Disentangling the role of laser coupling in directional breaking of molecules

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I. INTRODUCTION

As a primary step for steering chemical dynamics, directional bond breaking is one of the most fundamental and interesting phenomena in molecular dissociative ionization. It can be coherently controlled by using carrier-envelope phase (CEP) stabilized few-cycle [1–5] or two-color [6–10] ultrashort laser pulses. Previous studies have demonstrated that the directional bond breaking in dissociative ionization of a diatomic molecule with symmetric orbital profile along the molecular axis is governed by the pathway interference of the dissociating nuclear wave packets [11–15]. The most intensively investigated example is the dissociative single ionization of molecular hydrogen and its isotopes [1–6,8,11–19], which have symmetric electron distribution along the molecular axis.

For many heteronuclear diatomic molecules, the electron distributions around two nuclei are asymmetric; thus both the ionization and dissociation may be directional and contribute to the ultimate asymmetric fragment emission. For instance, CO is preferred to be ionized by the laser field pointing from C to O along the molecular axis [20–25]. The created molecular ion with biased orientation can subsequently be dissociated by the same asymmetric ultrashort laser pulse and may involve laser-induced coupling of electronic states, which in itself might result in an asymmetric fragment emission [26–31]. We note that the controllable directional strong-field dissociative ionization was recently demonstrated in multiply charged states [9,29,30], polyatomic [10,32] and hydrocarbon molecules [33–37], and further in two-dimensional space [38,39]. One essential aspect to thoroughly understand the directional dissociative ionization of complex molecules is to clearly distinguish individual contributions of the ionization and dissociation steps. It will also allow us to testify for the directional dissociation of a multielectron system the role of laser-induced coupling of various electronic states, which rules the directional dissociation of the one-electron molecule H2+ [17,18]. However, the coexistence of asymmetric ionization and asymmetric dissociation within a single femtosecond laser pulse blurs the contributions responsible for the ultimate directional emission of ionic fragments.

In this paper, using CO as a prototype, we conceived a strategy to disentangle the contributions from either step. As illustrated in Fig. 1, the single ionization created CO+ cation by an elliptically polarized pump pulse, i.e., CO + nhνpump → CO++ + e, is dissociated by a time-delayed linearly polarized two-color pulse into C+ and a neutral O atom, labeled as (C+, O). The distinguished momentum distributions of electrons governed by the light polarizations allow us to identify the ionization and the dissociation induced by the pump and probe pulses. By coincidentally detecting the released electron and C+, we can attribute the observed directional ejection of C+ upon the bond breaking of CO+ to the phase-dependent laser-induced coupling of various electronic states. The mechanism is numerically verified by solving the time-dependent Schrödinger equation (TDSE).

II. DISENTANGLING IONIZATION AND DISSOCIATION

The experiments were performed in an ultrahigh vacuum cold-target recoil ion momentum spectroscopy (COLTRIMS) setup [40,41], as illustrated in Fig. 1, driven by an elliptically polarized pump (ellipticity ～0.8) and linearly polarized...
two-color probe femtosecond laser pulses. A femtosecond laser pulse from a Ti:sapphire multipass amplifier (25 fs, 795 nm, 10 kHz) was split into pump and probe pulses. The pump pulse was adjusted to be elliptically polarized in the y-z plane with the major and minor axes along y and z axis, respectively. The two-color probe pulse was generated in a collinear scheme. Briefly, the z-direction polarized fundamental wave (FW) was frequency doubled in a 200-μm-thick β-barium borate (BBO) crystal to produce a second harmonic (SH) through the type-I phase matching. The polarization of the FW was rotated to be parallel to that of the SH along the y axis by using a dual-color wave plate. The time lag between the FW and SH waves of the two-color pulse was compensated by a birefringent α-BBO crystal. A pair of fused silica wedges were used to continuously vary the relative phase φ1 between the FW and SH waves of the two-color pulse. The pump and probe pulses were collinearly combined using a beam splitter, which were afterwards sent into the vacuum chamber and focused onto the molecular beam using a concave silver mirror with a focusing length of f = 75 mm. The molecular beam was generated by supersonically coexpanding a mixture of 10% CO and 90% He through a 30-μm nozzle with a driving pressure of 1.5 bar. The intensities of the pump pulse, the FW, and SH fields of the two-color probe pulse. On the other hand, electrons in the eprobe region correspond to the ionization by the pump pulse, while the created CO⁺ could be dissociated into the (C⁺, O) pair either by the pump pulse itself or later on by the time-delayed probe pulse. We are mostly interested in the latter case, i.e., the ionization by the pump pulse and dissociation by the probe pulse. This scenario excludes the influence of molecular orientation-dependent field ionization and thus reveals the role of laser-induced coupling of various electronic states of the molecular ion on the directional dissociation.

