependent parameter and $S_{dT^2}(Q)$ is calculated from the differences between simulated



FIG. 4. Time-resolved X-ray Solution Scattering (TR-XSS) difference signals and fit results. (a) TR-XSS signals as a function of time after laser excitation. (b) The residual after subtracting the structural model in Eq. (2). (c) A comparison of the TR-XSS signal (gray) at 2 ps, the structural model (black) and the residuals (bottom, black). The structural model (middle) consists of contributions from the water heating (red and purple) and a change in the I⁰ solvent cage estimated from the equilibrium MD simulations (blue).

scattering signals from equilibrium MD simulations of neat water at a range of temperatures. 72

To simulate the difference scattering signal arising from structural transformations in the solvent shell, $\Delta S_{cage}(Q,t)$, the steady-state x-ray scattering patterns for both aqueous I⁰ ($S_{I^0}(Q)$) and I⁻ ($S_{I^-}(Q)$) were calculated⁹⁵ using the pairwise RDFs from the two equilibrium MD simulations discussed above as input. The solvent cage term in our model is thus estimated as the difference of those equilibrium scattering signals S(Q) scaled with a time-dependent factor, $\Delta S_{cage}(Q, t) = \alpha(t)(S_{I^0}(Q) - S_{I^-}(Q))$. From these considerations and suppressing the Q-dependence for clarity of presentation, the total difference scattering signal in Eq. (1) is calculated through the expression

$$\Delta S_{total}(t) = \alpha(t)(S_{I^0} - S_{I^-}) + \Delta T(t)(\partial S/\partial T)|_{\rho} + \epsilon(t)S_{dT^2}, \quad (2)$$

thus rendering a full model with three parameters (α , ΔT , ϵ) for each time delay, where α denotes the relative completeness of the full solvent shell reorganization.

Figure 4 shows the difference scattering signals acquired at all investigated time delays (a) and the residuals (b) after modeling the signal with a combination of the three terms in Eq. (2). Figure 4(c) shows a representative example of the model fit to the data at 2 ps in more detail. Remarkably good agreement is observed from the residuals, supporting that the XSS signal is indeed monitoring the dynamics of the formation of the new solvent shell. Figure 8 shows the time evolution of the magnitude of the model components.

To further support this simple three-component analysis approach, a second, independent analysis was performed where the XSS difference signal from an experiment utilizing direct threephoton excitation of water in the same experimental setup was used. Within this scheme, the neat water heating signal was scaled to and subtracted from the XSS signals from the I⁻ experiment to remove the solvent-heating contribution. Figure 5(a) shows the difference signal following this solvent-subtraction, indicating the presence of a difference signal component that appears shortly after t_0 and then grows in magnitude over the next few hundreds of femtoseconds. Figures 5(b) and 5(c) show the result of a SVD of this solvent-subtracted residual difference signal, with panel (b) showing the two dominating left-singular vectors directly compared to the difference signal from bulk water heating (red curve/long dashes) and to the cage signal calculated from the equilibrium MD simulations (blue curve/short dashes) and with the corresponding time evolution (right-singular vectors) in panel (c). The high degree of similarity between the dominating SVD component of the solvent-subtracted difference signal (blue) and the difference signal calculated from equilibrium MD simulations before and after electron abstraction supports the use of this component in our data modeling and interpretation approach.

C. Femtosecond L₁ x-ray absorption near edge structure (XANES) spectroscopy

The time-resolved XANES spectrum around the iodide L_1 -edge detects both the appearance of the neutral I⁰ atom (or more precisely the opening of the 2s-5p pre-edge absorption channel) next to the absorption edge shift between the iodide anion (with its fully closed 5p shell) and the nascent iodine atom. The top of Fig. 6 shows the



FIG. 5. (a) Solvent-subtracted (i.e., cage-only) TR-XSS data from aqueous iodide excited at 400 nm calculated by subtraction of the solvent-only heating contribution [modeling the solvent term in Eq. (1)] identified in a reference experiment on neat water. (b) First two components from an SVD analysis of the data in panel (a). From comparison with the model components shown in Fig. 4, these can be identified as a difference signal arising primarily from solvent cage dynamics (blue) and the bulk solvent heating (red). (c) Time evolution of the two SVD components, indicating a clear onset and grow-in of component 1, with the time evolution in component 2 surmised to arise from over-subtraction of the water heating signal [i.e., solvent term in Eq. (1)].

spectrum of the laser-excited sample (after 200 fs) together with the ground state iodide spectrum. The bottom of Fig. 6 depicts a collection of transient XANES measurements (laser-excited minus unexcited XANES) for selected time delays. The transient spectra exhibit mainly a positive and a negative feature, near 5.184 and 5.192 keV, respectively. The latter is caused by a (blue) shift of the iodide L_1 absorption edge to higher energies. This is expected for the removal of one 5p electron after laser excitation, which slightly



