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Striking Isotopologue Dependent Photodissociation Dynamics of Water Molecules: The Signature of an Accidental Resonance

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Abstract

Investigations of the photofragmentation patterns of both light and heavy water at the state-to-state level are a pre-requisite for any thorough understanding of chemical processing and isotope heterogeneity in the interstellar medium (ISM). Here we reveal dynamical features of the dissociation of water molecules following excitation to the \( \tilde{C}(010) \) state using a tunable vacuum ultraviolet source in combination with the high resolution H(D)-atom Rydberg tagging time-of-flight technique. The action spectra for forming H(D) atoms and the OH(OD) product state distributions resulting from excitation to the \( \tilde{C}(010) \) states of H\(_2\)O and D\(_2\)O both show striking differences, which are attributable to the effects of an isotopologue-specific accidental resonance. Such accidental resonance induced state mixing may contribute to the D/H isotope heterogeneity in the Solar System. The present study provides an excellent example of competitive state-to-state non-adiabatic decay pathways involving at least five electronic states.
Introduction

Water, ubiquitous in the universe, absorbs light at all wavelengths ($\lambda$) in the vacuum ultraviolet (VUV) region below 190 nm, resulting in fragmentation leading to a hydrogen (H) atom and a hydroxyl (OH) radical and/or, at shorter wavelengths, the three constituent atoms. The single bond fission process is an important source of OH radicals in the interstellar medium (ISM). An in-depth understanding of the OH product quantum state population distributions from the photodissociation of water is thus an essential prerequisite for interstellar related chemistry modelling. Many distinctive features of the OH product state distributions arising from water photodissociation have already been identified, and explained as exemplars of a rich gamut of non-adiabatic dynamics, e.g., passage through conical intersections (CIs), Coriolis couplings, vibronic couplings, etc. Here we demonstrate another novel feature of the OH product state distribution arising in the VUV photodissociation of water, which is isotope-specific and can be attributed to an accidental vibronic resonance. Accidental resonances are often observed in molecular spectra – as perturbations in line positions, intensities or widths. However, the effects of accidental resonances in the parent absorption upon the subsequent fragmentation dynamics have rarely been recognized hitherto.

The photodissociation of H$_2$O has been the subject of extensive experimental and theoretical investigations over the past decades, so much so that H$_2$O (and its various isotopomers) is now viewed as a benchmark system for illustrating and understanding
non-adiabatic coupling pathways between potential energy surfaces (PESs). Excitation of H$_2$O at $\lambda \sim 160$ nm populates the lowest excited electronic ($\tilde{A}^1B_1$) state; the subsequent direct dissociation from this state yields an H atom plus a ground state OH($\chi^2\Pi$) radical with little internal excitation.$^{10-16}$ Absorption to the second excited singlet ($\tilde{B}^1A_1$) state maximizes at $\lambda \sim 128$ nm. This state displays a minor direct dissociation channel to H + OH($A^2\Sigma^+$) products, and major non-adiabatic pathways via two CIs with the ground ($\tilde{X}^1A_1$) state PES at linear HOH and HHO configurations that favor formation of highly rotationally excited OH($X$, $v$=0) products.$^{17-29}$ The next two excited states identified in absorption have predominant Rydberg character, and predissociate by non-adiabatic coupling to those lower (predominantly valence) states. The $\tilde{C}^1B_1$ state shows resolvable rotational structure at $\lambda \sim 124$ nm,$^{30-33}$ and previous studies have identified dramatic variations in the OH product state distributions and angular distributions that arise following excitation to different rotational levels of this state.$^{34}$ Dissociation from $\tilde{C}$ state rotational levels with $K_a' = 0$ (where $K_a$ describes the projection of the parent rotational angular momentum ($J$) onto the $a$ inertial axis) occurs exclusively by electronic coupling to the $\tilde{A}$ state and yields highly vibrationally excited OH($X$) products with vibrational quantum number $v'' \leq 13$ (henceforth termed “high $v$” products). A rival decay pathway opens for H$_2$O ($\tilde{C}$) molecules in levels with $K_a' > 0$. This relies on Coriolis-type (i.e. rotationally induced) coupling to the $\tilde{B}$ state PES, and results in both ground ($X$) and electronically excited ($A$) state OH fragments with very high rotational but minimal vibrational excitation (henceforth termed “high $N$” products).
Replacing H by deuterium (D) often leads to significant isotopic effects in processes that are dynamically controlled, particularly in cases where non-adiabatic dynamics are involved. Indeed, investigations of the photodissociation of heavy water (D₂O) following excitation to the same excited states have revealed qualitative variations from the H₂O data but, in all cases studied to date, the same dissociation mechanisms appear to prevail for both H₂O and D₂O. However, a recent ultrafast time-resolved photoelectron spectroscopy study revealed quantitatively different predissociation behaviors for the \( \tilde{C}(010) \) states (i.e. the states carrying one quantum of bending vibrational excitation, \( \nu_2' \)) of H₂O and D₂O. The present article unravels the state-to-state dynamics associated with this unusual isotopic effect, using tunable VUV pump and probe laser sources and the high resolution H/D atom Rydberg tagging technique. The distinctive OH and OD quantum state distributions determined in this work provide further exquisitely detailed illustrations of the sensitivity of the various possible non-adiabatic decay pathways to details of the excited state PESs and the inter-state coupling matrix elements.

