Isotope effects in dynamics of water isotopologues induced by core ionization at an x-ray free-electron laser

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Abstract

Dynamical response of water exposed to x-rays is of utmost importance in a wealth of science areas. We exposed isolated water isotopologues to short x-ray pulses from a free-electron laser and detected momenta of all produced ions in coincidence. By combining experimental results and theoretical modelling, we identify significant structural dynamics with characteristic isotope effects in H_2O^{2+} , D_2O^{2+} , and HDO^{2+} , such as asymmetric bond elongation and bondangle opening, leading to two-body or three-body fragmentation on a timescale of a few femtoseconds. A method to disentangle the sequences of events taking place upon the consecutive absorption of two x-ray photons is described. Methodological access to the structural properties and dynamics of dissociating water isotopologues exposed to x-rays opens the way to a broader understanding of aqueous systems of biological interest.

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I. INTRODUCTION

The dynamical behavior of water molecules exposed to x-rays is of utmost interest from both a fundamental point of view and with respect to the possible implications in a multiplicity of research fields. The relevance of radiation-induced dynamics ranges from radiation damage in living tissues [1] to problems in oncology treatments [2] and even interstellar phenomena [3]. High-energy radiation can induce a variety of processes in water, such as the production of multiply charged species and fragmentation processes yielding ions or radicals. In particular, if a core electron is ejected, a relaxation process occurs, in which a valence electron fills the core vacancy and another valence electron is emitted (Auger-Meitner decay). From gas-phase data, it is known that the resulting doubly charged water cation (H_2O^{2+}) eventually dissociates into fragments, which are mainly either a hydroxyl cation (OH^+) or an atomic oxygen (O), and protons (H^+) [4]. These dissociation processes have been studied by ion-mass spectroscopy [5] or ion/ion-electron coincidence measurements [4,6-9].

In a recent study [10], conducted at the Small Quantum Systems (SQS) instrument [11,12] of the European XFEL [13], we exploited the possibility of using very intense and very short x-ray pulses to induce sequential multiphoton absorption in isolated water molecules. We measured triple ion-ion-ion coincidences (H⁺/Oⁿ⁺/H⁺) and concentrated on the coincidence channel leading to two protons and a doubly charged oxygen ion, which can be almost entirely related to consecutive absorption of two photons and accompanying Auger-Meitner decay processes involving the O 1s inner shell. We stress that such dicationic states are the ones relevant in many situations where water molecules are exposed to x-rays or other high-energy radiation. Therefore, information on their dynamical evolution is crucial in a wealth of different fields of science.

By appropriate choice of experimental parameters and theoretical modelling, we demonstrated that with our approach we can follow the dynamics occurring between two consecutive photoabsorption events in the time span of approximately 0-25 femtoseconds. We were able to characterize both structural and temporal evolution of the above-mentioned dissociation events, achieving a complete characterization of both geometrical changes and timing of the events, concerning all dicationic states formed after core ionization and Auger-Meitner decay.

In particular, we demonstrated that, on a timescale of a few femtoseconds after core ionization and subsequent Auger-Meitner decay, water undergoes structural deformation such as asymmetric O—H bond stretching and/or opening of the bond angle all the way up to 180 degrees, eventually leading to rapid two-body or three-body fragmentation in asymmetric and/or unbent geometries.

In the present work, we extend the investigation to two water isotopologues, semi-deuterated water, HDO, and fully deuterated (heavy) water, D₂O. Using the same instrumentation, data analysis, and theoretical modelling, we are able to investigate in unprecedented depth the related isotope effects. In particular, we show that similar bond elongation and bond-angle-opening mechanisms are present for all three isotopologues as well, exhibiting, however, significant differences. Specifically, the dynamical processes are slower in D₂O and show

pronounced asymmetries in HDO due to the heavier mass of D as compared to H. By combining the unique characteristics of the photon beam generated by the European XFEL, the coincidence imaging abilities of the COLTRIMS reaction microscope, and an advanced theoretical modelling, we show which deep insight we can obtain into a photoinduced dynamical process.

II. METHODS

A. Experiment

The experiment was performed at the Small Quantum Systems (SQS) scientific instrument [11,12] of the European XFEL in Schenefeld, Germany [13]. The accelerator was operated at 14 GeV electron energy with a bunch charge of 250 pC, and an effective number of 970 photon pulses per second was generated for the present experiment in the SASE3 soft x-ray undulator. The x-ray pulses had an estimated duration of about 10–25 fs, and a maximum pulse energy of 4.4 mJ, measured after the last undulator [11,12,14,15]. The photon energy was set to hv=1000 eV, sufficient to extract all electrons from the investigated systems via consecutive photoionization and Auger-Meitner decay steps. The spectral bandwidth was determined as approximately 0.9% of the central photon energy. Data were collected using a COLTRIMS reaction microscope [16], with a supersonic gas jet consisting of a 50:50 mixture of H₂O and D_2O , leading to approximately 25% H_2O / 25% D_2O / 50% HDO vapor, and helium. The water vapor was generated by heating a reservoir (located outside of the vacuum chamber) to 40°C, while the gas line and the nozzle (consisting of an aperture of 200 µm diameter) were heated to 50°C and 70°C, respectively. The jet was intersected with the x-ray beam. Charged particles which were generated in the interaction region due to the photoionization and decay processes were then guided to a position- and time-sensitive detector [17]. From the measured position of impact of a particle and its time-of-flight, the particle's three-dimensional momentum vector was determined in an offline analysis. All ions generated in the photoreaction were measured in coincidence, which allowed to determine kinetic energies and relative emission angles. The detector employed for the measurement had an active diameter of 120 mm and consisted of a stack of two microchannel plates. The spectrometer used for guiding the charged particles to the detector consisted of an acceleration region with a strong electric field of E=213 V/cm (with a length of 153 mm) followed by a region with constant electric potential. The overall distance between the interaction region and the ion detector was 250 mm. We refer to Ref. [10] for further experimental details.