To extract the real pump-ionization probe-dissociation (C⁺, O) breakup events, we further testify the kinetic energy release (KER) and angular distribution of the emitted C⁺ fragments measured in coincidence with electrons in the eprobe region. As displayed in Fig. 3(a), as compared to those by only the pump pulse (black dotted curve, legend “pump only”) or only the probe pulse (gray dotted curve, legend “probe only”), the yield of the C⁺ is significantly enhanced for $E_{C^+} > 0.57$ eV when the probe pulse is sequentially applied following the pump pulse (black solid curve, legend “e_{pump} + e_{probe}” correlated to all photoelectrons). The significant enhancement and the similar positions of the KER peaks indicate that (C⁺, O) breakup is governed by the two-step process: the pump pulse singly ionizes CO, and the produced CO⁺ is later on dissociated by the probe pulse. Such a two-step process is further confirmed by gating on the momentum distribution of the electron measured in coincidence with the ion fragments.
As shown in Fig. 3(a), for electrons in the $e_{\text{probe}}$ region (blue dashed curve, legend “$e_{\text{probe}}$”), the C$^+$ shows a similar KER distribution to that produced by the probe pulse only. The enhancement at $E_{C^+} > 0.57$ eV is observed only for the (C$^+$, O) breakup when the electron is freed by the pump pulse and obtains momentum in the $e_{\text{pump}}$ region (red solid curve, legend “$e_{\text{pump}}$”). The successive dissociation of the pump-created CO$^+$ by the time-delayed probe pulse leads to the enhanced KER distribution of C$^+$.

More interestingly, as displayed in Figs. 3(b) and 3(c), the emitted C$^+$ correlated to different electrons shows different KER-dependent angular distributions. Associated with electrons in the $e_{\text{probe}}$ region, the C$^+$ mainly emits along the polarization direction of the linearly polarized two-color probe pulse, i.e., along $\phi_{C^+} = 0^\circ$ or $\pm 180^\circ$. However, associated with electrons in the $e_{\text{pump}}$ region, the angular distribution of the emitted C$^+$ strongly depends on $E_{C^+}$. As shown in Fig. 3(c), C$^+$ emits mainly along $\phi_{C^+} = 0^\circ$ or $\pm 180^\circ$ for $E_{C^+} < 0.57$ eV, which is similar to the data in Fig. 3(b), and also similar to that by the pump pulse only (data not shown here). Thus we conclude that this part is produced by the pump pulse itself. Furthermore, a noticeable dissociation of orthogonally and parallel oriented molecules with the energy peaked at $E_{C^+} = 0.6$ and 0.7 eV, respectively, is also observed. Figure 3(d) shows the corresponding angular distributions of the emitted C$^+$ for the high ($E_{C^+}\text{-H}$) 0.64–0.8 eV) and low ($E_{C^+}\text{-L}$) 0.57–0.64 eV) KER regions, respectively. The $E_{C^+}$-dependent preferred dissociation of the CO$^+$ cation with the molecular axis parallel or orthogonal to the laser polarization indicates different dissociation dynamics.