The first experiment involved scanning the VUV photolysis wavelength while recording the total H (D) atom yield (see Experimental Methods in supporting information), thereby obtaining action spectra for forming H (D) atoms from VUV photodissociation of H₂O (D₂O). Figure 1A shows the action spectrum from photolysis of H₂O in the wavelength range 121.7-122.4 nm, which spans transitions to the \( \tilde{C}(010) \) state and to the origin (000) level of the \( \tilde{D}^1A_1 \) state. The broad peak centered at \( \lambda \sim 122.1 \) nm and extending across this entire spectral window is
attributable to the $\tilde{D} \leftarrow \tilde{X}$ origin band; the width of this feature reflects the short predissociation lifetime of the $\tilde{D}$ (000) state. Note that this spectrum is distorted by an obvious dip in the VUV tuning curve at $\lambda \sim 122.09$ nm, which is also apparent in the action spectra for forming D atoms from both D$_2$O and CD$_4$ photolysis shown in Figure 1C. Analogy with the rotational fine structures of the $\tilde{C} \leftarrow \tilde{X}$ origin band allows the partially resolved peaks at $\lambda \sim 121.85$ nm to be assigned to the three strongest rotational lines in the jet-cooled $\tilde{C}$ (010) $\leftarrow \tilde{X}$ (000) transition – as shown in Figure 1B. This analysis provides an experimental estimate of the bending vibrational wavenumber for the $\tilde{C}$ state of H$_2$O, $v_2' = 1402\pm 3$ cm$^{-1}$, which is in good accord with the values determined in previous multiphoton spectroscopy studies and theoretical predictions.

In contrast, the action spectrum from the photodissociation of D$_2$O shows well resolved rotational structure associated with the $\tilde{C}$ (010) $\leftarrow \tilde{X}$ (000) transition (Figure 1C). The striking similarity between this spectrum and that of the $\tilde{C} \leftarrow \tilde{X}$ origin band (Figure 1D) allows straightforward assignment of the rotational fine structure in the $\tilde{C}$ (010) $\leftarrow \tilde{X}$ (000) spectrum and yields a value of $v_2' = 1040\pm 2$ cm$^{-1}$ for the $\tilde{C}$ state of D$_2$O. The widths of individual lines in the $\tilde{C}$ (010) $\leftarrow \tilde{X}$ (000) action spectra (Figures 1B and 1D) are $\sim 15$ cm$^{-1}$ and 1.2-1.7 cm$^{-1}$ for H$_2$O and D$_2$O, respectively. These widths correspond to predissociation lifetimes of $\sim 330$ fs for H$_2$O and 3.0-4.2 ps for D$_2$O, in good accord with the time resolved measurements of He et al. and with the conclusions from earlier analyses of the respective resonance enhanced multiphoton ionization spectra of H$_2$O and D$_2$O, wherein the striking
isotope effect was attributed to the effects of an accidental resonance between the $\tilde{C}(010)$ and $\tilde{D}(000)$ vibronic levels of H$_2$O. The $\tilde{C}(010)$ and $\tilde{D}(000)$ states of D$_2$O, in contrast, are $\sim 270$ cm$^{-1}$ out of resonance$^{30}$ and the mixing between the two states is thus negligible.

