Strong Field Ionization of Water II:  
Electronic and Nuclear Dynamics En Route to Double Ionization

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We investigate the role of nuclear motion and strong-field-induced electronic couplings during the double ionization of deuterated water using momentum-resolved coincidence spectroscopy. By examining the three-body dicaticonic dissociation channel, D^+/D^+/O, for both few- and multi-cycle laser pulses, strong evidence for intra-pulse dynamics is observed. The extracted angle- and energy-resolved double ionization yields are compared to classical trajectory simulations of the dissociation dynamics occurring from different electronic states of the dication. In contrast with measurements of single photon double ionization, pronounced departure from the expectations for vertical ionization is observed, even for pulses as short as 10 fs in duration. We outline numerous mechanisms by which the strong laser field can modify the nuclear wavefunction en-route to final states of the dication where molecular fragmentation occurs. Specifically, we consider the possibility of a coordinate-dependence to the strong-field ionization rate, intermediate nuclear motion in monocation states prior to double ionization, and near-resonant laser-induced dipole couplings in the ion. These results highlight the fact that, for small and light molecules such as D_2O, a vertical-transition treatment of the ionization dynamics is not sufficient to reproduce the features seen experimentally in the coincidence double-ionization data.

I. Introduction

Investigations into the mechanisms of strong-field ionization (SFI) remain an important cornerstone in ultrafast science research due to its central role in strong-field induced phenomena such as high-harmonic generation [1-3], laser-induced electron diffraction [4-6] and Coulomb-explosion imaging [7,8]. SFI has also been used as a “pump” for experiments that aim to study charge transfer or charge migration in molecular cations [9-12]. For molecular systems this continued interest is in part due to the complexity of the SFI process when compared with atomic systems [13,14]. Critical differences that are partially responsible for this are: the spatial arrangement of the nuclei that give rise to the molecular structure; the presence of internal degrees of freedom, vibrations and rotations; and the significantly higher density of electronic states that are typically found in these systems [15,16].

For many experiments, the ideal scenario is that SFI serves as a “sudden” transition, where the nuclei do not move during the pulse that induces the ionization [17-19]. The electron rearrangement that accompanies sudden transitions can be compared to the “vertical” transitions that describe weak-field perturbative ionization, where a portion of the ground state vibrational wave function is projected onto the electronic states of the mono- or dication. The notion of a vertical transition is based on the idea that the transition dipole moment for the coupling of two electronic states by a weak (perturbative) external field is roughly independent of nuclear coordinate over the extent of the initial wave function. This allows one to factor the coupling matrix element into an electronic term which is multiplied by an overlap of the initial vibrational wave function with the vibrational eigenstates of the upper electronic state, and thus the initial vibrational wave function is simply “vertically lifted” and projected (mapped) onto the excited state potential energy surface (PES). This assumption is frequently described in terms of the Franck-Condon principle [20].

Nuclear motion during the pulse is not the only mechanism for non-vertical ionization. Coordinate-dependent strong-field ionization rates [21,22] and impulsive Raman excitation in one of the electronic states could also result in deviations from vertical ionization [23,24].

Recent work that considered the single-photon double ionization of water showed excellent agreement between the measured and calculated momentum resolved yield of fragment ions assuming vertical ionization [25,26]. Here we explore the extent to which SFI with short pulses can be considered vertical. Using a combination of coincidence velocity map imaging of fragment ions and tra-
jectory calculations for the three body dissociation dynamics of the molecular dication, we demonstrate that even for very short pulses (< 10 fs), the ionization cannot be considered to be “vertical”, but involves reshaping (changes to the first and second moments of the distribution) of the vibrational wavefunction during the ionization dynamics. We discuss different contributions to the wave function reshaping during ionization.

Our measurements, calculations and analysis may help interpret previous work that made use of excited state dynamics [27, 29] and for vibrational wave packet holography [30]. In the case of probing excited state dynamics, while SFI produced qualitatively similar time dependent yields as weak field or single photon ionization, quantitative agreement was not possible [27]. We argue that this is due to variation in SFI rates with nuclear geometry and reshaping of the vibrational wave packet during SFI. While the holography measurements showed very nice agreement between the experiment and theory in terms of the interference fringe positions and visibility, details of the comparison were not perfect due to the limited ion imaging ability and the assumption of vertical ionization. Our work directly examines this assumption and can be thought of as characterizing the instrument response function associated with SFI as a probe. Particularly for early time delays in pump probe experiments, where the initial and final PESs vary significantly with nuclear coordinate, SFI can differ from vertical ionization significantly. Also, for high intensities, one can see enhancement of ionization to higher charge states through charge resonance enhanced ionization, which also violates the notion of vertical ionization [31, 32]. Finally, for pulses longer than 10 fs, experiments and simulations are greatly affected by laser-induced dynamic angular dependence of the ionization dynamics. We discuss different contributions to the laser-coupled ion dynamics simulation through inclusion of a source term in a density matrix approach [30].

II. Experiment

Two similar apparatuses were used to carry out the measurements. Since they are almost the same, we provide a description for the apparatus used to carry out the 10 fs pulse measurements (described in detail in previous work [41, 42]) and indicate any differences for the apparatus used for the 40 fs pulse measurements. Briefly, the output from a commercial amplified Ti:sapphire laser system (1 mJ, 780 nm, 1 kHz) is spectrally broadened using filamentation in Ar gas, and compressed to ~10 fs using chirped mirrors and an acousto-optic pulse shaper [43]. The laser pulses are directed into a vacuum chamber (base pressure of 10^{-10} mbar) and focused at the center of a velocity-map imaging (VMI) spectrometer using an in-vacuum concave silver mirror (f = 5 cm). The in situ intensity was calibrated by SFI of argon, measuring the classical 2U_p cutoff for field-ionized electrons from argon [44] at low pulse energies and extrapolating to higher ones (using a procedure outlined in Ref. [45] for the 40 fs measurements). The estimated intensities for the 10 fs and 40 fs measurements are 400 TW/cm^2 and 600 TW/cm^2, respectively.

