Strong field ionization to multiple electronic states in water

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High harmonic spectra show that laser-induced strong field ionization of water has a significant contribution from an inner-valence orbital. Our experiment uses the ratio of H$_2$O and D$_2$O high harmonic yields to isolate the characteristic nuclear motion of the molecular ionic states. The nuclear motion initiated via ionization of the highest occupied molecular orbital (HOMO) is small and is expected to lead to similar harmonic yields for the two isotopes. In contrast, ionization of the second least bound orbital (HOMO-1) exhibits itself via a strong bending motion which creates a significant isotope effect. We elaborate on this interpretation by simulating strong field ionization and high harmonic generation from the water isotopes using the time-dependent Schrödinger equation. We expect that this isotope marking scheme for probing excited ionic states in strong field processes can be generalized to other molecules.

The superposition of several ionic states resulting from strong field ionization of molecules has been observed via high harmonic generation (HHG). Tunnel ionization preferentially ionizes the HOMO. This leaves the molecule in the ground ionic state. Rotational or vibrational pre-excitation is typically used to force ionization from inner valence orbitals [1–3]. The population of ionic states is then probed by the recombination step of HHG [4–6]. In this contribution we show that strong field ionization of water can populate several ionic states without vibrational or rotational pre-excitation. This is done with an isotropic, unexcited ensemble by using isotope marked molecules. The excitation of lower lying valence states triggers an electronic wave packet with femtosecond to attosecond time scale dynamics [7], which was demonstrated recently for the case of fine-split ionic states of Krypton [8].

We investigate ionization of the inner valence orbital by comparing the high harmonic yields from different isotopes of water. As first understood by Lein and observed by Baker et al., nuclear dynamics occurring between ionization and recombination are mapped onto the harmonic spectrum and can be isolated by using isotope marked molecules [9, 10]. Thus, the presence of multiple ionic states in HHG can be determined if the different ionic states launch significantly different nuclear motion.

Figure 1 presents an intuitive picture of the theory elaborated in [9]. The ionization of the molecular orbital (either HOMO or HOMO-1 in Fig. 1(a)) launches a vibrational wave packet on an ionic potential energy surface in Fig. 1(c) (red and green, respectively). The harmonic light is emitted as the returning electron recombines with the molecular ion. Photoionization and HHG recombination are inverse processes [11]. The well known modulation of extreme ultraviolet photoionization cross sections by the Franck-Condon overlap (shown in Fig. 1(b) for H$_2$O) then requires the HHG recombination cross section to be modulated by the overlap of the ionic state nuclear wave packet and the neutral vibrational ground state. As the ionic state nuclear wave packet moves, it loses overlap with the ground state. The nuclear wave packet for the heavy isotope is slow, thus maintaining a stronger overlap than the lighter, faster isotope. This gives a larger harmonic yield for the heavier isotope. The timescale of the nuclear motion is mapped onto the energy of different harmonics [9]. Comparison of

![Figure 1: (a) Constant amplitude contours of the HOMO (1b$_1$) and HOMO-1 (3a$_1$) orbitals. (b) Photoelectron spectrum of H$_2$O reproduced from [12]. The transition to the ground ionic state (2B$_1$) occurs near 12.6 eV and populates only a few vibrational lines. In contrast, the transition to the first excited ionic state (2A$_1$), which occurs around 14.8 eV, populates many vibrational lines. (c) Potential energy curves of the neutral and first two ionic states. The minimum of the 2B$_1$ surface lies almost directly above that of the neutral, while the minimum of the 2A$_1$ surface is at 180$^\circ$.](image)
harmonic spectra from the different isotopes isolates the effect of nuclear motion, since their electronic structure is identical.

In H$_2$O, population of different ionic states can launch significantly different vibrational motion. Ionization of the HOMO, 1$b_1$, excites the molecule into the ionic ground state, $X^2B_1$. The resulting potential energy surface [Fig. 1(c)] is very similar to that of the neutral as can be seen from the exponentially decaying Franck-Condon progression of the $X^2B_1$ state in the photoelectron spectrum shown in Fig. 1(b). This is due to the non-bonding properties of the 1$b_1$ orbital evident by its shape: the 1$b_1$ is a single oxygen $p$ orbital that sticks out of the molecular plane [Fig. 1(a)]. In turn, only extremely weak nuclear dynamics are induced [12, 13]. In contrast, ionization of the HOMO-13$a_1$, which sends the molecule into the $A^2A_1$ state, strongly excites the bending mode [12, 13]. The equilibrium bond angle changes to 180° [Fig. 1(c)]. The 3$a_1$ has an intramolecular bonding character, see Fig. 1(a). The photoelectron spectrum illustrates the strong (weak) vibrational excitation upon ionization of the 3$a_1$ (1$b_1$) orbital. The vertical binding energies of the 3$a_1$ and 1$b_1$ orbitals are 14.8 and 12.6 eV, respectively [14].