We have measured TOF spectra of the H (D) atom products formed with the VUV photolysis laser tuned to each of the more intense rotational lines in the $\tilde{C}(010) \leftarrow \tilde{X}(000)$ bands of H$_2$O and D$_2$O. Knowing the distance travelled from the photodissociation region to the detector and the fragment masses, such TOF spectra can be converted into spectra of the total kinetic energy release (TKER). The TKER distributions obtained with the polarization vector $\mathbf{e}_{\text{phot}}$ aligned, respectively, parallel and perpendicular to the detection axis can then be used to construct 3-dimensional (3-D) flux diagrams of the H + OH (D + OD) fragments. Figure 2 shows such 3-D flux diagrams for the products formed following excitation to the same $\tilde{C}(010) \leftarrow \tilde{X}(000)$, $1_{10} \leftarrow 0_{00}$ transitions of H$_2$O and D$_2$O. Each diagram shows two groups of features: an inner group, associated with formation of levels of electronically excited $A$ state of OH (OD) fragments, and an outer group associated with ground $X$ state OH (OD) products. The obvious differences in the 3-D distributions displayed in Figure 2 clearly illustrate that H$_2$O and D$_2$O molecules exhibit different predissociation dynamics, even when excited via the same transition to the same $J_{KaKc}$ level of the respective $\tilde{C}(010)$ state.

Figure 3 displays TKER distributions from D$_2$O photolysis at 122.282, 122.235 and 121.954 nm with $\mathbf{e}_{\text{phot}}$ aligned parallel and perpendicular to the detection axis. These wavelengths excite, respectively, the $1_{01} \leftarrow 1_{11}$ and $1_{10} \leftarrow 0_{00}$ transitions of the $\tilde{C}$
(010) ← $\tilde{X}$ (000) band, and near the center of the $\tilde{D}$ ← $\tilde{X}$ origin band. All of these TKER distributions show progressions of sharp peaks. Given that the total energy and linear momentum must be conserved in the photodissociation process, we can write

$$h\nu + E_{\text{int}}(\text{D}_2\text{O}) - D_0(\text{D–OD}) = E_T(\text{D + OD}) + E_{\text{int}}(\text{OD}),$$

(1)

where the internal energy ($E_{\text{int}}$) distribution of the OD fragment can be determined from the TKER ($E_T$) distribution, $h\nu$ is the photolysis photon energy, $h$ is Planck’s constant and $D_0(\text{D–OD})$ is the bond dissociation energy. All of the sharp peaks in Figure 3 can be assigned to population of specific rovibrational levels of the $A$ and $X$ states of OD.