For the results presented in Fig.1, the measured ion momenta were transformed into a recoil frame of reference. This coordinate frame is defined by the emission direction of the O^{2+} ion (providing the x-axis of the coordinate frame) and the emission direction of one of the two emitted proton or deuteron ions. The latter spans the xy-plane together with the emission direction of the O^{2+} ion. A similar transformation was done for the results presented in Figs. 4 and 5, which uses a proton as the reference particle. In multicoincidence experiments, the

detection of fragments that do not belong to the same parent molecule can lead to false coincidences that affect the quality of the data. In order to reduce drastically such events, the sum momentum of the three measured ions was inspected for each photoreaction event. After subtraction of the average linear momentum of the absorbed photons and the average molecular beam velocity, the sum reflects only the recoil momentum due to the emitted electrons. We have considered only events where the sum momentum is less than < 20 a.u. as valid.

B. Simulation

The theoretical modelling of the multiple ionization and fragmentation dynamics has been described in Ref. [10]. Here we only give a brief overview of the methodology.

The dynamics of the water isotopologues was simulated using a kinetic Monte-Carlo approach. We conducted *ab-initio* molecular dynamics simulations, where, in each time step, we calculated on-the-fly potential energy gradients, photoionization cross sections, and Auger-Meitner decay rates for the current electronic state [18-20]. While the molecular geometry was propagated, photoionization, fluorescence, and Auger-Meitner decay transitions took place randomly according to computed probabilities. The electronic structure is calculated using the XMOLECULE electronic structure toolkit [18,20,21] employing the 6-31G(d,p) basis set [22] and the Hartree-Fock method. For open-shell configurations we employed the restricted open-shell Hartree-Fock method.

The initial conditions for the molecular dynamics trajectories were sampled from the Wigner distribution of the vibrational ground state of the neutral molecule. A time-step of 0.1 fs was used to propagate the molecular geometry and the simulation was conducted for a total time of 200 fs. We sampled at total number of 8,000 trajectories.

For the x-ray pulse, we employed a photon energy of 1 keV and a temporal pulse shape described by two Gaussians, both centered at t = 90 fs, with full widths at half maximum of 5 fs and 35 fs and relative weights of 0.6 and 0.4, respectively. A total fluence of 2×10^{11} photons/ μ m² was employed. For a more detailed discussion of the pulse parameters, we refer to Ref. [10].

In order to disentangle the fragmentation dynamics of HDO, additional, more simplified simulations were performed for three model scenarios. In scenario 1 (instantaneous Coulomb explosion) the molecular geometry was propagated using bare Coulomb forces, where proton/deuteron carries a charge of 1 and oxygen a charge of 2. In scenario 2 (delayed ionization), also bare Coulomb force were used, but the oxygen atom was kept neutral for the first 10 fs. In scenario 3 (sequential fragmentation), on top of the delayed ionization of scenario 2, an artificial bond potential between O and D fragments was employed for the first 10 fs using bond parameters from the SPC/FW force field [23] to mimic a sequential fragmentation.

III. RESULTS AND DISCUSSION

A. Newton diagrams

In Fig. 1 we present the momentum space data for all three water isotopologues as Newton diagrams. The final momenta of the three ionic fragments O^{2+} , $H(D)^{+}$, and $H(D)^{+}$ were detected in coincidence, as described above. The experimental diagrams are compared with simulations of the ionization and fragmentation dynamics conducted with the XMOLECULE toolkit [18,20,21].

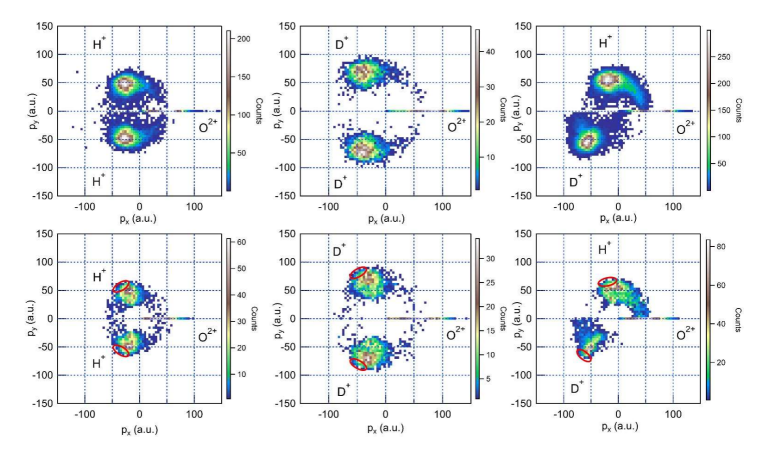


Figure 1. Newton diagrams (experiment, top and theory, bottom) of the ion momenta for O^{2+} and two H^+ (D^+) ions for H_2O (left column), D_2O (middle column), and HDO (right column), detected in coincidence. The oxygen momentum defines the x axis. In the theoretical plots, the red ovals mark the momentum values expected for an instantaneous Coulomb explosion from the neutral ground-state configuration.

The coincidence channel leading to two singly charged light particles (protons or deuterons) and an oxygen ion with charge 2+ allows us to follow in detail the evolution of doubly charged water molecular ions formed by the absorption of a first photon and subsequent Auger-Meitner relaxation in a time interval in-between two consecutive photoabsorption events.

All three Newton diagrams show two dominant peaks for the $H(D)^+$ momenta and a pronounced tail. The main peaks originate mostly from an almost instantaneous Coulomb explosion, caused by the absorption of two photons within a very short time interval: in all three systems, the final

overall charge state 4+ is reached rapidly and the intermediate molecular dication has not enough time to change its geometry. In contrast, the pronounced tail in the $H(D)^+$ momentum distribution pointing towards the momentum direction of O^{2+} can be attributed to a situation where the delay between the two photoabsorption events is sufficiently large, allowing the molecule to undergo some form of structural rearrangement after the first photoionization event and first Auger-Meitner decay, and before it is torn apart by the strong Coulomb repulsion induced by the second photoionization event and second Auger-Meitner decay.