Target D_2O molecules are expanded into a separate source chamber and subsequently skimmed to yield an effusive molecular beam [41]. This beam intersects the focused laser at the center of the electrostatic lens stack of the VMI spectrometer [41]. The extracted ions and electrons are recorded using a microchannel plate (MCP), phosphor screen, and TPX3CAM camera [41, 46] with 1.5 ns time resolution (a Roentdek hexanode detector, with a time resolution of <1 ns [47] was used for the 40 fs pulse measurements).

In order to ensure the low count rates required for coincident detection of all charged particles, a working pressure of 4×10^{-10} Torr was used throughout the experiment (with a base pressure of about 1×10^{-10} Torr). The adopted pressure resulted in an average event rate of approximately 0.8 per laser shot, which in turn corresponded to an event rate of <0.1 per shot for the double-ionization channels considered in this work. Presented in Table II are the experimentally extracted yields and branching ratios for the dissociation channels following strong-field double ionization. Two-body fragmentation into OD^+/D^+ is the dominant channel followed by three-body fragmentations into D^+/D^+ + O and D^+/D^+ + O_2. In the present work we focus our discussion on the three body D^+/D^+ + O fragmentation channel. A detailed analysis of fragmentation into OD^+/D^+ was included in an earlier publication, which investigated the roles of dynamic and geometric alignment during water double ionization [48]. We note that our extracted ratios differ significantly in some channels from reported values for single photon double ionization [48] and, more critically, SFI with higher intensity longer duration pulses [49].
TABLE I. Table of electronic states of the water dication in different conventions. Branching ratios are for the dissociation dynamics on the \textit{ab initio} potential surfaces with initial conditions from the Wigner phase space distribution of the ground vibrational state, calculated from $10^5$ trajectories on each potential surface, similar to the calculations in \cite{25, 26}.

| $C_s$ sym | spin state order | $C_{2v}$ sym | $C_{2v}$ config | 2-body branching ratio [\%] | 3-body branching ratio [\%] |
|-----------|----------------|--------------|----------------|-----------------------------|-----------------------------|
| $X^2A^+$ | $T_0$ | $^3B_1$ | $(3a_1b_1)^{-1}$ | 92 | 8 |
| $2^2A^+$ | $T_1$ | $^3A_2$ | $(1b_2b_1)^{-2}$ | 0 | 100 |
| $1^2A^+$ | $T_2$ | $^3B_2$ | $(1b_23a_1)^{-1}$ | 0 | 100 |
| $1^1A^+$ | $S_0$ | $1^1A_1$ | $(1b_1)^{-2}$ | 100 | 0 |
| $1^1A^0$ | $S_1$ | $^1B_1$ | $(3a_1b_1)^{-1}$ | 87 | 13 |
| $2^2A^0$ | $S_2$ | $2^1A_1$ | $(3a_1)^{-1}$ | 26 | 74 |
| $2^2A^+$ | $S_3$ | $^1A_2$ | $(1b_2b_1)^{-2}$ | 0 | 100 |
| $3^2A^+$ | $S_4$ | $^1B_2$ | $(1b_23a_1)^{-1}$ | 0 | 100 |
| $3^2A^+$ | $S_5$ | $3^1A_1$ | $(1b_2)^{-2}$ | 0 | 100 |

TABLE II. Table of branching ratio of relevant water dication dissociation channels. The detection efficiency of different ions are estimated to be 0.58(D$^+$), 0.2(O$^+$) and 0.3(OD$^+$).

thermore, we note the absence of the weak D$_2^+/O^+$ channel observed in Ref. \cite{49} for our data recorded with 10 fs pulses. A detailed investigation into the pulse duration dependence of double ionization branching ratios is beyond the scope of the present work.

III. Trajectory Calculations

Classical trajectory calculations simulate the conditions of single-photon double ionization of the ground state of D$_2$O in which the molecule undergoes a Franck-Condon transition to the doubly ionized excited state. The trajectories are propagated on the potential surfaces of the lowest nine states of the water dication. Those states are created, in the simple molecular orbital picture, by removing two electrons in all possible ways from the highest three filled molecular orbitals of the neutral as indicated in Table I, leaving the D$_2$O$^{++}$ ion in all possible spin states for each configuration. The surfaces were calculated earlier \cite{25, 50} with MOLPRO \cite{51, 52} using internally contracted multireference configuration interaction (icMRCI) methods at the configuration interaction singles and doubles (CISD) level, including the Davidson correction to the CI energy. The full dimensional surfaces were then fitted using a functional form developed by Gervais \textit{et al.} \cite{50}.

The trajectory calculations assumed the neutral water molecule is initially in its ground vibrational state with zero total rotational angular momentum. The initial conditions for $10^5$ trajectories on each potential surface were sampled from the Wigner phase space distribution for the computed normal modes of the ground electronic state in icMRCI calculations. The Phase space distribution has the form

$$W(\mathbf{Q}, \mathbf{P}) = \frac{1}{(\pi \hbar)^{3N-6}} \prod_{j=1}^{3N-6} e^{-\frac{\omega_j}{2}Q_j^2 - \frac{\omega_j}{2}P_j^2}$$

where $N$ is the number of atoms, $\omega_j$ are the associated frequencies, and the vectors $\mathbf{Q}$ and $\mathbf{P}$ are the normal mode coordinates and momenta, respectively, and is positive definite for the case of the ground state. A similar trajectory study was performed for double ionization of H$_2$O previously \cite{25} and compared extensively with experimental momentum images \cite{26} from single photon double ionization measurements using the cold target recoil ion momentum spectroscopy (COLTRIMS) technique \cite{53, 55}. That comparison validated the accuracy...
of this \textit{ab initio} treatment of the dissociation of the cation in all but its finest details. The present calculations differ only in the masses of the atoms (D versus H) and the larger number of trajectories on each surface \((10^5 \text{ versus } 10^3)\). They thus show small quantitative differences from the previous work, for example in the two-body versus three-body dissociation branching ratios.