We will now address the essential question of whether or not the laser-induced coupling of various electronic states plays a crucial role in the directional dissociation of the multielectron system. We trace the directional emission of C$^+$ as a function of $\phi_L$ of the two-color probe pulse.

As shown in Figs. 4(a)–4(c), the $\phi_L$-dependent yield of C$^+$ as a function of $\phi_{C^+}$ at different KERs measured in coincidence with electrons in the $e_{\text{probe}}$ region, (b) $e_{\text{pump}}$ region at $E_{C^+}\text{-H}$, and (c) $e_{\text{pump}}$ region at $E_{C^+}\text{-L}$. The corresponding asymmetries of the directional emission of C$^+$ versus $\phi_L$. The corresponding asymmetries of the directional emission of C$^+$ versus $\phi_L$ (blue square) and the numerically simulated asymmetries when the molecule orients parallel (orange dashed curve) to the field polarization. The corresponding asymmetries of the directional emission of C$^+$ versus $\phi_L$ and the numerically simulated asymmetries when the molecule orients orthogonal (olive dashed curve) to the field polarization.
We now analyze the directional emission of C\(^+\) in the pump-ionization probe-dissociation process. As shown in Figs. 4(b) and 4(e) (blue squares), directional emission of C\(^+\) for \(E_{C^+\text{II}}\) is clearly observed along the polarization direction of the two-color pulse. Note that there is no asymmetry of the molecular orientation in the initial ionization by the multicycle elliptically polarized pump pulse. The observed asymmetry in Fig. 4(e) (blue squares) should originate from the dissociation by the time-delayed two-color probe pulse. Since the electron has already been released by the elliptically polarized pump pulse in the ionization step, recollisional excitation of various electronic states dominate the observed asymmetry depending on the relative phase of the two-color pulse. This is also consistent with previous observations [28] that the emitted C\(^+\) at KER lower than 1 eV is mainly due to the laser-induced coupling of the bound and repulsive electronic states. As compared to the asymmetry displayed in Fig. 4(d) where both the ionization and dissociation steps contribute the asymmetry for the C\(^+\) emission, the \(\phi_\text{L}\)-dependent asymmetry (blue squares) in Fig. 4(e) is much smaller and slightly phase shifted. In addition to molecules oriented along the light polarization, asymmetric dissociation of CO\(^+\) is also observed for orthogonally oriented molecules around \(\phi_\text{C^+} = \pm 90^\circ\), i.e., C\(^+\) is preferentially emitted to \(\phi_\text{C^+} = 90^\circ - \delta_\phi\) (or \(-90^\circ + \delta_\phi\)) for \(\phi_\text{L} = 0\) as compared to \(\phi_\text{C^+} = 90^\circ + \delta_\phi\) (or \(-90^\circ - \delta_\phi\)) for \(\phi_\text{L} = \pi\), where \(\delta_\phi\) is a small angle. As shown in Figs. 4(e) and 4(f), the asymmetries for the parallel and orthogonally orientated molecules have different phases. We emphasize that the orthogonal orientation here only means that CO\(^+\) has an orientation angle very close to but outside of \(\pm 90^\circ\). Strictly speaking, C\(^+\) symmetrically emits with the exact angle \(\pm 90^\circ\).

### III. NUMERICAL SIMULATION

To explore how the directional emission of C\(^+\) is built in the dissociation of CO\(^+\), we numerically simulate the modeled TDSE (atomic units are used throughout unless indicated otherwise):