In our previous work [10], we demonstrated that the dynamical processes that can be related to the tail in the Newton diagram are of two different origins: an asymmetric bond elongation, whose limit is sequential two-body fragmentation into OH⁺/OD⁺ and D⁺/H⁺, and a bond-angle-opening trend, whose limit is a linear geometry or even an overbent configuration.

We now focus on the clear differences that we can immediately notice between the three Newton diagrams for H_2O , D_2O , and HDO.

In particular, the comparison between D_2O in Fig.1(b) and H_2O in Fig.1(a) evidences a less pronounced tail in the former. This can be easily explained with the consideration that both motions we mentioned before, i.e. bond elongation and bond-angle opening, are slower in D_2O due to the difference of mass between proton and deuteron. Therefore, during the same time interval, whose maximum length is set by the x-ray pulse duration, the percentage of molecular ions undergoing structural deformation is lower.

The comparison between H_2O in Fig.1(a) and HDO in Fig.1(c) points to two effects. On the one hand, the angles between the momenta of the three particles are not symmetric, as is the case for both H_2O and D_2O . On the other hand, there is a tail for both H^+ and D^+ momenta towards the O^{2+} momentum, but the two tails are of different length and intensity.

The different angles between the H^+ and D^+ momenta with respect to the O^{2^+} momentum (122° for the angle between H^+ and O^+ , 134° for the angle between D^+ and O^+ in momentum space) are simply a consequence of momentum conservation during the entire dissociation process. The peak values of the momentum distributions are close to the values one would expect from an instantaneous Coulomb explosion, marked as red ovals in the simulations (lower row in Fig. 1).

The difference in the tails of the momentum distributions in the Newton plots is more revealing of the different structural dynamics of the three ionized systems. The tail is clearly longer for H^+ , while the momenta of D^+ are more concentrated towards the value corresponding to instantaneous Coulomb explosion. Due to the mass difference of the fragments, the momenta generated by the Coulomb explosion is not equally distributed across the three ionic fragments. In addition, the bond length between O and H increases more quickly than the O-D bond length. Accordingly, a dynamical process such as bond elongation is more advanced for the O-H bond when the second photon eventually triggers the final Coulomb explosion. This asymmetry is further enhanced by the subtle interplay of the two light atoms for some of the reached intermediate dicationic states. It has been reported that the lowest dicationic states in HDO asymptotically lead to a fragmentation into $OH^+ + D^+$ or $OD^+ + H^+$, where the latter is much

more favorable by a ratio of 1:6 [24,25] due to the dynamical mass effect. If the delay between the two photoabsorption events is large enough, one may thus expect to see some intermediate stable DO⁺ and HO⁺ fragments with unequal ratio that are then ripped apart by the second fragmentation.

B. Scatter plots

Additional information can be obtained by choosing a different representation of the data, which can give more detailed insight into the processes than the Newton diagrams, in particular on specific aspects of the molecular dynamics.

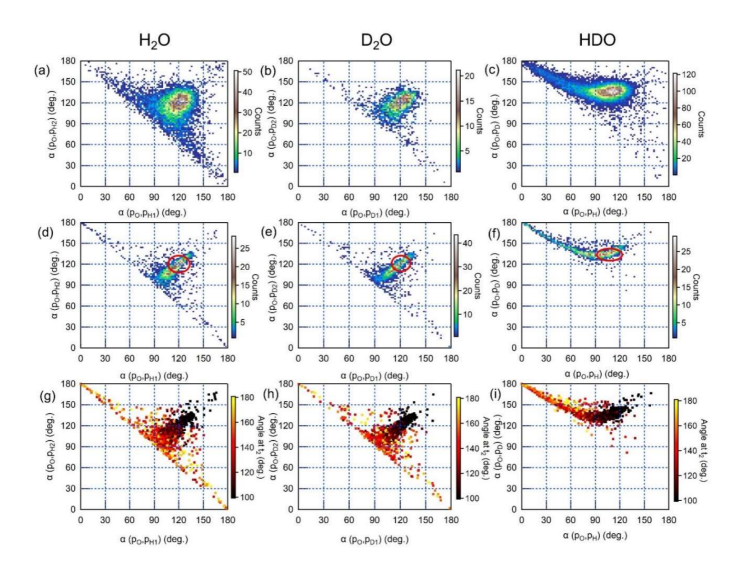


Figure 2. Scatter plots showing the angles of the final proton (deuteron) momenta with respect to the final oxygen momentum for H_2O (left column). D_2O (middle column), and HDO (right column). In each column, the top panel is the experiment, the middle panel is the simulation, and the bottom panel is the simulation where the color code represents the angles at the time of the second photoionization. The red ovals in the simulation panels indicate the value ranges for the event of a Coulomb explosion from the neutral ground-state configuration.

In Fig. 2, we therefore show proton/deuteron emission angles (experiment and simulations) with respect to the emission direction of the doubly charged oxygen ion for all three water isotopologues. From the correlation of the two angles shown in Figs. 2(a) and 2(d) (H₂O, experiment and simulation) and Figs. 2(b) and 2(e) (D₂O, experiment and simulation), it is clear that the majority of the emitted protons/deuterons is distributed at angles around $\approx 115^{\circ}$, which is close to what can be predicted for an immediate Coulomb explosion taking place in almost ground-state geometry (see the red ovals in the middle row of Fig. 2). However, in a significant number of events, the angles of the two protons/deuterons are strongly anticorrelated, and they exhibit an angle sum up to about 180°. In the simulations shown in Figs. 2(g) and 2(h), the events with large HOH-bond angle, or DOD-bond angle, at the time of the second photon absorption are highlighted in orange and yellow. In our previous work [6], we discussed that this signal must be attributed to an unbending motion, where the water molecule reaches an almost linear geometry at the point in time when the second photoabsorption event triggers the final Coulomb explosion.

An interesting observation in the present context is that this effect is clearly visible for both water and heavy water. However, there is no pronounced difference, indicating that the bond opening motion is marginally affected by the mass difference.