The potential energy curves along the symmetric stretch coordinate of the first 9 states of dication are shown in the Fig. 1. The state characters are labelled according to \(C_{2v}\) symmetry in that figure. Outlined in Table 1 are the equivalent labelling conventions in other symmetries, together with the dominant electronic configurations for these states near the equilibrium geometry of neutral molecule and the associated two- and three-body theoretical branching ratios for the present case of \(D_2O\). The criterion for categorizing a trajectory as three body in this work was that the R\(_{OD}\) distance of one deuterium be 200 Bohr or greater and that the other reach at least 50 Bohr. All others were categorized as two-body dissociation.

FIG. 2. Measured and calculated \(D^+\)/\(D^+\)/\(O\) yield as a function of angle between the two \(D^+\) ions’ momentum, \(\beta\), and the kinetic energy release (KER). Panel (a) Measured coincidence \(D^+\)/\(D^+\)/\(O\) yield as a function of \(\beta\) and KER for laser parameters described in the text; (b) Simulated \(D^+\)/\(D^+\)/\(O\) yield as a function of \(\beta\) and KER for equal population of each of the first nine states of the dication; (c) simulation \(\beta\)-KER with fit; (d) simulation \(\beta\)-KER with experimental resolution included. The table at the bottom lists the relative fitting populations for each states where the states are labeled with their \(C_{2v}\) symmetry due to lack of the spin information in the observable. Details are outlined in the text.

| State  | \(2^1A_1\) | \(2^1A_1\) | \(2^1A_1\) | \(3^1A_1\) |
|-------|--------------|--------------|--------------|--------------|
| Weight| 1.00         | 0.804        | 0.287        | 0.163        | 0.092        |

IV. Measurements

Given the agreement between the trajectory calculations and the single photon double ionization measurements described above, the question we address here is whether strong-field double ionization prepares a similar superposition of states of the dication, and whether the wave packet launched on each state via SFI is similar to the one launched by single photon (weak field) ionization - i.e., can the SFI process be thought of as vertical? We therefore compare the calculated double ionization yield as a function of \(\beta\) and KER with our measurements. Fig. 2 shows the measured and calculated \(D^+\)/\(D^+\)/\(O\) yield. Panel (a) shows the measured yield as a function of \(\beta\) and KER. Panel (b) shows the calculated yield vs \(\beta\) and KER assuming vertical ionization with equal population of the di-cation states. Panel (c) shows the same results as panel (b) with coefficients for the first 9 states of the dication fitted to achieve the best agreement with the measurements. Panel (d) shows the calculated results accounting for experimental broadening of the features due to the limited resolution of our VMI apparatus. Details of which are discussed below.

In the fitting procedure, due to the lack of the spin state information about the oxygen atom, states of the the same \(C_{2v}\) symmetry (e.g. \(1B_1\) and \(3B_1\)) have been grouped together since they are not separable in the \(\beta\)-KER plot. From now on we just use the word “state” to represent different \(C_{2v}\) symmetries. As can be seen from Fig. 2 each state populates a distinct region (“island”) of the \(\beta\)-KER plot. More details on the island assignments can be found in Refs. 25, 26 as well as in Appendix A. Based on these islands, a simple principle component analysis (PCA) procedure that minimizes the residual was used to reproduce the \(\beta\)-KER plot. Different states population have been fitted and the reconstructed \(\beta\)-KER plot is shown as Fig. 2(c). The fitted relative state population has been listed as the table in the figure as well. The coefficients from the fit are decreasing with increasing ionization potential, roughly in agreement with expectations from a simple quasi-static tunnel ionization model of Ammosov-Delone-Krainov (ADK theory). An interesting observation is that while the relative weights for states that involve removing electrons from different orbitals are in reasonable agreement with predictions from ADK theory, the relative weights for singlet states that involve removing two electrons from the same orbital with opposite spins are significantly higher than predictions based on ADK theory. This observation may indicate cooperative behavior, or non-sequential double ionization to states of the dication that involve removing two electrons from the same orbital.

The fitting in Fig. 2(c) only coarsely matches the measurements shown in Fig. 2(a). Although the experimental yield shows a center of mass similar to the calculations, the distinct state resolved features in the \(\beta\)-KER plot are not captured in the SFI results. An important
The sensitivity of the correlation between initial and final coordinates to electronic state is related to the breakdown of the axial recoil approximation and “slingshot” motion of the initial ground state wavefunction can result in different β distributions than those predicted for single photon ionization. The upper row in Fig. 3 which shows how the final β values depend on the initial coordinates, exhibits a strong state dependence. The \( ^3B_1 \) and \( ^2A_1 \) states show a positive correlation between the final β value and the initial opening angle, while the \( ^1A_2 \) state shows a negative correlation. The behavior with respect to the initial O-D distance is the opposite. This illustrates how reshaping or motion of the initial ground state wavefunction can result in different β distributions than those predicted for single photon ionization. The bottom row in Fig. 3 shows the KER dependence on the initial coordinates. Again, there is a significant dependence of the KER on initial bend angle and O-D distance, illustrating how reshaping or motion of the initial ground state wavefunction can distort the KER distributions predicted for single photon ionization. These plots motivate an examination of the different strong-field mechanisms that can result in non-vertical ionization and the measured β-KER plots. The sensitivity of the correlation between initial and final coordinates to electronic state is related to the breakdown of the axial recoil approximation and “slingshot” motion of the \( D^+ \) ions for the \( ^3B_1 \) and \( ^2A_1 \) states, as discussed in more detail in [25].

