Laser-induced dissociative recombination of carbon dioxide

Hongtao Hu, Seyedreza Larimian, Sonia Erattupuzha, Jin Wen, Markus Kitzler-Zeiler, and Xinhua Xie

1 Photonics Institute, Technische Universität Wien, A-1040 Vienna, Austria
2 Institute of Organic Chemistry and Biochemistry, AS CR, 166 10 Praha 6, Czech Republic
3 Institute of Theoretical Chemistry, University of Vienna, A-1090 Vienna, Austria
4 SwissFEL, Paul Scherrer Institute, 5232 Villigen PSI, Switzerland

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We experimentally investigate laser-induced dissociative recombination of CO₂ in linearly polarized strong laser fields with coincidence measurements. Our results show laser-induced dissociative processes with electron recombination after laser-induced double ionization. After double ionization, one electron can recombine to one of the two ionic fragments during laser-induced molecular dissociation. Our measurements reveal that the recombination probability of the second ionized electron is three times as high as that of the first ionized electron.

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Dissociative recombination refers to the process where a positively charged molecular ion captures a free electron, upon which a (highly) excited molecular complex is formed that subsequently dissociates into fragments [1,2]. It is one of the most important processes in plasmas, for instance in planetary atmospheres, fusion plasmas, and laser physics [3,4]. Most of experimental studies on dissociative recombination were performed at large facilities, such as heavy-ion storage rings in which the molecular ions are produced by electron-impact ionization [1,2].

In the past decades, femtosecond lasers became a versatile tool to reveal nuclear and electron dynamics of atoms and molecules [5–8], due to their ultrashort pulse durations and high peak intensities. When exposed to a strong laser field, molecules can become ionized or excited, which may cause further molecular reactions, such as dissociation and isomerization. Laser-induced dissociative recombination (LIDR) is one of such reactions, which has attracted the interest of researchers for a decade [9]. In the literature on strong-field laser science this process is also referred to dissociative frustrated ionization [10], where electrons are released from a molecule by the strong laser field and afterwards an electron recombines with one of the ionic fragments during the dissociation of the molecule. To the best of our knowledge, the research on LIDR only focused on homonuclear diatomic molecules, such as H₂, D₂, and Ar₂ [11–14].

In this work, we investigated the LIDR of a triatomic molecule CO₂ in linearly polarized laser fields with a reaction microscope. In comparison with a diatomic, the complexity of a triatomic molecule is significantly higher due to the increasing of molecular motion modes and electron configurations [15]. Dissociative recombination of CO₂ is expected to play an important role in the interaction of CO₂ with photons and electrons [16]. It is valuable to gain knowledge on dissociative recombination of CO₂ for further understanding of laser-induced reaction in molecular complex. In this paper, we are trying to understand the dynamics of the LIDR of CO₂ through answering the following two questions: (1) how does the LIDR happen in CO₂ and (2) which of the two released electrons contributes more to the recombination in the LIDR process?

In the experiment, coincidence measurements of three-dimensional momenta of resulting particles are achieved using a reaction microscope [17]. A schematic of the coincidence measurement of the LIDR process is shown in Fig. 1(a). Charged particles, e.g., CO⁺ and electrons, are guided to the multihit position- and time-sensitive detectors by a homogeneous dc field of 10.5 V/cm and a uniform magnetic field of 12 G. Excited neutral particles, e.g., O*, with an initial momentum vector pointing towards the detector are registered with a detection acceptance angle of about 70° [9,11,18,19]. In the off-line data analysis, momentum conservation conditions between detected particles are applied to minimize background signals. In the measurements, linearly polarized 25-fs, 790-nm laser pulses (along z direction) are delivered from a Ti:sapphire laser amplifier with a repetition rate of 5 kHz. The laser beam is focused onto the CO₂ gas jet in the interaction chamber with a spherical mirror which has a focal length of 60 mm. CO₂ molecules are introduced to the interaction chamber from a supersonic gas jet system which consists of a gas nozzle followed by a skimmer. The laser peak intensity at the focus is about 5 × 10¹⁴ W/cm², so that CO₂ molecules can be singly or doubly ionized and subsequently dissociate into fragments. More details about the experimental setup can be found in our previous publications [20–24].