FIG. 6. Time-resolved XANES at the I L₁-edge. Top: X-ray absorption probe spectrum with (black triangles) and without (gray squares) the laser pump pulse. After laser excitation, a distinct pre-edge feature appears at 5.184 keV due to the created vacancy in the 5p orbital, which then opens up the new 2s-5p transition. The change in oxidation state (from -1 for $|^-$, to 0 for $|^0$) causes an absorption edge shift (starting around 5.188 keV) toward higher energies. Bottom: Transient XANES at selected time delays, showing the prompt appearance and slower decay of both transient features, the positive transient related to the 2s-5p transition at 5.184 keV and the negative transient related to the absorption edge blue shift at 5.192 keV.

reduces the shielding of the 2s orbital for the iodine nucleus, yielding a stronger binding energy.^{44,45} The positive transient feature at 5.184 keV arises from a new bound-bound transition: Having a closed-shell 5p⁶ electron configuration, I⁻ cannot exhibit the 2s-5p transition, but excitation of one 5p electron opens up the allowed 2s-5p transition at 5.184 keV.^{44,45} In addition, this 2s-5p transition occurs at slightly different energies for the chemical species I₂ and I₃, being thus sensitive to the different iodine-related species.^{43,44}

Earlier picosecond time-resolved XANES experiments at the I L1-edge under similar pump laser conditions and identical reactant concentration (100 mM) found mainly I⁰ and the solvated electron as photoproducts after 50 ps,⁴⁴ which is in agreement with diffusion-driven time scales for generating subsequent products, e.g., I_2^- or I_3^- . Consequently, on shorter time scales, the 2s-5p transition provides a quantitative measure of the concentration of generated iodine atoms. This is important as the excitation yield cannot be independently derived from laser-only measurements, which are only sensitive to solvated electrons and cannot distinguish between solute and solvent generated electrons. In this work, we thus extract the iodide excitation yield by comparing the oscillator strength of the iodine 2s-5p transition after ~200 fs with the related 2s-5p transition strengths measured for solid I2 and solvated $I_3^-(aq)$ (for details, see the supplementary material, Sec. IV and Fig. S12). The excitation yield right after photoexcitation determined by this procedure is 32(4)%, or in other words, 32(4) mM aqueous I^0 with 68(4) mM aqueous I⁻ are in the 0.1M solution right after excitation.

D. Kinetics from femtosecond x-ray absorption near edge structure (XANES) spectroscopy

Figure 7 shows the temporal evolution of the 2s-5p transition as a signature of the electron removal from the filled 5p orbital of I⁻ (for data analysis and reduction, see the supplementary material, Sec. III). One observes a fast rise around time zero with an Instrument Response Function (IRF) of 110(7) fs (inset in Fig. 7), which is slightly smaller than the expected time resolution due to the velocity mismatch between 400 nm optical and ~5.2 keV x-ray pulses through a 0.14 mm thick sample (the 0.1 mm flat sheet jet rotated by 45°), while the expected pulse-width governed IRF amounts to about 70 fs (for 50 fs fwhm laser and x-ray pulse widths). This hints toward the nonlinear excitation of the sample, which dominantly occurs within the first parts of the irradiated volume, thus effectively reducing the sample thickness of the excited state species. Right after signal buildup governed by the IRF, we observe a signal decrease within the first picosecond, which flattens out toward 10 ps. We compared our experimental kinetic traces to the theoretical predictions of three distinctly different models, which had been previously used to describe the geminate recombination of photo-generated electrons in TA experiments on halides in



FIG. 7. Kinetic traces of the iodine 2s-5p transition intensity together with three different theoretical models describing the geminate iodine-electron recombination dynamics: Model (a), assuming only the contact pair as primary photoproduct (10 $(:e^{-})^{2}$ (blue curve), model (b)³⁷ ⁶ (green curve) based on a semianalytical theory for diffusion-controlled reactions in a potential well⁹ (which is a resemblance of the contact pair idea), and model (c) using only diffusionlimited recombination of photo-generated geminate pairs (red curve)² with a best-fitted average electron ejection distance of $\langle r_0 \rangle = 7.4 \pm 1.5$ Å and literature values for (i) the combined electron and iodine diffusion constant, $D' = 5.6 \cdot 10^{-4} \text{ Å}^2/\text{fs}$ at room temperature,³⁶ and (ii) a fixed reaction radius $r_{xn} = 5 \text{ Å}.^{36}$ Hereby, one dataset containing time points up to 10 ps is averaged at 10 ps and plotted together with the average from a series of datasets covering the first picosecond (details are in the supplementary material, Sec. III C and Fig. S6).

aqueous solution^{26,32,33,36–39,43,44} (Fig. 7). This treatment is valid since our time delays of $\Delta t \leq 10$ ps allow the iodine atom and the electron as the only two species generated in solution.⁴⁴ We denote these models here as (a) the (I⁰:e⁻) contact pair model^{26,32,33,43,44} in conjunction with MD simulations^{51–56} (blue curve), (b) the diffusion in a potential well model,^{37–39,96} which is based on a semi-analytical theory for diffusion-controlled reactions in a potential well developed by Shushin^{97–99} (this includes the contact or caged pair idea) (green curve), and (c) the straightforward diffusion model following electron ejection^{26,32,33} based on Refs. 100 and 101 (red curve). Details of all these models including their mathematical description and extracted parameters can be found in the supplementary material, Sec. III D. Theity of each of the models (a)–(c) was identified via their minimized χ^2 -value in comparison to those of our I 2s-5p kinetic traces, i.e., (a) $\chi^2 = 3.43$, (b) $\chi^2 = 2.82$, and (c) $\chi^2 = 1.51$, which thus favors model (c) over the other two models as the most accurate description of the measured kinetics.