The calculated β-KER plot shown in Fig. 2(b) and...
Fig. 2 are results of applying one-photon perturbation theory to describe the transition from the initial neutral directly into the dication states, which is then followed by classical propagation to compute the final fragment energies and angles. In this treatment, the initial Wigner function launched on the dication states is then simply the Wigner function corresponding to the normal-mode ground state of the neutral species, that is, the Franck-Condon wave packet. While this treatment is applicable to weak-field ionization where one-photon perturbation theory is applicable, additional effects are present during SFI that modifies the initial neutral ground-state wave packet before it arrives on the dication surfaces. First, the SFI rate can depend strongly on the nuclear coordinates, an effect which can reshape the initial ground state nuclear wave packet during ionization. Second, multiple ionization via SFI is typically a sequential process where there is a time delay between the ejection of each liberated electron, thereby giving the nuclei a chance to relax and rearrange in between different ionization events. Finally, since there is still a strong-field present during this time delay, near-resonant laser-driven electronic transitions can occur, causing additional non-ionizing electronic transitions that reshape the nuclear wave packets while the molecule is in an intermediate ionic state.

We now outline each of these effects in more detail. We do not attempt a complete treatment of D$_2$O double ionization in strong-fields with all degrees of freedom active, which although desirable, represents a massive theoretical and computational task. Rather, we limit ourselves to outlining each effect using simplified reduced-dimensionality models. Since ionization to both the X and A cation states is expected to occur [30], modifications and dynamics arising from ionization to both X and A are used to exemplify these effects.

**Coordinate-dependence of the SFI rate**: SFI with low-frequency fields can be envisioned as a quasistatic tunnel ionization process where the ionization rate depends exponentially on the ionization potential (IP) through the Keldysh tunneling rate $\Gamma(R, t)$ [57, 58]. In this description, ionization occurs in short sub-cycle bursts near the peaks of the laser oscillations. With the assumption that the nuclei remain stationary during a single ionization burst, the ionized wave packet after the burst can be written as

$$\Psi_K(R, t) = \Gamma(R, t)\Psi_0(R),$$  

where $\Psi_0(R)$ is the initial nuclear wave packet before ionization and R stands for all nuclear coordinates. If ionized from the neutral at equilibrium, $\Psi_0(R)$ is often called the Franck-Condon wave packet. The Keldysh rate is given by

$$\Gamma(R, t) = \mathcal{P} \exp \left[ -\frac{2}{3} \frac{(2 \text{IP}(R))^{3/2}}{|F(t)|} \right]$$  

where $\mathcal{P}$ is a slowly-varying (i.e., non-exponential) prefactor that depends weakly on IP, $F$, and R. Here $F$ denotes the amplitude of the electric field. In molecular systems, the $\mathcal{P}$ prefactor would also encode the molecular orientation dependence of SFI as well as other molecular effect such as enhanced ionization, and accurate computation of $\mathcal{P}$ would require some form of ab initio numerical simulation of the ionization process. In the following we set $\mathcal{P} = 1$ for simplicity and consider only the effects of the Keldysh exponent. Eq. (2) shows that the Keldysh rate can modify the spatial structure of Franck-Condon wave packet through the coordinate-dependence of the IP.

Fig. 4 shows examples of the Keldysh rate modifying the initial Franck-Condon wave packet in D$_2$O. Fig. 4a shows the IPs for the first ionization step from the neutral to the X state of the ion along the bend coordinate $\theta$ together with cuts through the nuclear wave function both with and without applying the Keldysh weighting applied. Fig. 4b plots the same but for the neutral to
A state. While very little change in the wave function occurs for X ionization along this coordinate, it can be seen that the Keldysh weighting has the effect of shifting the Franck-Condon wave packet along the bend coordinate. Panels c and d plot similar cuts for two transitions of the second ionization steps, X → 1^1A_1 (S_0) and X → 3^1A_2 (T_1), but now taken along the symmetric stretch coordinate. Due to the increased magnitude and steep coordinate dependence of the IPs for these transitions, the Keldysh-induced shifts of the wave function are more pronounced compared to the previous single-ionization examples.

**Few-cycle nuclear motion in the ion:** Following the first ionization event, the nuclear wave packet can begin to move on the cationic surfaces before the second ionization occurs. This intermediate motion can change the nuclear wave packets before being projected onto the dicationic states. From our investigations of the cationic surfaces, the dominant motion is expected to be along the bend coordinate. In reality the motion in the cation occurs in the presence of the laser field, but we first consider the effects of field-free motion, which alone can already cause pronounced reshaping of the wave packets. Laser-driven motion in the cation is considered below.

The bending wavepacket dynamics on the X^2B_1 and A^2A_1 cationic surfaces is simulated using the following simplified model of D_2O^+. First, the bond lengths are fixed at the neutral equilibrium values, R_{eq}, throughout the dynamics. Second, the bending is restricted to a single plane of motion, where the overall rotational motion about the center-of-mass is not considered during these wavepacket simulation. Finally, the mass of the oxygen atom is assumed to be infinite, which significantly simplifies the corresponding kinetic energy operator. With these restrictions, the Hamiltonian of the model system is written (in atomic units) as

\[
\hat{H}(\theta,t) = \begin{pmatrix}
-\frac{1}{2\mu R_{eq}^2} \frac{\partial^2}{\partial \theta^2} + V_X(\theta) & 0 \\
0 & -\frac{1}{2\mu R_{eq}^2} \frac{\partial^2}{\partial \theta^2} + V_A(\theta)
\end{pmatrix},
\]

where \( \mu = m_D/2 \) is the reduced mass of the bend coordinate, \( m_D \) is the mass of atomic deuterium, while \( V_X(\theta) \) and \( V_A(\theta) \) are the potential energy surfaces of the X^2B_1 and A^2A_1 states. This Hamiltonian is used to solve the time-dependent Schrödinger equation (TDSE) \( i\partial_t \Psi(\theta,t) = \hat{H}(\theta,t) \Psi(\theta,t) \). In order to investigate the effect of wave-packet motion as separate from the Keldysh effects discussed above, we populate both the X and A surfaces during the ionization event starting from the unchanged Franck-Condon wave packet, and hence the initial condition for \( \Psi(\theta,t) \) at the moment of ionization is taken to be the bending ground state, \( \psi_0(\theta) \), on the neutral surface. The TDSE is solved using the Fourier-split-operator technique [50].