The very weak vibrational wave packet excited via population of the ionic ground state indicates that HHG from the 1$b_1$ should not display a strong isotope effect. This is confirmed by a recent simulation, which considered ionization solely from the 1$b_1$ and found a negligible difference between H$_2$O and D$_2$O high harmonics for vibrationally unexcited neutral molecules [15].

Several experiments have shown that inner valence orbitals can contribute significantly to high harmonic yields [1-3, 16]. We expect the inner valence 3$a_1$ orbital to contribute to the high harmonics of water. The broad vibrational wave packet launched via ionization of the 3$a_1$ orbital causes a rapid decrease in the overlap between the ionic and ground state nuclear wave functions. This results in a reduction of the HHG yield and significant difference between H$_2$O and D$_2$O harmonics. Water is therefore an ideal test molecule to determine inner valence contributions via nuclear motion. Harmonic generation in water has been demonstrated before in both liquid and gas phase samples [17, 18].

We generate harmonics with a commercial Ti:Sapphire laser (pulse duration 30 fs, pulse energy 100 µJ, central wavelength 800 nm, repetition rate 1 kHz). A 40 cm focal length mirror focuses the beam to a full width at half maximum of about 60 µm. The harmonics between 20 and 70 eV pass through an Al filter (thickness 100 nm) onto a toroidal flat-field grating, which disperses the radiation onto an extreme ultraviolet sensitive CCD camera. The gas cell is approximately 0.5 mm thick and has entrance/exit holes of approximate diameter 0.2 mm. Water and heavy water reservoirs kept at room temperature are connected to the cell by a fine-dosing valve to keep the cell gas pressure of about 3 mbar (number density $10^{17}$ cm$^{-3}$). The laser intensity is changed by inserting pellicle beam-splitters into the beam. An iris placed just upstream of the grating rejects the long trajectory harmonics [19].

Figure 2(a) shows the experimental ratios of heavy water to water spectra for several intensities. The error bars include the standard error of the mean of the individual datasets, but are dominated by a systematic effect characterized by taking two different sets of spectra at different times. (b) Simulated ratios. Markers represent simulations including both the 1$b_1$ and 3$a_1$ orbitals for several intensities. Lines represent simulations that solely include the 1$b_1$. See text for description of the theoretical error bars.
effectively keeping the slope of the curve constant.

These results show that the D₂O/H₂O harmonic ratio gets progressively larger for longer electron travel times. Thus strong field ionization launches significant and fast nuclear motion in water. As discussed above, the photoionization studies as well as the disagreement with the 1b₁-based simulation in [15] suggest that ionization of the 3a₁ is the dominant source of this motion.

To model the experiment, we extend the theoretical approach used in [15] by taking into account multiple electronic states from different molecular orbitals. Furthermore, we solve the time-dependent Schrödinger equation under the single-active-electron approximation (SAE) with fixed nuclei separately for each orbital. Our approach has been used successfully for H₂ [20] as well as N₂, O₂, and CO₂ [21]. The resulting spectrum is then multiplied by the nuclear autocorrelation function to include the nuclear motion.

We now briefly outline this theoretical method, for details see references [20, 21]. Field-free Kohn-Sham orbitals are obtained using the LB94 exchange-correlation potential [22]. They are expressed in a multi-centered basis [23], which consists of a set of (typically atom-centered) local spheres (bases) defined as the product of spherical harmonics and a radial part expressed in B-splines. Molecular symmetry (for water C₂ᵥ) is accounted for by implicitly generating equivalent local basis sets at symmetry-equivalent positions. A large central sphere positioned at the charge center of the molecule defines an additional large B-spline basis. It overlaps with all atomic-centered spheres and serves for an improved description of the chemical bonding and the molecular electronic continuum. The latter is discretized with a density determined by the size of the central sphere. The radius of the central sphere is set to r₀ = 121.5 a.u., the maximum number of angular momenta used is l max = 12 and the experimentally determined equilibrium geometry (R OH = 0.958 Å, ∠HOH = 104.5° [24]) is applied. The time-dependent electronic wavefunction |Ψ α(t)⟩ is represented as the sum of the Kohn-Sham orbitals multiplied by time-dependent coefficients. The index α refers to the orbital that is initially occupied. All Kohn-Sham orbitals are propagated in the velocity gauge during their interaction with a 10 cycle cos² laser pulse centered at 800 nm. The HHG spectrum for a single molecular orientation is then:

\[ \bar{S}_α(ω) = | \int_{-∞}^{+∞} dt e^{iωt} \vec{d}_α(t)f(t) |^2 \]  \hspace{1cm} (1)

where \( \vec{d}_α(t) = \langle Ψ_α(t)|\vec{p} + \vec{A}(t)|Ψ_α(t)\rangle \). The canonical momentum \( \vec{p} \) and the vector potential \( \vec{A}(t) \) are expressed in a.u. The damping function \( f(t) \) reduces \( \vec{d}_α(t) \) at the extreme times to minimize edge effects in the Fourier transform.