Briefly, the contact pair model (a) implies the formation of an electron-iodine contact pair directly after CTTS photoexcitation with a rate k_p , followed by electron escape into solution by overcoming the free energy barrier with a rate k_d or nonadiabatic recombination to form the ground state halide with a rate k_n .^{26,32,33,43,44} The iodine atom survival kinetics using the published rate constants after 400 nm multiphoton excitation ($k_p^{-1} = 200$ fs, $k_d^{-1} = 59.4 \pm 6.7$ ps and $k_n^{-1} = 97.9 \pm 7.2$ ps; see the supplementary material of Ref. 44) does not match the data, which decays faster than this model predicts (Fig. 7, blue curve).

Model (b) relies on a semi-analytical theory for diffusioncontrolled reactions in the presence of a potential well. $^{97-99,102}$ It expands on the previous contact pair model, viewing the photoexcited system as two states: (1) one with the iodine atom and electron bound in a potential well [U(r), r] is the distance between the geminate pair] due to the attractive interaction between the geminate pair ("bound state," arising from the polarization of the halide atom by the electron, resembling the idea of a contact or close pair) and (2) the other state with both partners outside the range of the potential well.^{38,39} The size of the potential well is characterized by the Onsager radius, a, at which $U(r) = -k_B T$. The former state is described by the population of the geminate pair inside the well and the latter is represented by the geminate pair spatial distribution function.³⁹ The two states interact with one another at the boundary defined by the Onsager radius. The only way the population of the geminate pair decays is via passage to the "bound" state, where it can recombine with a rate W_r to reform iodide.39 Alternatively, it can escape from the potential well with a dissociation rate W_d . The calculated iodine atom survival kinetics using the published model parameters applying to 389 nm two-photon excitation (6.4 eV) of aqueous $I^{-38,39}$ (Fig. 7, green curve) $(W^{-1} = 14 \text{ ps and } p_d = 0.216 \text{ with } W = W_r + W_d \text{ and } W_r + W_r +$ $p_d = W_d/W$; for details including all model parameters, see the supplementary material, Sec. III D 2) does not match our experimental data, which again decays faster than the model prediction. Note that this model and also model (c) (see discussion below) rely on the mutual diffusion coefficient of the iodine-electron pair and assume thermalization is complete for the nascent species.⁹⁶ This does not strictly apply within the first picosecond, and we will come back to the uncertainties associated with this treatment in Sec. V.

Model (c) is a pure diffusion-limited geminate recombination model following photoejection into the solvent without assuming any contact or caged pair formation.^{26,32,33} This model shows the best agreement with our data, especially within the first picosecond (Fig. 7, red trace). The main model fit parameter is the average ejection distance of the excited 5p electrons ($\langle r_0 \rangle$, which is excessenergy dependent). Using this straightforward diffusion approach, the separated iodine and electron are promptly created after photoexcitation with a mutual distance r_0 ,⁵⁹ assuming thermalization is complete for the nascent species.³² This separated pair then undergoes three-dimensional diffusion in the solvent without any interparticle potential or any external force.³² When both species re-encounter each other at a certain reaction radius, r_{xn} , recombination occurs promptly.⁵⁹

In this model, the survival probability Ω of the electron against geminate recombination (which is identical to the iodine survival probability in our experiments) is given by³²

$$\Omega(r_0, t) = 1 - \frac{r_{xn}}{r_0} \cdot \operatorname{erfc}\left(\frac{r_0 - r_{xn}}{\sqrt{4D't}}\right),\tag{3}$$

where D' is the sum of the diffusion coefficients of the involved species (here, I^0 and e^-), t is the elapsed time after initial creation of the recombination partners, and erfc(z) is the complementary error function [= 1 - erf(z)]. The recombination radius r_{xn} is connected to D' and the bimolecular rate constant k for a diffusion-limited reaction via $k = 4\pi D' r_{xn}$. Fixing D' and r_{xn} to literature values,³⁶ we are left with only one adjustable parameter: the initial average electron ejection distance $\langle r_0 \rangle$ for a certain assumed distribution function of the initial distances. Although photoionization results in a relatively narrow distribution of initial kinetic energies of electrons, the ensemble of ejected electrons will exhibit a relatively broad distribution of distances.³² Thus, a distribution function of initial distances instead of a single distance value, r_0 , is required. For the ensemble of ejected electrons, both exponential and Gaussian distributions are readily used, in particular for the description of their observed kinetics in TA experiments.^{26,32,33,59} These imply initial ballistic and diffusive formation of the pair, respectively. The survival probability is then obtained by convolution of the selected distribution function with Eq. (3). For an exponential distribution of distances, this yields the following integral for the survival probability:32