The first three panels of Fig. 5 show the effects of intermediate field-free few-cycle motion on the X and A cation surfaces. Fig. 5f shows the Wigner function of the initial Franck-Condon wave packet. This is the initial state of the bending coordinate that is used in the β-KER trajectory simulations. Figs. 5g and c plot Wigner functions after this initial state has propagated field-free for a time corresponding to 2 laser cycles of the 780 nm field (2×2.6 fs) on the X and A states, respectively. The X-state Wigner has undergone a little acceleration and motion as can be seen by the small shift of the center of the Wigner function, but there is still significant overlap between the propagated and initial Wigner functions in this case. However, the 2-cycle field-free propagation on the A-state substantially modifies the initial Wigner function, which now has effectively zero overlap with the initial state. The remaining panels of Fig. 5 pertain to laser-driven motion in the ion, which is outlined in the following.

**Near-resonant dipole coupling in the ion:** At 780 nm, the laser induces a near-resonance one-photon coupling between the X and A states. This coupling is included in the wave packet simulation by adding off-diagonal dipole
FIG. 6. Time-resolved simulations of the strong-field coupling between the X state of the water cation and the A state in the presence of (a) the 10 fs pulse with $\lambda_0 = 780$ nm and $I_0 = 400$ TW/cm$^2$ and (b) the 40 fs pulse with central wavelength $\lambda_0 = 800$ nm and peak intensity $I_0 = 600$ TW/cm$^2$ (parameters chosen to match the experiments). In each case, the cation is initiated in the X state (red/blue solid line) at a local peak in electric field (solid green line) that roughly corresponds to the point at which the pulse intensity is half its maximum value: $t/\tau \approx 0.5$. Due to the presence of the field, the X population couples to the A state (red/blue dashed line). Plotted below each of these figures is the probability distribution over bend angle, $\theta$, for the mixture of states displayed above, shown separately for (c) the case of a 10 fs pulse (blue) and (d) the case of a 40 fs pulse (red). In each case, dotted gray lines denote the equilibrium bend-angles for the X and A states. In all four panels, green shading roughly indicates the window in time over which the second ionization event in sequential double-ionization is expected to occur.

terms to the Hamiltonian, which then becomes

$$\hat{H}(\theta, t) = \begin{bmatrix} -\frac{1}{2\mu R_\theta^2} \frac{\partial^2}{\partial \theta^2} + V_X(\theta) - F(t) \cdot d_{XA}(\theta) & -F(t) \cdot d_{XA}(\theta) - \frac{1}{2\mu R_\theta^2} \frac{\partial^2}{\partial \theta^2} + V_A(\theta) \end{bmatrix},$$

(5)

where $d_{XA}(\theta)$ is the transition dipole between these states, and $F(t)$ is the electric field of the laser which is chosen to be parallel to $d_{XA}(\theta)$, i.e., perpendicular to the molecular plane. The electric field of the laser is taken to have a Gaussian envelope

$$F(t) = F_0 \exp \left[ -4 \ln 2 \left( \frac{t}{\tau \sqrt{2}} \right)^2 \right] \cos(\omega_0 t)$$

(6)

where $\omega_0$ is the carrier frequency, $F_0$ is the peak electric field magnitude, and $\tau$ is the full width at half maximum of the corresponding intensity envelope $|F(t)|^2$.

The effects of the dipole coupling on the Wigner functions can be seen in Figs. 5(d)-(g), while a more complete picture of the laser-driven dynamics is shown in Fig. 6. We first consider the Wigner functions. Figs. 5(d) and (e) plot the Wigner functions on the X and A states respectively after the system was initialized with the Franck-Condon wave packet on the X state at the peak ($t=0$) of a laser pulse with $\tau = 10$ fs and intensity of 400 TW/cm$^2$, and allowed to propagate for 2 cycles of the laser. Initially only the X state is populated, but due to the near-resonant dipole coupling some population is transferred to the A state, which is discussed further below. In addition to the modifications due to field-free propagation, seen in Figs. 5(b) and (c), both of the Wigner functions on the X and A states have acquired additional structures after the laser-driven propagation. These structures arise due to light-induced potentials created by the strong near-resonant laser field that modify the field-free potential energy surfaces, an effect known as bond softening [60, 61]. Additionally, the transfer of population from one state to the other, which occurs through cascaded one-photon transitions between the X and A states induced by the strong near-resonant laser, also contributes to these structures. Figs. 5(f) and (g) plot Wigner functions for the same scenario but now with the initial population starting on the A state. Again, it is seen that new structure not present in the Franck-Condon wave packet have developed.

In order to put things into perspective, we compare the amount of X to A coupling that occurs in a 10 fs pulse to a significantly longer pulse with a duration of 40 fs. Fig. 6 shows a broader picture of the laser-driven dynamics. The top panels show the population of ground and first excited states of the monocation together with the laser field as a function of time for 10 fs (top left)
and 40 fs (top right) laser pulses. The bottom two panels show the corresponding probability density as a function of D-O-D angle and time. The figures illustrate the fact that while population transfer and nuclear dynamics can take place during a 10 fs pulse, they have a much more dramatic effect for a 40 fs pulse.