Since the experimental sample is unaligned, we average simulated spectra from many different angles between the molecular axis and the laser polarization. This is performed for each orbital separately. Both coherent and incoherent averaging give essentially the same ratio \( D₂O/H₂O \). Averaging over only the three main axes of water (including the main ionization axes for the HOMO, HOMO-1 and HOMO-2) approximates the ratio obtained with about 80 orientations very well. Following Lein [9], the influence of nuclear dynamics is included only by the autocorrelation function \( C_α(t) \) defined as

\[ C_α(t) = \langle φ_α(0)|φ_α^*(t) \rangle \] \hspace{1cm} (2)

where \( |φ_α⟩ \) denotes the vibrational ground state of H₂O and \( |φ_α^*(t)⟩ \) denotes the time-dependent nuclear wave packet launched on H₂O⁺. Data for the autocorrelation functions is taken from Reutt et al [13]. Finally, we form the incoherent sum of the orientation-averaged HHG contributions \( S^{(D₂O/H₂O)}(ω) \) from the HOMO \( S_{1b₁}(ω) \) and the HOMO-1 \( S_{3a₁}(ω) \) (the HHG contribution of the HOMO-2 1b₂ is found to be negligible) multiplied by the respective autocorrelation function

\[ S^{(D₂O/H₂O)}(ω) = \sum_{α=1b₁,3a₁} |C_α^{(D₂O/H₂O)}(ω)|^2 S_α(ω) \] \hspace{1cm} (3)

where we map the time of the autocorrelation function to energy by using equation (12) in [9]. This restricts the spectrum to energies below the cutoff. The ratio \( D₂O/H₂O \) is then obtained by \( S^{(D₂O)}(ω)/S^{(H₂O)}(ω) \).

Uncertainty in the theoretical ratios was determined by varying several parameters. First, three different damping functions were used. In each case the high harmonic spectra are additionally convolved with a Gaussian distribution function for four different variances chosen to maintain the basic peak structure of the spectra. Finally, the ratios using both peak heights and areas were computed for each case.

Figure 2(b) shows simulations using the same intensities as the experiment. In agreement with [15], the isotope effect calculated using only the 1b₁ orbital (shown with lines) is very small in comparison to the measurement. The two orbital simulation (shown with markers) gives a much larger effect, confirming our qualitative estimate that the 3a₁ plays an important role. For example, at 45 eV, the ratio predicted by the two orbital simulation, approximately 1.10, is much closer to the measured value of 1.15 than that of our single orbital simulation, 1.04.

The intensity dependence of the harmonic spectra is also telling. As stated above, the experimental slope had no discernible dependence on intensity. In contrast, increasing the intensity in the single orbital simulation decreases the isotope ratio for each harmonic, effectively reducing the slope of the curve. This is expected from the single orbital picture due to the one-to-one mapping between travel time and isotope ratio. In agreement with
the measurement, the two orbital simulation does not exhibit a discernible change in slope with intensity, even though the intensity range used is extremely large.

Examination of the simulations suggests that the independence of the slope on intensity stems from an angular ionization saturation effect. Prior to the laser pulse, the angular probability distribution is isotropic. For a given molecule, the primary ionization direction is determined by the intensity range used is extremely large. The intensity range used is extremely large. The intensity range used is extremely large.

The HHG spectrum of the 3a1 orbital by itself provides an estimate of the upper bound for the isotope effect and is shown in the inset in Fig. 3. A maximum D2O/H2O ratio of approximately 1.4 results in this case.

We have shown, using water, that strong field ionization populates an excited electronic state without vibrational/rotational pre-excitation. This is done with an isotropic ensemble by using isotope marked molecules. We modeled the multi-orbital system by employing a well tested SAE-TDSE simulation framework and show that both the 1b1 and 3a1 orbitals contribute. The simulation predicts a magnitude of the isotope effect somewhat smaller than is measured but exhibits the same characteristic intensity dependence. The single orbital simulation predicts a much smaller effect and a different intensity dependence than is measured. The simulation suggests that the measured independence of the ratio on intensity is due to angular ionization saturation, which increases the relative contribution of the lower valence orbital at high intensities.

These results provide a new tool for assessing contributions of different orbitals from strong field ionization. In addition, we show that ionization from inner valence orbitals can also excite nuclear dynamics. This extends the technique used in [9, 10, 25] to a larger range of molecules. Possible improvements of the theoretical model include, e.g., an incorporation of the relative phase between the two orbitals [2] and geometry-dependent ionization yields [20].

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