The excited level populated at the first of these wavelengths has $K_a' = 0$ and, as Figure 3A shows, the TKER spectrum is invariant to the alignment of $\mathbf{e}_{\text{phot}}$ (i.e. the recoil velocity distribution is isotropic) and the fine structure indicates that the OD products are formed in their ground ($X$) state, in a very wide spread of vibrational states (up to $v'' = 18$), and with little rotational excitation. Both the energy disposal and the isotropy of the product recoil are very similar to those found when exciting the same rotational transition within the $\tilde{C}$ (000) ← $\tilde{X}$ (000) origin band, but totally different to that observed when exciting to the $\tilde{D}$ (000) state – which yields ‘high $N$’ OD products (Figure 3C). The TKER spectra obtained following excitation of the $1_{10}<\rightarrow 0_{00}$ transition (i.e. populating a $\tilde{C}$ (010) state level with $K_a' = 1$) are different again. As Fig. 3B shows, the spectrum recorded in the perpendicular direction is dominated by ‘high $v$’ OD($X$) products, whereas ‘high $N$’ OD ($A$) and OD($X$) products dominate the TKER spectrum recorded in the parallel direction. Again, these very
obvious differences are reminiscent of those reported previously when exciting the same rotational transition in the \( \tilde{C} \leftarrow \tilde{X} \) origin band of D\(_2\)O (and H\(_2\)O).\(^{34}\) These parallels suggest that the predissociation of \( \tilde{C} (010) \) and \( \tilde{C} (000) \) state D\(_2\)O molecules follow essentially the same mechanisms – namely a Coriolis-induced pathway to the \( \tilde{B} \) state PES and an electronic coupling to the \( \tilde{A} \) state PES as summarized in \textbf{scheme 1}. The former yields ‘\textit{high N}’ OD(X) and OD(A) products whereas the latter yields ‘\textit{high v}’ OD(X) products. The products of these two pathways show opposite recoil anisotropies and, for \( \tilde{C} (010) \) levels with \( K_{a} = 1 \), the respective signals suggest that the two processes occur with comparable efficiencies.

\[
\begin{align*}
\text{D}_2\text{O}(\tilde{C}(010)) & \xrightarrow{\text{Coriolis Coupling}} \text{D}_2\text{O}(\tilde{B}) & \rightarrow & \text{D}+\text{OD}(A, \text{ high } N) \\
& \xrightarrow{\text{Electronic Coupling}} \text{D}_2\text{O}(\tilde{A}) & \rightarrow & \text{D}+\text{OD}(X, \text{ high } N)
\end{align*}
\]

\textbf{Scheme 1}

The situation with H\(_2\)O is very different. \textbf{Figure 4} shows the TKER distributions obtained when exciting each of the three \( \tilde{C} (010) \leftarrow \tilde{X} (000) \) transitions identified in \textbf{Figure 1B}. TKER distributions were also measured with the photolysis wavelength tuned to the valley between the peaks at 121.851 and 121.814 nm in the action spectrum, and to 122.117 nm (where absorption is ascribed exclusively to the \( \tilde{D} \leftarrow \tilde{X} \) origin transition). These are shown in the supplementary information (\textit{Figures S1} and \textit{S2}). The TKER distributions obtained at all five wavelengths are essentially identical, showing progressions of sharp peaks that can be assigned to population of \textit{high N} levels of the \textit{A} and \textit{X} states of OH.
The TKER distributions and the recoil anisotropies appear insensitive to whether the initial excitation is to any of the three \( \tilde{C} (010) \) state rotational levels or to the \( \tilde{D} (000) \) state. Note that each of the probed \( \tilde{C} (010) \) levels of H\(_2\)O has \( K_a' \) (and thus \( \langle J_a^2 \rangle = 1 \)). Unfortunately, the linestrengths of the \( 0_{00} \leftarrow 1_{10} \) and \( 1_{01} \leftarrow 1_{11} \) transitions (which populate levels with \( \langle J_a^2 \rangle = 0 \)) are too weak to be identified under the stronger \( \tilde{D} \leftarrow \tilde{X} \) origin band, and any transitions to levels with \( K_a' = 2 \) are too heavily predissociated to be resolved.\(^{30}\) What is immediately obvious, however, is that – in contrast to D\(_2\)O – the fragmentation of H\(_2\)O \( \tilde{C} (010) \) molecules with \( K_a' = 1 \) yields high \( N \) products, i.e. the fragmentation dynamics are indicative of eventual nuclear motion on the \( \tilde{B} \) state PES, which are different from that when exciting the same rotational transition within the \( \tilde{C} (000) \leftarrow \tilde{X} (000) \) origin band (Figures S3). The TKER spectra in the perpendicular direction show no indication of any ‘high \( \nu \)’ OH(X) products such as would be expected if non-adiabatic coupling from these levels to the \( \tilde{A} \) state PES was a competitive process.