The calculations described above suggest that in a longer pulse, dynamics in the monocation en route to the dication during a longer pulse also leads to significant unbending and a $\beta$-KER plot that is shifted to larger angles. We therefore compare the $\beta$-KER plot for 10 and 40 fs pulses as a test of the conclusions of the theoretical results. Fig. 7 shows this comparison, which bears out the predictions based on the calculations above. We note that the comparison shows that 40 fs tend to have larger $\beta$ angles and also the lower KER. The former is what we expected from the simulation while the latter is not easy to explain. Nevertheless, this comparison provides strong evidence that while there is rough agreement between the measured and calculated momentum resolved fragment ion yields for sub 10-fs pulses assuming vertical ionization, reshaping of the wave packet and nuclear dynamics during ionization can lead to significant differences between strong- and weak-field ionization. And one needs to understand the strong-field dynamics in order to predict the reaction products in an SFI measurement. Finally, a weak diffuse feature at higher KER ($\sim$ 20 eV) and smaller $\beta$ values is observed for 40-fs pulses, which is notably absent for the short pulse case. This feature can readily be assigned to trication Coulomb explosion and is discussed in detail in Appendix C.

VI. Conclusion

In conclusion, while we find rough agreement between measurements of the $D^+$/D$^+$/O yield as a function of $\beta$ and KER for double ionization of D$_2$O with 10 fs laser pulses and calculations of the same yield assuming weak-field/vertical ionization, there are significant differences in the details. As summarized in the Fig. 8, these differences may be ascribed to wave packet reshaping by the strong-field of the laser pulse, and nuclear dynamics during the pulse. These result in SFI being a non-vertical transition, with an understanding of the strong-field dynamics being important for the calculation of the fragment ion yield momentum distributions.

Our calculations of the wave packet reshaping suggest that the coordinate dependence of the tunnel ionization rate plays an important role in reshaping the initial wave packet, particularly for the second ionization step in a sequential double ionization process. This is a result of the dication having a larger coordinate dependence to the ionization potential. Calculations of the monocation dynamics in the presence of the strong laser field indicate that field dressed nuclear dynamics can also reshape the vibrational wave packet. The reshaped wave packet is then projected onto the dication potential energy surfaces toward the end of the pulse. These observations are supported by measurements of the double ionization yield as a function of $\beta$ and KER for 10 fs and 40 fs pulses. This comparison has significant bearing on the use of SFI as a probe of molecular structure and dynamics, and indicates that while very short pulses (< 10 fs) can minimize nuclear dynamics during the pulse, wave packet reshaping by the strong field of the laser can still result in significant distortion of the initial wave function of the molecule. We believe this work will have important consequences for pump-probe techniques, such as time-resolved Coulomb explosion imaging, that aim to track nuclear dynamics during excited state photochemical processes.

These results showcase the need for a more comprehensive theoretical description of SFI processes, which include the role of nuclear motion occurring during the ionizing laser pulse duration. As outlined in the introduction, such nuclear dynamics could have implications for computations of SFI from multi-cycle pulses when invoking the frozen-nuclei approximation, such as recent state-of-the-art time-dependent R-matrix ionization computations for H$_2$O that highlighted modifications to the angular dependence of ionization when significant laser coupling in the ion is present for the frozen-nuclei case. Fur-
ther experimental investigations into the pulse shape and intensity dependence of SFI processes in water may help to target future modelling of these dynamics.

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A. Individual dication state contributions to \(\beta\)-KER plot

In order to try to connect the observed dissociation features in our strong-field double ionization experiment to specific dication states we adopted a similar methodology to Refs. 25, 26. In those studies, the \(H^+ / H^+ / O\) yield following single photon double ionization of \(H_2O\) were compared against classical trajectory simulations on various electronic states of water dication. By comparing theoretical kinetic energy releases (KERs) and relative angle correlations (between the two \(H^+\) fragments) to the experimental data, it was unambiguously shown that specific features could be connected to dissociation occurring on different electronic states of the dication.

Classical trajectory simulations of three-body dissociation of \(D_2O^{2+}\) into \(D^+ / D^+ / O\) were performed using a methodology briefly outlined in Sec. III. In Fig. 3 theoretical KERs and \(D^+ / D^+\) relative angles, \(\beta\), from these calculations are shown alongside the experimental data. In order to connect the features observed in theory to the dication states that are expected undergo three-body decay (see Table I), Fig. 9 shows the \(\beta\)-KER plot with annotated regions corresponding to specific final states. For clarity we adopt the same state specific color scheme outlined in Fig. 3.

B. Alternative representations of the \(D^+ / D^+ / O\) channel

In Sec. II it was briefly outlined that by exploiting the TPX3CAM camera in conjunction with a voltage switching VMI apparatus it is possible to coincidentally detect all charge particles (electrons and ions) following strong-field double ionization, provided a suitably low number (<1) of double-ionization events per laser shot is achieved. For the ions, the time-stamping capabilities of the camera permit the three-dimensional vector momenta of all fragments to be extracted. This information permits observables, such as the relative angles between fragments or total KERs, to determined. As is outlined in the main text of manuscript, as well as Appendix A, a particularly useful way to view the data associated with the \(D^+ / D^+ / O\) channel is to consider the relative angle between the two \(D^+\) ions, \(\beta\), as a function of KER. We note, however, that there exists a large number of possible representations of the data due to three-dimensional correlated information extracted during the experiment. To highlight this, Fig. II shows various cuts of the full data set along the KER, \(\beta\), \(E_{D_1}\) and \(E_{share}\) coordinates. Here \(E_{D_1}\) and \(E_{share}\) refer to the kinetic energy of the one of deuterons and the sharing of kinetic energy between both the deuterons, respectively. In Fig. III the equivalent plots for the theoretical simulations outlined in Sec. III are presented.