\[
\begin{align*}
\frac{\partial}{\partial t} & \begin{pmatrix} 
\chi_1(R,t) \\
\chi_2(R,t) \\
\chi_3(R,t) \\
\vdots \\
\chi_{6}(R,t)
\end{pmatrix} = \\
& \begin{pmatrix} 
T_{\text{mac}} + V_1(R) & \mu_{12} \cdot \vec{E}(t) & \mu_{13} \cdot \vec{E}(t) & \cdots \\
\mu_{12} \cdot \vec{E}(t) & T_{\text{mac}} + V_2(R) & \mu_{23} \cdot \vec{E}(t) & \cdots \\
\mu_{13} \cdot \vec{E}(t) & \mu_{23} \cdot \vec{E}(t) & T_{\text{mac}} + V_3(R) & \cdots \\
\vdots & \vdots & \vdots & \ddots
\end{pmatrix}
\end{align*}
\]

where six electronic states are included to describe the dissociative dynamics, \(\chi_{1-3}\) and \(\chi_{4-6}\) are the associated nuclear wave packets for the three \(\Sigma\) states, and three \(\Pi\) states shown in Fig. 5 from bottom to up, respectively. \(T_{\text{mac}}\) is the nuclear kinetic energy operator, the potential energy curves \(V_i(R)\) and the \(i\)-dependent dipole coupling matrix elements \(\vec{\mu}_{ij}\) (1 \(\leq i \leq 6, 1 \leq j \leq 6, i \neq j\)) are calculated by MOLPRO [49] with the multireference configuration interaction method based on the aug-cc-pVQZ basis set [50]. The reference configurations are all electronic configurations generated from \([1\sigma^2,2\sigma^2,3\sigma^0,2,4\pi^0,2,1\pi^0,4,5\sigma^0,2,2\pi^0,2,6\sigma^0,2,7\pi^0,0]\), and the calculations are performed under \(C_{2v}\) symmetry [50].

The two-color probe pulse \(E(t)\) is written as

\[
E(t) = E_1\cos(\omega_1 t)\exp\left[-4\ln 2 \left(\frac{t - \left(\frac{\tau_2}{2}\right)}{\tau_1^2}\right)^2\right] \\
\quad + E_2\cos(\omega_2(t - \Delta t))\exp\left[-4\ln 2 \left(\frac{t - \left(\frac{\Delta t - \frac{\tau_2}{2}}{\tau_2^2}\right)}{\tau_2^2}\right)^2\right],
\]

where the laser parameters are the same with that used in the experiment, and \(\Delta t\) is the time delay. We used the Crank-Nicolson method to propagate the wave packets. The time and spatial steps are \(\Delta t = 0.1\) a.u. and \(\Delta R = 0.02\) a.u., and the simulation convergence has been tested by using denser time-spatial grids. The simulation box is big enough to hold all wave packets in all simulations.

To reproduce the main observations of the experiment, we focused on two cases with the molecular axis parallel or orthogonal to the polarization of the two-color pulse. For C\(^+\) emitted around the angle \(\phi_{C^+} = 0^\circ\) or \(180^\circ\), the molecular axis of CO\(^+\) is parallel to the polarization axis of the probe pulse.

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**FIG. 5.** The potential energy surfaces of involved electronic states of CO\(^+\) calculated by MOLPRO, where the Franck-Condon ionization region of the ground state CO is indicated by the grayed bar. The red (FW photon) and blue (SH photon) arrows indicate the transition pathways for the dissociation of CO\(^+\) with molecular axis parallel and orthogonal (labeled as \(S_{\text{par}}\) and \(S_{\text{ort}}\)) to the polarization of the two-color field.
when dissociation starts. We started from the vibrational states \(v = 3\) or \(4\) of \(A^2\Pi\) produced by two-\(\omega_{\text{pump}}\)-photon resonant excitation from the \(X^2\Sigma^+\) state separately, which are obtained by the imaginary time propagation, and simulated Eq. (1) by only keeping all \(2\Pi\) states, and added the dissociative \(C^+\) energy spectra incoherently after the interaction, i.e., the nuclear wave packet propagating on the \(A^2\Pi\) curve undergoes a one-\(\omega_{\text{SH}}\)-photon and a two-\(\omega_{\text{FW}}\)-photon transition to the \(D^2\Pi\) curve. Note that the simulation from the coherent sum of the vibrational states \(v = 3\) or \(4\) gives almost the same result with the incoherent simulation, since the dissociation fragments from \(v = 3\) or \(4\) end with clearly different energies. The initial populations of these two vibrational states are assumed equally, and the final dissociation probabilities from these two vibrational states are adjusted by the relative weights obtained from experimental measurements. After the interaction with the probe pulse, we used the windows operator [51] to extract the energy spectra,