$$\Omega(t) = \int_{r_{xn}}^{\infty} \frac{e^{-r_0/b}}{8\pi b^3} \cdot \Omega(r_0, t) \cdot 4\pi r_0^2 \, dr_0, \tag{4}$$

b is defined via the average radius $\langle r_0 \rangle = 3b$, and the root-meansquared radius is $\sqrt{\langle r_0^2 \rangle} = 2 \cdot \sqrt{3} \cdot b$.³² The survival probabilities for the different electron ejection distributions with identical $\langle r_0 \rangle$ are actually quite similar³² (for details, see the supplementary material, Fig. S9). The entire time trace data in Fig. 7 is then fitted by a convolution of Eq. (4) with the Gaussian-shaped instrument response function,

$$f_{2s-5p}(t) = \text{IRF}(\sigma = 47 \text{ fs}) * \begin{cases} 0, & t \le t_0, \\ \Omega(t), & t > t_0. \end{cases}$$
(5)

The red curve in Fig. 7 shows the best fit to our data using fixed parameters $D' = 5.6 \cdot 10^{-4} \text{ Å}^2/\text{fs}$ (room temperature) and $r_{xn} = 5 \text{ Å}$ from literature³⁶ while allowing the width σ of the instrument response function (IRF), time zero (t₀) as well as the main model fit parameter, the average ejection distance $\langle r_0 \rangle$ of the excited 5p electron, to be free. We obtain excellent agreement with our data and extract a fitted average electron ejection distance of $\langle r_0 \rangle = 7.4 \pm 1.5 \text{ Å}$ in agreement with literature values derived from fitting model (b) to TA data for 389 nm two-photon excitation conditions ($\langle r_0 \rangle = 8.0 \text{ Å}$ for 6.4 eV).³⁹ Using the entire range of reported values for $D'^{9,32,33,36,105}$ and for $r_{xn}^{32,33,36}$ yields $\langle r_0 \rangle$ values within the narrow range of 10%–20% around 7.4 Å (for details, see the supplementary material, Sec. III D 3 and Fig. S11).

E. Combined femtosecond XANES and XSS time traces

The entire XANES datasets were treated with the LCLS timing tool correction beforehand (for details, see the supplementary material, Sec. III B), which was then equally applied to the back-toback measurements of the transient XSS data. Given that the timing



FIG. 8. Results from the structural model fit via Eq. (1) to the extracted XSS signals [blue circles for the nascent iodine atom and its rearranging solvation shell, I^0 (aq)* and red squares for the bulk heat signal Δ T] and the measured amplitude of the 5.184 keV transient (I^0 2s-5p transition) from the TR-XANES data (green, small solid circles). The solid lines represent the fit curves for each data curve, the 2s-5p XANES trace using the pure diffusion model (c) [Eq. (5)]. (a) The results from the first 7 ps after laser excitation, scaled to the same amplitude after 10 ps for both XANES and XSS. (b) The same data zooming into the first psecond.

tool settings were the same for both XANES and XSS studies, we can now correct the transient XSS timing to match the same time scale as the XANES. This then allows us to compare both signals on a common time scale (Fig. 8).

Figure 8 shows the solvent temperature increase (ΔT , red squares) and the solvent cage contribution [I⁰(aq)*, blue circles] obtained from applying the structural model in Eq. (1) to the time-resolved XSS data independently at all time delays. The solid lines show how these contributions can be well captured by an IRF broadened single exponential grow-in at $t = t_0$,

$$f_{XSS}(t) = \text{IRF}(\sigma = 40 \text{ fs}) * \begin{cases} 0, & t \le t_0, \\ A \cdot \left(1 - e^{\frac{t - t_0}{\tau}}\right), & t > t_0, \end{cases}$$
(6)

with all parameters except the IRF width σ allowed as free fit parameters. The solvent temperature increase shows a time constant of $\tau = 0.57 \pm 0.07$ ps and a grow-in starting at $t_0 = -10 \pm 30$ fs. The time evolution is very similar to the neat water experiment measured subsequent to the I⁻ experiment (see the supplementary material, Figs. S3 and S4). The best-fit time constant of the grow-in of the solvent cage term is found to be $\tau = 0.35 \pm 0.04$ ps, with the grow-in starting at a time zero of $t_0 = 0.10 \pm 0.03$ ps.