Similar to the discussion of the \(\beta\)-KER plot in the main text there exists some coarse agreement between theory and experiment in several of the panels. However, it is apparent that the well-resolved features in Fig. II are absent in the experimental data. This is likely due to the mechanisms outlined in Sec. V. A particularly noteworthy region of disagreement is observed in the panel where the yield is plotted as a function of \(E_{share}\) and \(\beta\). A weak feature extending over all angles at \(E_{share} = 0\) and 1 is seen experimentally but has no analogous signature in the theory. In Fig. II a zoomed \(\beta\)-\(E_{share}\) plot is presented.

FIG. 9. Transparent theoretical \(\beta\)-KER plot generated from the simulations outlined in Sec. III. Annotations of the specific states associated with each distinct spectral feature are shown (ovals). A state dependent color coding consistent with Fig. 3 is adopted for clarity.
to highlight this discrepancy. We attribute this large angular spread in \( \beta \) to a process by which \( \text{D}_2\text{O}^{2+} \) initially undergoes two-body decay into \( \text{D}^+/\text{OD}^+ \) but subsequently fragments into three-bodies (\( \text{D}^+/\text{D}^+/\text{O} \)) via a sequential break-up mechanism. A corresponding feature was observed in the \( \text{H}^+/\text{H}^+/\text{O} \) yield following single-photon double ionization \cite{[26]} and has also been observed in SFI induced fragmentation of triatomic molecules such as OCS \cite{[62]}.

C. Signatures of trication formation in 40-fs pulse data

The comparison of the \( \beta \)-KER plot for the cases of short (10 fs) and long (40 fs) pulse durations (see Fig. \ref{fig:7}) provided experimental support for the role of nuclear motion occurring during double ionization process. These data highlighted two significant differences. The first is a shifting of the feature observed at \( \sim 7.5 \) eV to higher \( \beta \) angles. This, as discussed in the main text, can be attributed to unbending dynamics that result in the water dication undergoing fragmentation from geometries close to linear. A second, higher KER, feature is discernible in the 40 fs data but is notably absent for the case of shorter pulses.

In Fig. \ref{fig:13} we present \( \beta \)-angle-integrated KER spectra for the two pulse durations. A striking difference is observed between the spectra, with the majority of the 40 fs counts originating from the feature centred at \( \sim 20 \) eV. Given the significantly higher KER, a likely origin of this is fragmentation occurring from a more highly charged water ion and a missing coincidence partner ion (due to the limited detection efficiency of our apparatus). In order to verify this, we examined the \( \text{D}^+/\text{D}^+/\text{O}^+ \) coincidence channel and can unambiguously assign the origin of the higher KER feature to such a process.

\begin{thebibliography}{99}

\bibitem{[1]} P.-M. Paul, E. S. Toma, P. Breger, G. Mullot, F. Augé, P. Balco, H. G. Muller, and P. Agostini, Observation of a train of attosecond pulses from high harmonic generation, Science \textbf{292}, 1689 (2001).

\bibitem{[2]} T. Popmintchev, M.-C. Chen, D. Popmintchev, P. Arpin, S. Brown, S. Ališauskas, G. Andriukaitis, T. Baldiunas, O. D. Mücke, A. Pugzlys, \textit{et al.}, Bright coherent ultrahigh harmonics in the keV X-ray regime from mid-infrared femtosecond lasers, Science \textbf{336}, 1287 (2012).

\bibitem{[3]} J. Marangos, Development of high harmonic generation spectroscopy of organic molecules and biomolecules, J. Phys. B: At. Mol. Phys. \textbf{49}, 132001 (2016).

\bibitem{[4]} M. Meckel, D. Comtois, D. Zeidler, A. Staudte, D. Pavičić, H. Bandulet, H. Pépin, J. Kieffer, R. Dörner, D. Villeneuve, \textit{et al.}, Laser-induced electron tunneling and diffraction, Science \textbf{320}, 1478 (2008).

\bibitem{[5]} K. Amini, M. Sclafani, T. Steinele, A.-T. Le, A. Sanchez, C. Müller, J. Steinhof, L. Yue, J. R. S. Saavedra, M. Hemmer, \textit{et al.}, Imaging the Remmer–Teller effect using laser-induced electron diffraction, Proc. Natl. Acad. Sci. \textbf{116}, 8173 (2019).

\bibitem{[6]} C. I. Blaga, J. Xu, A. D. DiChiara, E. Sistrunk, K. Zhang, P. Agostini, T. A. Miller, L. F. DiMauro, and C. Lin, Imaging ultrafast molecular dynamics with laser-induced electron diffraction, \textit{Nature} \textbf{483}, 194 (2012).

\bibitem{[7]} H. Stapelfeldt, E. Constant, H. Sakai, and P. B. Corkum, Time-resolved coulomb explosion imaging: A method to measure structure and dynamics of molecular nuclear wave packets, Phys. Rev. A \textbf{58}, 426 (1998).

\bibitem{[8]} I. Bocharova, R. Karimi, E. F. Penka, J.-P. Brichta, P. Lassonde, X. Fu, J.-C. Kieffer, A. D. Bandrauk, I. Litvinyuk, J. Sanderson, \textit{et al.}, Charge resonance enhanced ionization of CO2 probed by laser coulomb explosion imaging, Phys. Rev. Lett. \textbf{107}, 063201 (2011).

\bibitem{[9]} P. M. Kraus, B. Mignolet, D. Baykusheva, A. Rupenyan, L. Horný, E. F. Penka, G. Grassi, O. I. Tolstikhin, J. Schneider, F. Jensen, \textit{et al.}, Measurement and laser control of attosecond charge migration in ionized iodoacetylene, Science \textbf{350}, 790 (2015).