The comparison with HDO shows significant differences with respect to both H_2O and D_2O . The corresponding plots are shown in Figs. 2(c), (experimental), 2(f) (simulation), and 2(i) (simulation with color code indicating the angle at the time of the second photon absorption). The maximum density is centered around the values of the two angles that correspond to an instantaneous Coulomb explosion: $\approx 105^{\circ}$ for H^+ and $\approx 135^{\circ}$ for D^+ . In addition, there is a tail towards larger angles for D^+ , up to a limit of 180° , and smaller angles for H^+ , down to a limit of 0° .

As mentioned before, the repulsion between O^{2+} and D^{+} gives a stronger kick to O^{2+} than the repulsion between O^{2+} and O^{2+} are simple argument based on the difference of mass, the oxygen ion tends to be emitted back-to-back with respect to the deuterium one. Furthermore, the O^{2+} bond is likely to stretch more rapidly than the O^{2+} one. In the limit of a molecule reaching a linear geometry after the absorption of the first photon and undergoing sequential fragmentation, the O^{2+} bond is ruptured first and the O^{2+} fragment undergoes Coulomb explosion upon arrival of the second photon. Therefore, the O^{2+} and O^{2+} fragments move in the same direction, while the O^{2+} and O^{2+} fragments move back to back. In the scatter plot, a whole distribution of angle combination values corresponding to all intermediate situations is visible.

In Fig. 3, we show another type of scatter plot, displaying the absolute momenta of the two light ions. Figs 3(a) and 3(d) show the momenta for H_2O , experiment and simulation, and Figs. 3(b) and 3(e) the momenta for D_2O , experiment and simulation. Whereas most of the proton/deuteron momenta are clearly symmetric, i.e., they show similar absolute values, a significant fraction of them is asymmetric. In Figs. 3(g), and 3(h), the events in which the two particles have asymmetric OH or OD bond lengths at the time of the absorption of the second photon are shown in orange and yellow.

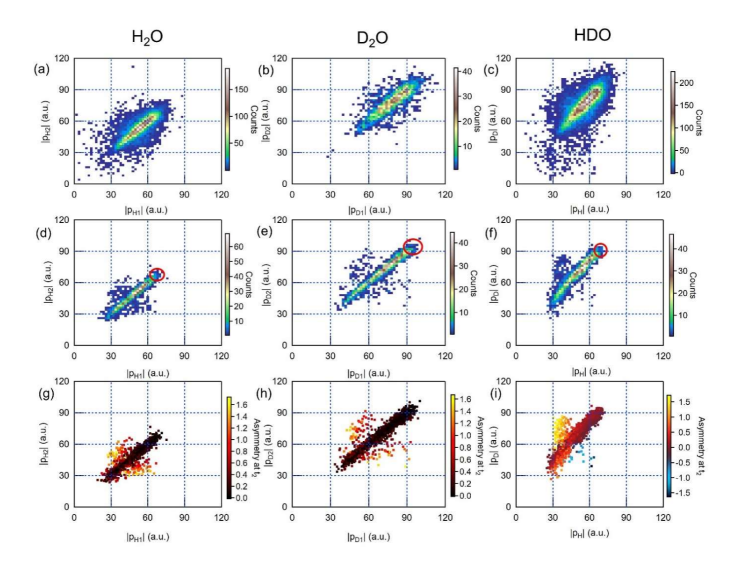


Figure 3. Scatter plots showing the magnitudes of the final proton/deuteron momenta for H_2O (left column), D_2O (middle column), and HDO (right column). In each column, the top panel is the experiment, the middle panel is the simulation, and the bottom panel is the simulation where the structural asymmetry at the time of the second photoionization is reflected by a color code. Asymmetry for H_2O and D_2O is defined as $2|d_{OX1}-d_{OX}|/|d_{OX1}+d_{OX2}|$, where X stands for H and D, respectively. For HDO, asymmetry is given by $2(d_{OH}-d_{OD})/(d_{OH}+d_{OD})$. Thus, positive asymmetry indicates a breakup into $OD^+ + H^+$, whereas negative asymmetry indicates a breakup into $OH^+ + D^+$. The red ovals in the simulation panels indicate the value ranges for the event of a Coulomb explosion from the neutral ground-state configuration.

From this analysis, we can clearly assign asymmetric proton/deuteron momenta to asymmetric bond elongation after the first core photoionization and Auger-Meitner decay. Eventually, this asymmetric bond elongation may correspond asymptotically to a fragmentation channel of H_2O^{2+} into OH^+ and H^+ , which has been reported in the literature for the three lowest dicationic states [7,8,26-28] and accounts for about 25–30% of the total Auger-Meitner yield [29,30].

The situation is again rather different for HDO. The absolute momenta are in general asymmetric and most of them follow a linear relation. The slope of this linear relation can be

understood if one considers the mutual Coulomb repulsion of O^{2+} and D^+/H^+ in a symmetric configuration independently, in which case

$$\frac{|p_D|}{|p_H|} = \alpha = \sqrt{\frac{m_D}{m_H}} \sqrt{\frac{m_O + m_H}{m_O + m_D}} \cong 1.37.$$
 (1)

There are many points beyond this linear relation, suggesting a fragmentation from an asymmetric geometry. This interpretation is supported by the analysis of the simulation data in Fig. 3(i), where a color code indicates the relative interatomic distance $2(d_{OD}-d_{OH})/(d_{OD}+d_{OH})$ when the second photon is absorbed.

For the cases outside of the linear relationship, the deuteron momentum tends to be much higher than the momentum of the corresponding proton. Only for a few events the proton momentum is significantly larger than the corresponding deuteron momentum. We quantify the degree of asymmetry by the fraction of data where $|p_D|/|p_H| \gtrsim \alpha \pm 0.2$ and find a ratio of 0.3 (simulation) and 0.16 (experiment). As expected, the deuteration effect on the asymmetry found here is considerably smaller than reported before (1:6), which is related to the fact that we address here the overall yield of dicationic states and do not limit ourselves to the three lowest dicationic states. Nevertheless, the deuteration-induced asymmetry can be considered significant also for the overall Auger-Meitner yield.