V. DISCUSSION

A. Molecular dynamics (MD) results

Turning first to the MD simulation results, previous theoretical and experimental studies have determined the first peak of the I-O RDFs of I⁻ to be in the range of 3.55-3.76 Å,^{106,107} corresponding well with our equilibrium-simulation results with an I-O distance of 3.65 and 3.75 Å for the I⁻ and I⁰, respectively. Dynamics simulations performed by Pham et al.44 showed the same trends as seen in Figs. 1 and 2, with a disordering and expansion of the first solvent shell, although this "cage expansion" was larger in these studies both for their QM/MM MD simulations (~0.3 Å) and the classical MD simulations (~0.7 Å). We note that in a previous study, Pham et al.44 reported RDFs calculated from pure density functional theory (DFT) simulations of iodide in water, which would result in a somewhat smaller cage expansion, if used instead of QM/MM MD. Thus, our MM treatment of iodine seems to capture the magnitude of the cage expansion. Moreover, our usage of classical MD simulations (e.g., instead of QM/MM approaches) and the obtained results for comparison to our timeresolved data is also motivated by recent, more advanced theoretical investigations.¹⁰

In the work by Pham *et al.*,⁴⁴ an L₃ EXAFS spectrum confirmed the general trends in the simulations, although the analysis did not allow for estimating the magnitude of the cage expansion nor its time scale. From the kinetic traces shown in Fig. 2 and the SVD analysis presented in Fig. 3, we find that following photoabstraction of the electron, the sub-100 fs structural response of the caging solvent is a symmetric broadening of the first peak in the I–O RDF followed by a peak shift and broadening to longer distances. We interpret these observations as arising from the dissolution of a very structured I⁻ solvation cage held in place by electrostatic forces between the charged solute and the dipolar solvent molecules. At the moment of photoabstraction, the excess electron is ejected into the bulk, leaving the water molecules without H-bonds to the solute, and therefore they rotate into new configurations determined more by dispersive (Van der Waals) forces. This reconfiguration of the H-bond network takes <100 fs (Fig. S1 in the supplementary material, showing the $g_{\rm IH}$ dynamics), during which the O-atoms are slightly displaced from the preexcitation equilibrium configuration, leading to the observed slight broadening of the I–O peak. Following this rotational reconfiguration to a less well-defined structural motif, the water molecules as described by the position of the O-atoms redistribute from the second shell of I⁻ into the first and second shells of I⁰, forming a new solvent cage structure around I⁰.

B. Time-resolved x-ray solution scattering (TR-XSS)

Referring first to Fig. 4, the solvent-subtracted time-resolved difference scattering signal following the $I^- \rightarrow I^0 + e^-$ photoabstraction was dominated by a single component. This component strongly resembles the simulated XSS signal calculated from the (equilibrium) MD calculations of the H₂O solvation shell reorganization around I⁰. Utilizing this simulated difference signal in a three-component model, Fig. 4 shows excellent agreement between our data and model at all the time delays investigated. The residual from applying this simple model (solvation cage expansion and a temperature increase) to the x-ray difference scattering signal is rather small [Fig. 4(b)], indicating that changes in scattering arising from the presence of solvated electrons and OH/H₃O⁺ species due to three-photon ionization of water is minimal. From the delayed onset and the time evolution shown in Fig. 8 and by comparison to the nonequilibrium MD simulation results shown in Fig. 2, this difference scattering signal component can be assigned as arising primarily from dynamics involving the O-atoms of the H₂O molecules.

The XSS measurements provide a direct handle on (in particular) the I-O RDFs, allowing us to follow the structure as the new solvation shell forms. XSS is a structural probe not relying on the details in the electronic state of the solute molecule and can be compared directly to simulations of the solvent shell structure, which provide a valuable complementary input. The time constant of the solvation shell reconfiguration of $\tau = 0.35 \pm 0.04$ ps in the XSS measurements corresponds very well to the $\tau \sim 0.4$ ps time scale found in our classical nonequilibrium MD simulations. Moreover, our assignment of this component to the solvent shell reconfiguration, i.e., to a new, looser structured solvation shell around the iodine atom, is also supported by QM/MM simulations,⁴⁴ which predict a similar time scale for this process: The I-O RDFs from QM/MM simulations in Ref. 44 derive a breakup of the first solvation shell with most molecules moving away within 200-300 fs [Fig. 5(a), Ref. 44], leading to a formation of a structured hydrogen bond network on a time scale of ~3 ps, a hallmark of the formation of a hydrophobic cavity around the neutral iodine (Fig. 6 of Ref. 44). The ~100 fs delayed onset on the other hand we ascribe to the time required for disruption of the existing H-bond network structure of the caging solvent molecules, after which the O-atoms move outward, giving rise to the difference scattering signal observed. Thus, overall, the combination of results from classical liquid phase MD simulations and experimental (structural) observables presented in this paper provides valuable benchmark values for the

solvation dynamics in a simple water solvent shell, induced by a point-like change of electronic charge.