\[
P(E) = \sum_k \left| \overline{\chi_k(E)} \right|^2,
\]

where \(\overline{\chi_k(E)} = \frac{2}{\pi \delta_E} \langle \chi_k | R^+ R | \chi_k \rangle \) and \(R = \frac{\delta_F^2}{(E - H_0)^2 + i \delta_F} \) with \(E\) the total energy, \(H_0\) the field free Hamiltonian, and \(\delta_F\) associated with the energy resolution. The simulated \(C^+\) energy spectra are shown in Fig. 6 by the gray dashed curve, which qualitatively agrees well with the experimental observations (olive dotted curve). The calculated asymmetry parameter of \(C^+\) is shown in Fig. 4(e) (orange dashed curve). The low-energy peaks in the range \(0.2\)–\(0.5\) eV shown in Fig. 3(a) are contributed by other vibrational states of \(A^2\Pi\), which are vertically populated when the HOMO-1 electron in CO is directly removed by the pump pulse.

For \(C^+\) emitted around the angle \(\phi_{C^+} = 90^\circ\), we aligned CO+ with an angle of \(80^\circ\) in calculations. We tested that the conclusion does not change substantially if the aligned angle varies between \(75^\circ\) and \(85^\circ\). In this case, all transitions between these six electronic states are allowed. We chose \(v = 0\) of the \(X^2\Sigma^+\) state of \(C^+\) as the initial state, which is also almost the ground nuclear state of CO. The component of the laser field perpendicular to the molecular axis triggers the transition between \(2\Sigma^+\) and \(2\Pi\), and the parallel component of the laser field to the molecular axis induces the transition between all \(2\Pi\) states, i.e., the nuclear wave packet propagating on the \(X^2\Sigma^+\) curve undergoes a one-\(\omega_{\text{SH}}\)-photon transition to \(A^2\Pi\), then undergoes a one-\(\omega_{\text{SH}}\)-photon and two-\(\omega_{\text{FW}}\)-photon transition to the \(D^2\Pi\) curve. We followed the same steps as calculating \(C^+\) emitted along the angle \(\phi_{C^+} = 0^\circ\) or \(180^\circ\), and show the calculated energy spectrum and asymmetry in Fig. 6 (blue solid curve) and Fig. 4(f) (olive dashed curve), respectively. To fully reproduce the angular distribution as shown in Figs. 3(b) and 3(c), a numerical model which describes the nuclear wave packet in two-dimensional space is to be developed. Nevertheless, the good qualitative agreement of the energy spectra and asymmetries between experimental measurements and theoretical calculations confirms that the laser coupling between different electronic states dominates the asymmetric \(C^+\) emission.

**IV. CONCLUSION**

In summary, taking CO as a prototype, a straightforward and robust strategy to disentangling the coexisting ionization and dissociation contributions to the directional dissociative ionization of a multielectron molecule is demonstrated. Our quantum simulations confirm that the laser coupling among different electronic states in \(CO^+\) plays an important role for the directional \(C^+\) fragment emission. Depending on the KER of the nuclear fragments, directional dissociation of \(CO^+\) oriented orthogonally to the light polarization is observed as a function of the relative phase of the two-color ultrashort laser pulse. The strong-field dissociative ionization of molecules is complex where many effects may be involved and entangled with each other. Our experimental technique disentangles the dissociative ionization into the ionization and dissociation steps, providing a powerful tool to investigate even more complex molecular reactions in strong laser fields.

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