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C. Native-frame analysis

In the discussion above, we have addressed the issues of asymmetric bond breaking and unbending motion, which are structural changes common to all three molecules under investigation.

While we can clearly identify an asymmetric bond elongation, from the above analysis based on Newton diagrams and scatter plots it is difficult to clearly separate possible sequential steps. One of those could be a first complete bond rupture after the absorption of the first photon and then a Coulomb explosion of the remaining diatomic fragment after the absorption of the second photon. We can state that the complete sequential fragmentation is the limit case, and in the above figures we see a distribution of all possible intermediate states.

In a previous work of some of the present authors, it was demonstrated that it is possible to distinguish sequential from concerted fragmentation in a triatomic molecule, CS₂, on the ground of the different KER ranges for the two processes [31]. Furthermore, a method called "native-frame analysis" has been recently reported in the literature [8,32,33]. The basic principle is to follow step-by-step a fragmentation process by creating "native frames" associated with each stage of the sequential dissociation. By using a combination of the momentum acquired by an undissociated ionic fragment in a first step and the momenta of fragments produced in a second step, it is possible to remove the momentum acquired in the first step and obtain the momentum distribution of the fragments produced in the second step, which is what is called "native frame" approach. It is an effective way to extract distinct

information on the dissociation dynamics and to obtain the momentum distributions for one specific fragmentation process.

We concentrated our analysis on HDO, where the two chemical bonds are more likely to elongate and then break on a different timescale. Assuming a two-step fragmentation where the O-H bond breaks and the proton leaves the molecule first, the final momentum distribution is a combination of the momentum acquired by the undissociated OD^{3+} fragment during the first step of the breakup, \mathbf{p}_{OD} =- \mathbf{p}_{H} , and the momentum acquired by the O and D ions during the second step, $\mathbf{p'}_{\mathrm{O}}$ =- $\mathbf{p'}_{\mathrm{D}}$. Therefore, the final momenta \mathbf{p}_{O} and \mathbf{p}_{D} of the O and the D fragment after the Coulomb explosion can be written as:

$$\mathbf{p}_{O} = \mathbf{p'}_{O} - \frac{m_{O}}{m_{O} + m_{D}} \times \mathbf{p}_{H},$$

$$\mathbf{p}_{D} = \mathbf{p'}_{D} - \frac{m_{D}}{m_{O} + m_{D}} \times \mathbf{p}_{H}$$
(2)

Accordingly, the momenta acquired in the second step of an assumed two-step fragmentation ($\mathbf{p'}_{O}$ and $\mathbf{p'}_{D}$) can be obtained from the final state momenta \mathbf{p}_{O} and \mathbf{p}_{D} by subtracting the recoil imparted by the emitted proton on the O and D ions in the first fragmentation step.

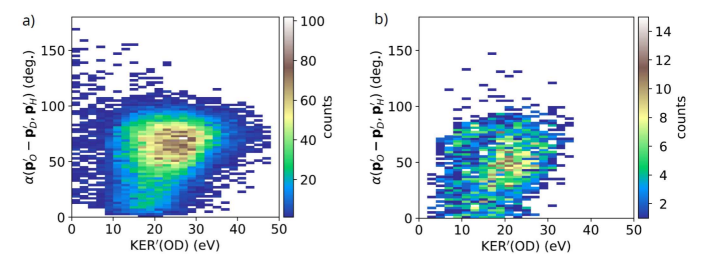


Figure 4. Angle between the momenta of the hypothesized OD³⁺ intermediate and H⁺ fragments versus intermediate KER for OD³⁺ in HDO (see equations in the text). Left: Experiment. Right: Theory.

A first step along the native-frame-analysis scheme to shed further light on the sequential-versus-concerted dissociation is to plot the relative angle between the momenta of the H⁺ fragment and the OD³⁺ intermediate species as a function of the KER of the OD intermediate. We show such experimental and theoretical plots in Fig.4. In case of sequential ionization, we

expect a broad distribution in the OD-H angle, up to covering the full range of 0 to 180 degrees. This would indicate full rotational turns of the OD intermediate after the emission of the proton. In contrast, if a sequential dissociation does not occur, or it is a minor channel, such angle would span a more limited range. This is actually the case in Fig.4. These results indicate that the OD fragment is not free to rotate, and hints at a dominant concerted fragmentation process.

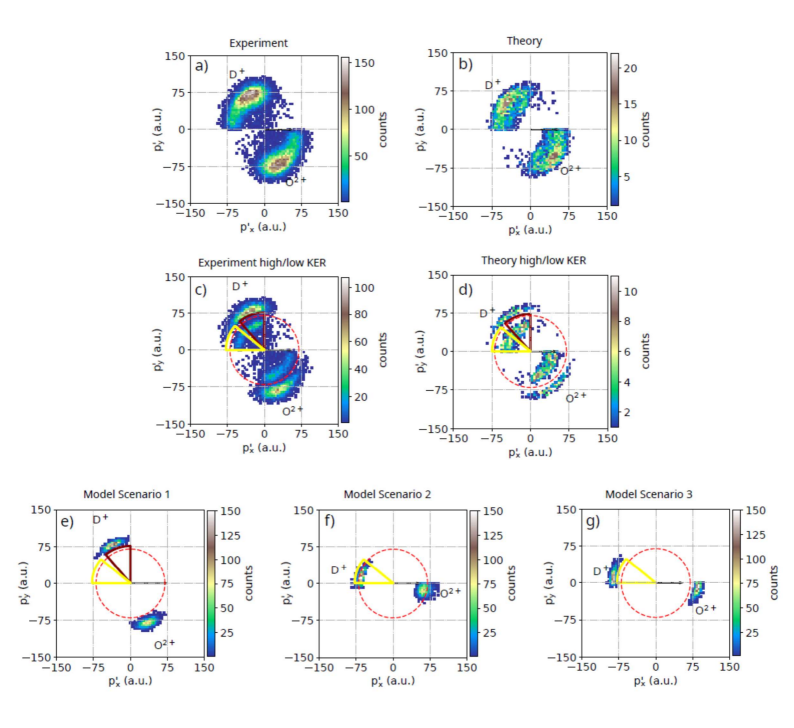


Figure 5. Native frame plots for O^{2+}/D^+ . The momentum vector of the H^+ fragment is oriented along the horizontal axis. 5(a) and 5(b): total momenta, experiment and theory; 5(c) and 5(d): corresponding plots, but gated on ranges of high and low KER, Here, low KER means KER < 30 eV, high KER means KER > 60 eV; 5(e), 5(f) and 5(g), three different model scenarios (simulations) (see text for details).