\bibitem{[10]} M. Kürbel, R. Siemering, C. Burger, N. G. Kling, H. Li, A. Alnaser, B. Bergues, S. Zherbtsov, A. Azzeer, I. Ben Itzhak, \textit{et al.}, Steering proton migration in hydrocarbons using intense few-cycle laser fields, Phys. Rev. Lett. \textbf{116}, 193001 (2016).

\bibitem{[11]} M. Sabbar, Y.-J. Chen, A. K. Pymer, Z.-H. Loh, S. G. Sayres, S. Pabst, R. Santra, and S. R. Leone, State-resolved attosecond reversible and irreversible dynamics in strong optical fields, Nat. Phys. \textbf{13}, 472 (2017).

\bibitem{[12]} K. Ramasesha, S. R. Leone, and D. M. Neumark, Real-time probing of electron dynamics using attosecond time-resolved spectroscopy, Annu. Rev. Phys. Chem. \textbf{67}, 41 (2016).

\bibitem{[13]} T. K. Kjeldsen and L. B. Madsen, Strong-field ionization of diatomic molecules and companion atoms: Strong-field approximation and tunneling theory including nuclear motion, Phys. Rev. A \textbf{71}, 023411 (2005).

\bibitem{[14]} K. Lin, X. Jia, Z. Yu, F. He, J. Ma, H. Li, X. Gong, Q. Song, Q. Ji, W. Zhang, \textit{et al.}, Comparison study of strong-field ionization of molecules and atoms by bicircular two-color femtosecond laser pulses, Phys. Rev. Lett. \textbf{119}, 203202 (2017).

\bibitem{[15]} M. Lezius, V. Blanchet, D. Rayner, D. Villeneuve, A. Stolow, and M. Y. Ivanov, Nonadiabatic multielectron dynamics in strong field molecular ionization, Phys. Rev. Lett. \textbf{86}, 51 (2001).

\bibitem{[16]} T. K. Kjeldsen, C. Z. Bissgaard, L. B. Madsen, and H. Stapelfeldt, Influence of molecular symmetry on strong-field ionization: Studies on ethylene, benzene, fluoro-benzene, and chloro-fluoro-benzene, Phys. Rev. A \textbf{71}, 013418 (2005).

\bibitem{[17]} L. Holmgaard, J. L. Hansen, L. Kalhej, S. L. Kragh, H. Stapelfeldt, F. Filsinger, J. Küpper, G. Meijer,
FIG. 10. Representations of the multi-dimensional D⁺/D⁺/O coincidence data along various coordinates. Shown are KER, β, E_D1 and E_share cuts of the data along each of these coordinates in a matrix style plot. Here the diagonal represents the one-dimensional normalized line-out along each of the aforementioned axes.

FIG. 11. Cuts of theoretical results along the same coordinates outlined in Fig. 10 for the D⁺/D⁺/O fragmentation channel in the classical trajectory simulations. Details of constraints used to classify the trajectories into specific channels are outlined in Sec. III.

D. Dimitrovski, M. Abu-Samha, et al., Photoelectron angular distributions from strong-field ionization of oriented molecules, Nat. Phys. 6, 428 (2010).
[18] F. Schell, A. E. Boguslavskiy, C. P. Schulz, S. Patchkovskii, M. J. Vrakking, A. Stolow, and J. Mikosch, Sequential and direct ionic excitation in the strong-field ionization of 1-butene molecules, Phys. Chem. Chem. Phys. 20, 14708 (2018).
[19] P. Sándor, V. Tagliamonti, A. Zhao, T. Rozgonyi, M. Ruckenbauer, P. Marquetand, and T. Weinacht, Strong field molecular ionization in the impulsive limit: Freezing vibrations with short pulses, Phys. Rev. Lett.
FIG. 12. Zoomed version of the $\beta$-E$_{\text{share}}$ plots shown in Figs. 10 and 11. A broad feature in angle is observed at $E_{\text{share}} < 0.2$ experimentally (a), which in notably absent in the theoretical results (b). This feature can be assigned to a sequential dissociation pathway involving predissociation of OD$^+$ as discussed in the main text.

FIG. 13. Kinetic Energy Release (KER) distribution of the D$^+$/D$^+$ dissociation channel for 10 (blue) and 40 fs pulse durations (red) and D$^+$/D$^+$/O$^+$ only for 40 fs pulse (yellow). The two D$^+$/D$^+$ lineouts correspond to $\beta$-angle-integrated versions of the data shown in Fig. 7. The overlap between D$^+$/D$^+$/O$^+$ channel and the D$^+$/D$^+$ in the 40 fs data indicates the high KER region are coming from events missing an O$^+$.
of H$_2$O in ultrashort intense near-IR laser fields by the time-dependent adiabatic state method and the time-dependent configuration interaction method, Chem. Phys. Lett. 742, 137165 (2020).

35. A. Howard, C. Cheng, R. Forbes, G. McCracken, W. Mills, V. Makhija, M. Spanner, T. Weinacht, and P. Buckbaurn, Strong-field ionization of water: Nuclear dynamics revealed by varying the pulse duration, Phys. Rev. A 103, 043120 (2021).

36. C. Cheng, R. Forbes, A. J. Howard, M. Spanner, P. H. Buckbaurn, and T. Weinacht, Momentum-resolved above-threshold ionization of deuterated water, Phys. Rev. A 102, 052813 (2020).

37. J. Benda, J. Gorfinikiel, Z. Maˇ s´ in, G. Armstrong, A. Brown, D. Clarke, H. van der Hart, and J. Wragg, Perturbative and nonperturbative photoionization of H$_2$ and H$_2$O using the molecular R-matrix-with-time method, Phys. Rev. A 102, 052826 (2020).

38. M. Richter, M. Lytova, F. Morales, S. Haessler, M. A. Dugan, J. X. Tull, and W. S. Warren, High-