C. Time-resolved x-ray absorption near-edge structure (TR-XANES)

Concerning the sub-picosecond I L1 transient XANES maximum and minimum features (Fig. 6), Pham et al. reported that these exhibit a weak broadening (typically less than 1 eV) on the high energy side with respect to the 50 ps transient.⁴⁴ They assigned it to the creation of an intermediate $I^0(OH_2)$ complex on a sub-10 ps timescale, supported by quantum-based simulations. The signal quality in their synchrotron measurements, however, using a low intensity time-slicing x-ray source (before the advent of powerful XFEL sources) was not sufficient to unambiguously identify this broadening. In our LCLS experiments with superior signal quality, there are some differences between the shapes of the I XANES L1 transient spectra at different time delays, but these remain within the noise of the measurement; thus, we do not observe a similar L1 transient shift or broadening to higher energies on the sub-10 ps time scale. This indicates that the shape of the transient XANES is entirely controlled by the edge shift and appearance of the 2s-5p transition feature upon oxidation of I⁻. In this direction, additional computational investigations simulating the evolution of the I L₁ XANES trace and the magnitude of the effect of a potential I^0 -(OH₂) formation on the I L1 absorption transient would be beneficial. This could also allow to completely rule out such a complex formation using XAS. In addition, it would be interesting to measure (more) XAS transients with 1–10 ps time delays as the I⁰-(OH₂) complex has a predicted lifetime of 3-4 ps.⁴⁴ Further investigations into this direction by us are planned.

Our experimentally measured iodine 2s-5p transient XANES kinetic trace (Fig. 7) monitors the early recombination dynamics <10 ps from the iodine (I) point of view and this without being obscured by the ultrafast blue shift of the absorption band of the solvated electron as in most previous TA measurements." The theoretical models used for comparison to this kinetics (Fig. 7) only consider geminate iodine-electron recombination, i.e., nongeminate recombination is not taken into account. This treatment is justified by the fact that TA experiments monitoring the e⁻ absorption spectrum after two-photon excitation of I⁻ at 389 and 400 nm, respectively, observed kinetic traces that were independent of the iodide concentration for c < 1M and could be modeled by pure geminate recombination approaches.^{37–39,43} Moreover, in our experiments, geminate recombination is expected to be dominating: An initial 100 mM aqueous iodide solution yields an average distance d between the I⁻ anions of d = 25.5 Å and for the experimentally deduced excitation yield of 32%, i.e., a concentration of the photogenerated I^0 atoms of 32 mM, the average distance of the I^0 atoms is d = 37 Å. According to the TA results, 389 nm two-photon excitation (6.4 eV) results in an average electron ejection distance of $\langle r_0 \rangle \sim 8.0$ Å^{38,39} as derived from a fit of model (b) to the data. In our experiment, the electrons are thus expected to be initially ejected much closer to the parent I atom than to a non-geminate recombination partner and this is supported by the best model fit to our data (purely diffusion-limited model in Fig. 7), yielding an average electron ejection distance of $\langle r_0 \rangle = 7.4 \pm 1.5$ Å.

The survival probability calculated by the three different model approaches follows our measured 2s-5p transient XANES kinetic

trace best for a purely diffusion-limited geminate recombination [model (c)^{26,32,33,59}], while the approaches including the concept of a contact or caged pair [model (a)^{26,32,33,44}] or "bound state" [model (b)^{37–39,96}] always predict a slower decay within the first picosecond (Fig. 7).

For time delays out to 100 ps, the TA kinetic traces recorded upon low-energetic photoexcitation of I⁻ at 255 nm have been well described by the contact pair model (a).^{26,32,33} Although this model had been previously successfully used to describe the recombination kinetics observed after 400 nm multiphoton excitation of aqueous iodide⁴⁴ (as in our experiment), we expect that this model—which assumes low-energy CTTS excitation conditions-will not accurately describe our data. Pump excitation energy-dependent TA^{38,39} and time-resolved photoelectron spectroscopy studies⁴⁸ indicate that excitation into the lowest-energy I⁻ band, e.g., as for a one-photon 255 nm absorption, populates a low-energy CTTS state from which the electron separates adiabatically, thus being ejected with little excess energy and, therefore, close to the iodine parent, where it is assumed to form a contact pair.^{26,32,33} In contrast, under 400 nm multiphoton excitation conditions, different electron ejection mechanisms via nonadiabatic coupling into the water conduction band (two-photon process) or even direct ejection into this band (three-photon process) were deduced.^{38,39} These yield larger electron ejection distances, which thus calls for a more diffusion-driven kinetics time scales.^{38,39} Nevertheless, a comparison of our data to model (a) is highly valuable as it does not only allow us to probe this expected behavior but also to test if our XAS data allow us to rigidly distinguish between different recombination models. This is equally important for further planned XAS experiments following the dynamics after one-photon CTTS excitation of iodide. For comparison, we applied both sets of rate constants derived from the different TA data, i.e., $k_p^{-1} = 200$ fs, $k_d^{-1} = 70$ ps and $k_n^{-1} = 33.0$ ps for 255 nm one-photon excitation³² and $k_p^{-1} = 200$ fs, $k_d^{-1} = 59.4 \pm 6.7$ ps and $k_n^{-1} = 97.9 \pm 7.2$ ps for 400 nm multiphoton excitation,44 to model our experimental iodine 2s-5p XANES kinetic trace from LCLS, and we obtained a large discrepancy with our data in both cases. When we allow for free fitting of all model parameters to our data, we find a reasonable agreement, but now with extremely large rate constants ($k_d^{-1} = 1.36$ ps and $k_n^{-1} = 5.6$ ps) in comparison to Refs. 32 and 44, which questions the need for including such a contact pair in the first place. In summary, fitting our data to the contact pair model yields very short lifetimes for the contact pair itself, in violation of the published long lifetimes of tens of picosecond.26,