In Fig. 5, we show the momentum-space results of the native-frame analysis of the HDO fragmentation, together with the theoretical simulations for three different possible scenarios. The corresponding plots are similar to the regular Newton diagrams shown in Fig.1, but now we depict the momenta of the two ions gathered in the second step of an assumed two-step fragmentation, i.e. the distribution of $\mathbf{p'}_0$ and $\mathbf{p'}_D$ in Eq. 2. In addition, we chose the momentum vector of the escaping \mathbf{H}^+ ion \mathbf{p}_{H} as the reference-axis, i.e., the x-axis in the figure

In general, in a triatomic system the momentum distribution associated with a sequential breakup is distributed along a circle, as the diatomic fragment left behind in the first breakup rotates with respect to the momentum vector of the departed atomic ionic fragment. The circle is typically considered as a clear sign of sequential fragmentation (see e.g. the plot for the CS²⁺ fragment in CS₂ [33].

In the plot of the experimental HDO data, Fig. 5(a), the momentum distributions align along quarter circle segments rather than a full circle. This seems to indicate again that that the molecular dynamics do not consist of two independent dissociation steps, such as $HDO^{2+} \rightarrow OD^{+} + H^{+}$ after the absorption of a first photon, involving core photoionization and Auger-Meitner decay, followed by further fragmentation of $OD^{3+} \rightarrow O^{2+} + D^{+}$ after the absorption of a second photon, involving core ionization and a second Auger-Meitner decay in the OD^{+} fragment. We can identify nuclear motion, i.e., the structural deformation of the molecule, but no clear evidence of a clean sequential process. In other words, the features in the diagrams are linked to fragmentation after asymmetric / angular dynamics in the dication.

Our results may appear in contradiction with what is reported in the literature concerning a two-step dissociation in HDO, identified with a similar multicoincidence analysis [8]. However, we stress the point that in [8] the experiments are based upon one-photon double-valence photoionization, and there is no timing information concerning how long after photoionization the sequential breakup takes place. As mentioned before, under our experimental conditions we can follow all processes occurring between two photoabsorption events, within a maximum time span of about 25 fs (the estimated pulse duration). We gain the additional information that the sequential breakup (as identified by the native frame analysis) "takes time", and it is unlikely to occur on a very short timescale.

The same native frame analysis was performed on the simulation data, Fig. 5(b) and by considering different limit cases based on pure Coulomb forces (details are given in the Method section).

In particular, in Figs. 5(c) and 5(d) we show the results from the experiment and the simulation by gating on a subset of the data excluding events with an intermediate KER of 30eV<KER<60eV. Additionally, we marked some value ranges of the plot which in the bottom panel correspond to the following different scenarios.

- Fig. 5(e), model 1: Instantaneous Coulomb explosion with charges O²⁺, H⁺, and D⁺;
- Fig. 5(f), model 2: First dissociation due to Coulomb charges of O, H⁺, and D⁺, and then after 10 fs switch to O²⁺, H⁺, and D⁺, i.e., delayed photoionization;
- Fig, 5(g), model 3: only release of H⁺ keeping OD⁺ connected via a harmonic spring, and after 10 fs breakage of the OD fragment into O²⁺ and D⁺, i.e., two-step fragmentation.

Model scenario 1, which corresponds to instantaneous Coulomb explosion, corresponds also to high momenta (and high KER), as already discussed concerning the Newton diagrams. Scenario 3 yields similar D⁺ and O²⁺ momentum magnitudes as in scenario 1. Compared to that, scenario 2 yields slightly lower D⁺ and O²⁺ momentum magnitudes. We can conclude that the main part part of the features observed in the experiment and the full simulation corresponds to what one sees in scenario 1: immediate Coulomb explosion with charges O²⁺, H⁺, and D⁺. The circle structure could be interpreted as a fragmentation of sequential nature, at least partially (extreme case, scenario 3, after 10 fs). Considering a continuum of delay times for both scenarios, the half-circle structure would emerge. But also, just a delayed Coulomb explosion (extreme case, scenario 2: O neutral, H⁺, D⁺, and further charging up after 10 fs) gives a very similar picture.

We can conclude that in the present case the native-frame analysis is effective in showing that the contribution of a pure two-step sequential dissociation (i.e., involving full rotations between the two fragmentation steps) is minor, at least in the time interval available, and different scenarios should also be considered.

D. HDO dynamics

To better visualize the dynamical processes triggered by the photoabsorption and fragmentation processes of the HDO molecule, we now discuss snapshots from movies of the calculated trajectories. Those movies are available online.

Figure 6 shows the time evolution of the ensemble of HDO trajectories in position and momentum space. The coordinates are shown in the recoil frame, i.e., the frame where the final oxygen momenta are occurring along the x-axis. Snapshots are shown before (t=-30 fs) and after the center of the pulse (t=30 fs). For the latter, the momentum coordinates are similar to the Newton diagram shown in Fig.1, which represents the momentum coordinates in the recoil frame for $t \to \infty$. The earlier snapshot shows instead the HDO molecule having negligible momentum, but a broad distribution of orientations in position space. The hydrogen atom tends to point towards the upper right and the deuterium atom tends to point toward the lower left direction.