The striking isotope-dependent dynamics of the \( \tilde{C} (010), K_a' = 1 \) levels of H\(_2\)O and D\(_2\)O are directly attributable to the accidental resonance between the \( \tilde{C} (010) \) and \( \tilde{D} (000) \) states of H\(_2\)O. As noted above, the \( \tilde{D} \) and \( \tilde{B} \) states share a common \((^1A_1)\) symmetry, and the \( \tilde{D} (000) \) states of both isotopomers are efficiently predissociated by electronic mixing with the \( \tilde{B} \) state continuum. The \( \tilde{C} \) state, in contrast, has \(^1B_1\) symmetry. As shown in Scheme 1, \( \tilde{C} \) state molecules can decay by electronic (same symmetry) coupling to the \( \tilde{A} \) state continuum or, if prepared with appropriate nuclear motion (a-axis rotation), by coupling to the \( \tilde{B} \) state continuum. The two decay pathways have comparable probabilities for D\(_2\)O molecules with \( K_a' = 1 \) in
either the $\tilde{C}(000)$ or $\tilde{C}(010)$ states, and for $\text{H}_2\text{O}$, $\tilde{C}(000)$ molecules with $K_a'=1$. However, the accidental resonance with the $\tilde{D}(000)$ state greatly boosts the efficiency of Coriolis-coupling from the $\tilde{C}(010)$ state, $K_a'=1$ levels of $\text{H}_2\text{O}$ to the $\tilde{B}$ state continuum, as illustrated in Scheme 2, to the extent that this pathway to ‘high $N$’ products overwhelms the potential rival electronic coupling route to ‘high $v$’ $\text{OH}(X)$ products.

![Scheme 2](image)

**Scheme 2**

A further feature of the $\text{OH}(X, v=0)$ rotational distributions displayed in Figure 4 is the clear intensity alternation in the populations of successive $N$ levels. These alternations are reminiscent of those observed in the $\text{OH}(X)$ fragments arising in the 121.6 nm photolysis of $\text{H}_2\text{O}_5$ and, as in that case, can be attributed to dynamical interferences between dissociation pathways via the CIs between the $\tilde{B}$ and $\tilde{X}$ state PESs at linear $\text{H}–\text{O}–\text{H}$ and $\text{H}–\text{H}–\text{O}$ geometries.

The dissociation mechanisms shown in schemes 1 and 2 highlight a clear yet unprecedented isotope dependence in the photodissociation of $\tilde{C}(010)$ state water molecules. Previous studies have revealed qualitatively different dissociation behaviors for $\text{H}_2\text{O}$ and $\text{D}_2\text{O}$ molecules following excitation to the same electronic states, most of which effects could be attributed to mass changes.\textsuperscript{35-37} Here, we reveal a much more dramatic isotope effect, caused by accidental resonance, which leads to different dissociation rates, different dissociation mechanisms, and different product
energy disposals following excitation to a common excited state of H$_2$O and D$_2$O. Given the high density of excited vibronic levels of H$_2$O and D$_2$O accessible at shorter excitation wavelengths$^{40}$ it would be surprising if the accidental resonance – and the striking isotope dependent fragmentation dynamics – identified in the present study were to be the only example available following VUV photoexcitation of water molecules.