Model (b), based on a semi-analytical theory for diffusioncontrolled reactions in the presence of a potential well, $^{97-99,102}$ which includes the concept of a contact or caged ($I^0 : e^-$) pair, was implemented to better describe the electron survival probability for the entire range of photoexcitation energies used in I^- photodetachment studies.³⁷⁻³⁹ It assumes an increase in the average electron ejection distance with increasing photoexcitation energy.³⁹ While the model could accurately describe the iodine–electron recombination kinetics found in TA experiments after photoexcitation of I^- with energies below 6.4 eV and above 8.0 eV, for intermediate excitation energies in the 6.4–8.0 eV range, the observed experimental signal on the scale of tens to hundreds of picoseconds decayed faster than the model fit.^{38,39} We also observe a qualitatively similar disagreement with our data (our experimental signal decays faster than the model fit), which now covers the first few picoseconds when using the model parameters derived from TA studies for 389 nm two-photon excitation (6.4 eV) of aqueous iodide.³⁷⁻³⁹ Chen and Bradforth concluded that a possible explanation could be a more complicated excitation energy-dependent initial electron ejection length distribution.^{38,39} Again, free fitting of all model parameters does yield reasonable agreement, however, with an extremely large combined rate constant $W = W_d + W_r$ for dissociation of the caged $(I^0:e^-)$ species (W_d) and recombination of this species to reform the parent I⁻ (W_r), corresponding to a short lifetime of $W^{-1} = 0.8$ ps. Again, the reported presence of the electron trapped inside a nearby potential well with a lifetime of tens of picoseconds ($W^{-1} = 14$ ps^{37–39}) is not consistent with our data, and the ultrashort lifetimes from the fit pose again the question if the concept of the "bound state" (potential well resembling the contact pair idea) is actually required.

We note that possible non-geminate recombination or additional electrons generated by multiphoton excitation of solvent water molecules would not explain the deviation of models (a) and (b) from our kinetic data. Both processes would lead to an increase in the overall number of recombination events, but within both models, non-geminate/geminate recombination would have to proceed through the caged pair or similar "bound state" with its lifetime of tens of picoseconds; however, our data analysis is in disagreement with such a long-lived state.

Concerning model (c), we find excellent agreement to our data, suggesting that on the earliest time scales, the observed recombination dynamics can be modeled purely by three-dimensional diffusion. This contrasts the results from optical transient absorption (TA) measurements after low-energy 255 nm photoexcitation of aqueous iodide, which could not be accurately described by this model,^{32,33} but these measurements also concentrated on longer time scales up to 500 ps after photoexcitation where, e.g., subsequent chemical processes, i.e., formation of further photochemical reaction I_2^- etc. species, already plays a significant role.⁴⁴ Moreover, as described above for the 400 nm excitation conditions in our experiment, we expect a slightly different electron ejection mechanism,³ and this can potentially alter the recombination pathways. The best fit to our data yields an average electron ejection distance of $\langle r_0 \rangle$ = 7.4 ± 1.5 Å. This is in the range of the earlier reported ~8 Å average electron ejection distance derived from TA experiments following two-photon excitation at 389 nm (6.4 eV), where the average electron ejection distance was inferred from fits of model (b) to the experimental data.^{38,39} In any case, this points to a two-photon excitation process (6.2 eV) in our experiment, excluding significant three-photon (9.3 eV) excitation contributions: The latter should yield larger average initial ejection distances above >40 Å.^{38,39} This is also supported by the fact that in TA experiments with excitation energies above 8.2 eV,^{38,39} no iodine-electron recombination was observed on the time scales below <500 ps, but in our experiment, we observe a fast recombination already within the first few picoseconds, i.e., in agreement with an excitation energy below 8.2 eV, i.e., a two-photon process.