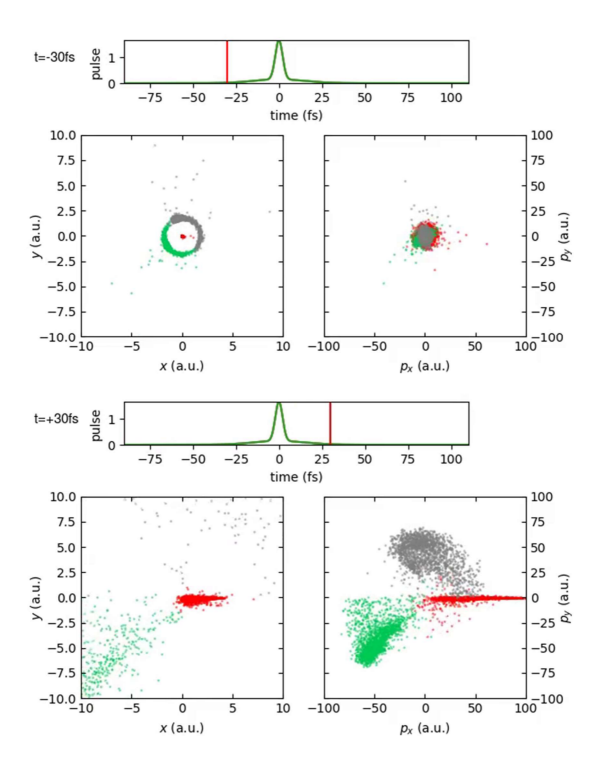


Figure 6. Position and momentum space coordinate in recoil frame for the ensemble of HDO molecules as a function of time. Left panels show position, right panels show momentum space in the molecular plane. Grey points show the positions and momenta of hydrogen atoms, green dots show the positions and momenta of the deuterium atoms and red dots show the position momenta of oxygen atoms. The two horizontal "time" plots show the current time and the green curve depicts the pulse envelope. (Multimedia available online)

In Fig. 7, the same evolution is shown in the molecular frame, i.e., in which the positions and momenta are sampled according to a Wigner distribution in the molecular rest frame. Here the water molecule is initially oriented so that oxygen points on average to the positive x-axis.

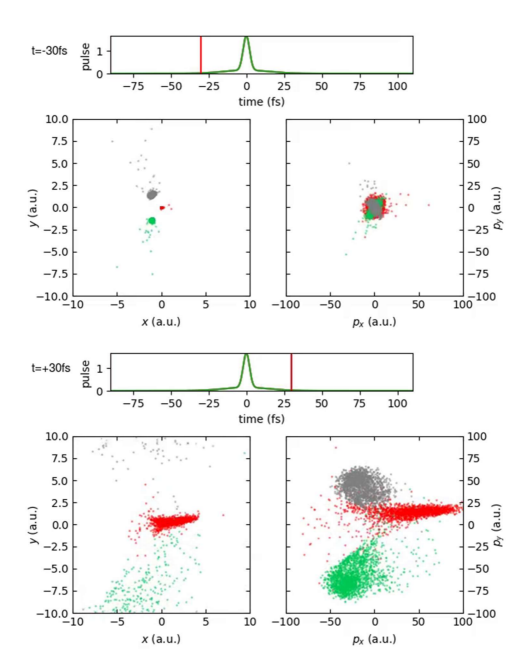


Figure 7. Position and momentum space coordinate in molecular frame for the ensemble of HDO molecules as a function of time. Left panels show position, right panels show momentum space in the molecular plane. Grey points show the positions and momenta of hydrogen atoms, green dots show the positions and momenta of the deuterium atoms and red dots show the positions and momenta of oxygen atoms. The two horizontal "time" plots show the current time and the green curve depicts the pulse envelope. (Multimedia available online)

The discrepancies between Fig. 6 and Fig. 7 are evident. At t=-30 fs, the position distribution resembles the geometry of the water molecule, whereas at t=30 fs, the momenta show different distribution as in the Newton diagram.

The discrepancies between recoil and molecular frame arise largely due to the strong bending dynamics that has a noticeable impact on the final oxygen momentum, and makes a clear interpretation of Newton diagrams in Fig. 1 in terms of fragmentation dynamics challenging. As we also have reported before for H₂O [10], in some cases even overbending occurs, where the HOH angle grows beyond 180 degree, which can be seen in Fig. 7 by the number of trajectories ending up with a final oxygen momentum into negative x-axis direction.

Overall, it becomes visible that the dynamics induced by sequential core ionization involves large variations, caused by the different ionization timings and the different dicationic electronic configurations visited.

To further highlight this large variation, we selected trajectories out of the ensemble, where specific dynamical patterns show up in considerable strength. Figures 8, 9, and 10 show trajectories where symmetric dissociation along the bonds, unbending motion, and strongly asymmetric fragmentation appear, respectively. The color change of the oxygen atom depicts the time at which photoabsorption occurs.

In Fig. 8, the first photoabsorption event triggers elongation of the OH and OD bond and eventually dissociation along these directions into neutral O, D⁺, and H⁺. The second photoabsorption event then ionizes the oxygen atom and accelerates the fragmentation.

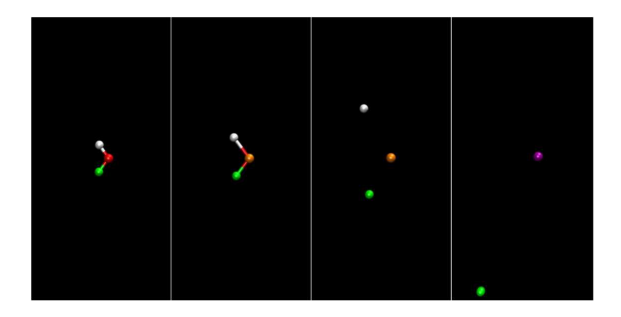


Figure 8. Selected trajectory with dissociation along bonds. White color denotes the hydrogen atom, green colour the deuterium atom. The oxygen atom is depicted in red, pink and orange colors, where the change in color denotes the timings of the two consecutive photon absorptions, and namely red: no photoabsorption, pink: absorption of the first photon, orange: absorption of the second photon (Multimedia available online)

Figure 9 shows and example for strong unbending dynamics. Here the three atoms reach an almost linear geometry when the second photon is absorbed, resulting in a back-to-back emission of D^+ and H^+ as discussed in Fig. 2.