The photodissociation of water is invoked as an important photochemical process in interstellar molecular clouds.$^{41}$ Thus the isotope-specific dissociation process identified in the present study may have implications in modelling chemistry prevailing in the interstellar medium. For example, self-shielding has been proposed to account for the observed isotope heterogeneity within the Solar System,$^{42,43}$ but recent investigations of the VUV photodissociation of CO support the view that perturbation-dominated state mixing dynamics will also introduce isotopologue-dependent dissociation probabilities.$^{44,45}$ The strikingly different fragmentation dynamics of H$_2$O and D$_2$O observed in the present work represent an extreme example of such accidental resonance induced state mixing and may be expected to contribute to the D/H isotope heterogeneity in the Solar System.

In summary, detailed dynamical features of the photodissociation of $\tilde{C}$ (010) state H$_2$O and D$_2$O molecules have been determined using a tunable VUV source in combination with the high resolution H/D atom Rydberg tagging technique. The predissociation mechanisms of H$_2$O and D$_2$O following excitation to the $\tilde{C}$ (010) state show a striking isotopologue dependence. The OH product state distributions are
determined by the accidental resonance mediated non-adiabatic coupling pathway, but this pathway has negligible influence on the OD product state distributions. The present study provides an excellent example of competitive state-to-state non-adiabatic couplings and decay pathways involving at least five electronic PESs.

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Figures and Captions

**Figure 1** One-photon action spectra for forming (A) H atoms from H$_2$O and (C) D atoms from D$_2$O following excitation at wavelengths around 122 nm. The resolved structures at $\lambda \sim 121.8$ nm for H$_2$O and $\lambda \sim 122.2$ nm for D$_2$O are attributable to rotational lines of the respective $\tilde{C}(010)-\tilde{X}(000)$ transitions. The broad features centered at $\lambda \sim 122.1$ nm for H$_2$O and $\sim 121.9$ nm for D$_2$O are the respective $\tilde{D}-\tilde{X}$ origin bands, the profiles of which are distorted by a dip in the VUV tuning curve at $\lambda \sim 122.09$ nm (which is also evident in the action spectrum for forming D atoms from one photon photolysis of CD$_4$ shown in panel (C)). The extended rotational structures and line assignments of the $\tilde{C}(010)-\tilde{X}(000)$ transitions of H$_2$O and D$_2$O are shown in panels (B) and (D), along with comparisons with the respective $\tilde{C}-\tilde{X}$ origin bands. The red arrows in panels (A) and (B) indicate the excitation wavelengths used when measuring the TKER spectra reported in Figures 4, S1 and S2.
Figure 2 3D contour plots of the (A) H + OH products from the photodissociation of H$_2$O and (B) the D + OD products from the photodissociation of D$_2$O via the $1_{10} \leftarrow 0_{00}$ lines of the respective $\tilde{C} (010) \rightarrow \tilde{X} (000)$ transitions. The double headed arrow in (A) shows the alignment of $\varepsilon_{\text{phot}}$. The outer rings in both plots are associated with formation of rovibrational states of the ground ($\tilde{X}$) state OH radical products, while the inner structures are primarily due to OH/OD(A) products.
**Figure 3** TKER spectra of the D + OD products from the photodissociation of D$_2$O at (A) 122.282, (B) 122.235 and (C) 121.954 nm, resonant with the 1$_{01}$ ← 1$_{11}$ and 1$_{10}$ ← 0$_{00}$ lines of the $\tilde{C}$ (010) − $\tilde{X}$ (000) transition and the center of the $\tilde{D}$ − $\tilde{X}$ origin band, respectively, with $\varepsilon_{\text{phot}}$ aligned parallel (black) and perpendicular (red) to the detection axis.
Figure 4 TKER spectra of the H + OH products from photodissociation of H₂O at (A) 121.881, (B) 121.851 and (C) 121.814 nm, resonant with the $1_1^1 \leftarrow 1_0^1$, $1_1^0 \leftarrow 0_0^0$ and $2_1^1 \leftarrow 1_0^1$ rotational lines of the $\tilde{C}(010) - \tilde{X}(000)$ transition, respectively, with $\varepsilon_{\text{phot}}$ aligned parallel (black) and perpendicular (red) to the detection axis.
References