We note that the results of models (b) and (c) (both including a diffusion treatment) do not critically depend on the actual choice of the diffusion constants themselves (see also the supplementary material, Sec. III D 3 and Fig. S11). This is important since different combined room-temperature iodine–electron diffusion coefficients

have been reported ($D' = 5.8-8.0 \cdot 10^{-4} \text{ Å}^2/\text{fs},^{32,33,36,37}$) and also since any model relying on diffusion coefficients requires thermal equilibrium conditions, which we do not have on the few picosecond timescale.^{32,33,37} However, if we assume, e.g., a temperature increase by a factor of three for the early time scales, the (now timedependent) diffusion coefficient would change by a factor of 2.3 (from $D' = 5.8 \cdot 10^{-4}$ to $D' = 1.35 \cdot 10^{-3} \text{ Å}^2/\text{fs}$),³⁶ and the resulting average electron ejection distance $\langle r_0 \rangle$ will only change by less than 1 Å (or <15%) [see the supplementary material, Fig. S11 for model (c)].

VI. CONCLUSIONS

The structural solvent reorganization and geminate recombination dynamics following 400 nm two-photon electron photodetachment from aqueous iodide, $I^- \rightarrow I^0 + e^-$, was revisited with new structural tools (XANES, XSS) on the femtosecond time scale. The present study delivers new insight into the processes occurring within the first picosecond, which are obscured in optical transient absorption (TA) measurements due to (i) the dynamic Stokes shift of the nascent electron absorption band in the visible-infrared region and (ii) which observed only details of the ejected electron instead of the iodine atom itself.

The TR-XANES results monitored the prompt generation of iodine atoms and identified hereby both time zero and the photoexcitation yield quantitatively (34% under the chosen laser conditions). The kinetic analysis is compatible with electrons ejected on average $\langle r_0 \rangle = 7.4 \pm 1.5$ Å away from the parent iodide. The decay in our XANES kinetic trace observed within the first picosecond is not accurately described with models invoking the existence of a so-called contact pair (I⁰ : e⁻) or a similar "bound state," but it is quantitatively in line with a pure diffusion of the separated partners, I⁰ and e⁻, toward geminate recombination.

The timing extracted from the TR-XANES measurements allowed us to precisely define time zero also for the subsequent femtosecond XSS experiments, and the ~100 fs delayed onset of the iodine-water cage response becomes visible, which we ascribe to the time required for disruption of the existing H-bond network structure of the caging solvent molecules. The classical MD simulations support such a delayed response of the caging water shell reconfiguration by roughly 0.1 ps, after which the initially tight eightfold coordinated solvent shell expands slightly and simultaneously reorients the dipole moment vectors and allows more water molecules (~22) to form the new, looser structured solvation shell around the iodine atom. The time constant of the solvation shell reconfiguration of $\tau = 0.35 \pm 0.04$ ps in the XSS measurements corresponds very well with the $\tau \sim 0.4$ ps time scale found in our classical nonequilibrium MD simulations, and it is also in reasonable agreement with previous QM/MM MD simulations, which predict a cage expansion within 200-300 fs leading to the formation of a structured hydrogen bond network on a time scale of ~3 ps.4

The results presented in this paper deliver a first direct visualization of the structural dynamics of the H_2O solvent shell following light-induced electron detachment from aqueous atomic ions. Here, the solute promptly changes its hydrophilic nature into a hydrophobic neutral atom. Combining XSS with XANES allows us to get a deeper insight into the femtosecond guest-host interactions

of reacting solutes. This marks a first step toward an ultrafast mechanistic and electronic understanding of the elementary steps in photochemically driven solvation dynamics.

We note that it would be highly interesting to compare the ultrafast structural changes observed with x-ray tools after multiphoton excitation of aqueous I⁻ (as in the current manuscript) to results from otherwise identical one-photon excitation (e.g., 255 nm) measurements that directly excite the first CTTS state. This would allow to rigidly compare the different reaction pathways and related models, e.g., the contact pair model. For such experiments, producing 255 nm with high intensities can be a challenge by itself and they need to be implemented at free electron laser end stations, which, in turn, have to provide x-ray laser beams in a challenging energy range (e.g., L1-edge: ~5.2 keV). For CTTS studies, the homologue aqueous bromide seems equally interesting: Not only does it have energetically higher absorption bands allowing for excitation into the first CTTS state with 200 nm radiation, which can be produced via fourth harmonic generation from an ultrafast, e.g., Ti:Sa laser system and a K absorption edge in the 13.5 keV region, but also studies on aqueous Br⁻ would allow us to access the (theoretically predicted¹⁰⁸) differences in the photoexcitation dynamics in the homologue series of the halides. Further studies in this direction are underway.

SUPPLEMENTARY MATERIAL

See the supplementary material for further information on MD-simulated time-resolved radial distribution functions, the experimental setup, the time-resolved XSS experiments on neat water, the time-resolved I L₁ XANES data, and the detailed procedure regarding their analysis, including the mathematical description of the three theoretical approaches used to model the recorded XANES kinetic trace as well as a derivation of the absorption cross section of the nascent 2s-5p transition.

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AUTHOR DECLARATIONS

Conflict of Interest

The authors have no conflicts to disclose

Author Contributions

Peter Vester and Katharina Kubicek contributed equally to this work.

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DATA AVAILABILITY

The data that support the findings of this study are available from the corresponding author upon reasonable request.

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