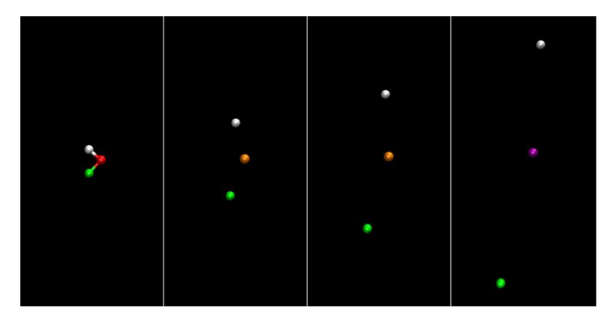


Figure 9. Selected trajectory with large unbending motion along the bonds. White color denotes the hydrogen atom, green color the deuterium atom. The oxygen atom is depicted in red, pink and orange colours, where the change in color denotes the timings of the two consecutive photon absorptions, and namely red: no photoabsorption, pink: absorption of the first photon, orange: absorption of the second photon. (Multimedia available online)

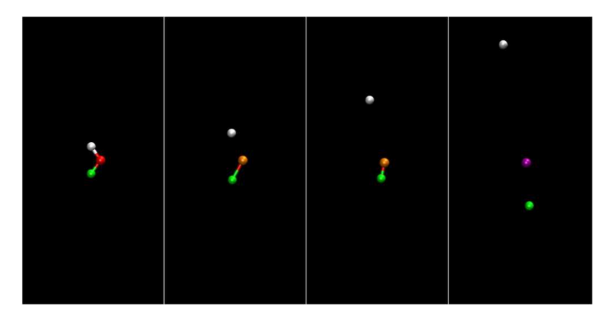


Figure 10. Selected trajectory with strongly asymmetric fragmentation. White color denotes the hydrogen atom, green color the deuterium atom. The oxygen atom is depicted in red, pink and orange colours, where the change in colour denotes the timings of the two consecutive photon absorptions, and namely red: no photoabsorption, pink: absorption of the first photon, orange: absorption of the second photon. (Multimedia available online)

The strong asymmetry displayed in Fig. 10 is an example for a quasi-sequential fragmentation. The first photoabsorption event triggers the release of H^+ while the remaining OD^+ fragment stays bound. Only after the second photoabsorption event, OD^+ is ripped apart into O^{2+} and D^+ . This dynamical pattern leaves a significant larger momentum on H^+ than on D^+ , as highlighted in Fig. 3.

V. CONCLUSION

We have investigated the dynamics triggered by core-shell ionization in three water isotopologues, H₂O, D₂O, and HDO. We show in unprecedented depth isotope effects along the series. In particular, similar structural changes such as asymmetric bond elongation and bond-angle-opening mechanisms occur for all three systems, but with significant differences. Due to the larger mass, dynamical patterns are slower in D₂O. Due to the asymmetry in mass for HDO structural asymmetry arises in the dynamics. A method to identify the sequences of events taking place upon the consecutive absorption of two x-ray photons, the native frame analysis, is also performed for HDO. It indicates that a clear two-step dissociation cannot be identified, possibly because a full sequential fragmentation takes more time than offered by the pulse duration of the ionizing x-ray beam of <25 fs. By combining the use of short and intense x-ray pulses with a detection method able to reveal structural changes in great detail, and including the insight obtained from an advanced theoretical modelling, we have shown differences and similarities in structural dynamics along a series of isotopologues, and gained a deep insight into x-ray induced processes.

VI. ACKNOWLEDGMENTS

We acknowledge European XFEL in Schenefeld, Germany, for provision of x-ray free-electron laser beam time at the SQS instrument and would like to thank the staff for their assistance (proposal # 2150). L.I. and R.S. acknowledge support from the Cluster of Excellence "CUI: Advanced Imaging of Matter" of the Deutsche Forschungsgemeinschaft (DFG)-EXC 2056project ID 390715994. M.I. acknowledges funding from the BMBF (Grant number by 13K22CHA). T.Maz. acknowledge support M.M. and the Forschungsgemeinschaft (DFG) (SFB-925, project No.170620586). Th. W. was supported by the U.S. Department of Energy (DOE), Office of Science, Basic Energy Sciences (BES) under Award No. DE-AC02-05CH11231. M. S., R. G., T.Mar., O. T. and I. I. acknowledge the financial support of the French CNRS GotoXFEL program. N.V. and T. Mar. acknowledge funding from the European Union's Horizon 2020 research and innovation program under the Marie Skłodowska-Curie grant agreement No [860553]. F.T. acknowledges support by the MaxWater initiative of the Max-Planck-Gesellschaft. G.S. acknowledges financial support from the Deutsche Forschungsgemeinschaft Research Training Group DynCAM (RTG 2717) and grant 429805582 (project SA 3470/4-1). D.R. was supported by the DOE Office of Science, Office of Basic Energy Sciences, Chemical Sciences, Geosciences, and Biosciences Division under Contract No. DE-SC0019451. R.F. acknowledges financial support of the Swedish Research Council and the Knut and Alice Wallenberg Foundation, Sweden.

Conflict of Interest

The authors have no conflicts to disclose.

Authors' Contributions

M. N. P. and M. S. originated the project. M. I., T. Maz., R. B., Th. W., S. E., P. G., N. R., T. Mar., N. V., O. T., I. I., J. N., E. K., F. T., M. G., R. F., G. S., D. R., M. Ma., M. Me., M. S., T. P., T. J. and M. N. P. participated in the experimental work, T. J. and R. G. analysed the experimental data. L. I., and R. S. performed the simulations. R. G., L. I., T. P., T. J. and M. N. P. wrote the paper with input from all authors.

Data Availability

Data recorded for the experiment at the European XFEL are available at [34].

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