We now turn to the question how the LIDR happens in CO₂. The two possible LIDR processes can be written
as \( \text{CO}_2 \rightarrow \text{CO}^+ + \text{O}^+ + e \) (Ch4) or \( \text{CO}_2 \rightarrow \text{CO}^+ + \text{O}^+ + 2e \) (Ch5) (see Table I for channel definitions). With the reaction microscope, we achieve complete coincidence detection of the three particles (an ion, a neutral, and an electron) generated in these LIDR processes. Figure 1(b) shows the measured photoion photoion coincidence (PIPICO) distribution. The sharp parabolic PIPICO lines are signals of two-body dissociation channels as indicated in the figure, among which the LIDR channel Ch4 can be clearly identified. The weak LIDR channel Ch5 is not visible on the intensity scale of Fig. 1(b). However, after applying coincidence selection on the momentum sums of the two particles \( |p_1 + p_2| < 4 \text{ au} \), \( |p_1 + p_2| < 8 \text{ au} \), and \( |p_1 + p_2| < 3 \text{ au} \), the two LIDR processes can be selected out from the PIPICO distribution, as shown in Fig. 1(c). The simulated PIPICO lines in Fig. 1(c) represent for the corresponding processes, which confirms the observation of the two LIDR processes. The branching ratios of related processes are summarized in Table I. In the following, we will focus on Ch4 first.

As shown in the inset of Fig. 1(b), no LIDR signals are detected with circularly polarized light. This observation contradicts a previous experimental observation of hydrogen [25], in which Rydberg states are formed via resonant multiphoton excitation by irradiation of the circularly polarized laser fields. The observation of no LIDR signals with circular pulses confirms that Rydberg states are formed through frustrated double ionization in this work, which is a rescattering-like process [9,10,23,24,26]. This process will be strongly suppressed by circularly polarized laser fields [6,27–29].

To gain insight into the quantum states involved in our observations of laser-induced dissociation of CO\(_2\), we performed quantum chemical simulations to obtain the potential energy curves of the relevant states. Multiconfiguration complete-active-space self-consistent field theory (MC-CASSCF) should be employed for an accurate description of the electronic structure of CO\(_2\) in excited states. To simulate the dissociation processes of CO\(_2\), CO\(_2^+\), and CO\(_2^{++}\) along the C-O stretching coordinates in highly-excited states, a multistate complete active space perturbation theory of second order (MS-CASPT2) [30], which further considered dynamic correlations, was carried out using Molcas 8.2 [31]. In the active spaces of CASSCF calculations, 10, 9, and 8 electrons in 15 orbitals are included in CO\(_2\), CO\(_2^+\), and CO\(_2^{++}\) molecules respectively, when they were all kept in \( C_2V \) symmetry. In searching for potential energy curves along C-O bonds of CO\(_2^+\), the state-averaged CASSCF wave functions for 20 singlet excited states were used in MS-CASPT2 calculations with ANO-RCC-VTZP basis set, while O-C-O valence angles were constrained to 180\(^\circ\). The calculated potential energy curves (PECs) of CO\(_2\), CO\(_2^+\), and CO\(_2^{++}\) along the C-O bonds are presented in Fig. 2.

When CO\(_2\) interacts with a strong laser field, single or double ionization may occur by removing of one or two valence electrons. Since the energy gaps between different molecular orbitals (MOs) are rather small, direct removal of electrons from a low-lying MO is possible, resulting in electronically excited states [32,33]. As shown in Fig. 2, removal of one or two electrons from the HOMO of CO\(_2\) leads to stable ground states of the CO\(_2^+\) or CO\(_2^{++}\), observed as Ch1 or Ch2. On the other hand, dissociative electronically excited states of cation or dication can be reached through removal of at least one electron from low-lying MOs, yielding Ch3, Ch6, and Ch7. In the strong field interaction, high-lying Rydberg states can be populated through the so-called frustrated field ionization [9,10,23,26]. In case of molecules, such Rydberg states can be dissociative and close to the ionization threshold.

![Figure 1](image-url)

**FIG. 1.** (a) LIDR of CO\(_2\) induced by the intense laser field, and the coincidence detection of resulting particles with a reaction microscope. (b) Measured PIPICO distribution of CO\(_2\) in linearly polarized laser field. The blue curve at the bottom of (b) is the time-of-flight spectrum of the experiment. The insert of (b) shows the PIPICO distribution for the measurement with circular polarized laser pulses in the range marked by the pink rectangle. (c) Coincidence selection of LIDR processes in PIPICO distribution with corresponding simulated PIPICO lines.

**TABLE I.** The branching ratios of resulting channels from CO\(_2\) in strong laser fields. The peak intensity of the circular pulse was 2.5\( \times 10^{14} \) W/cm\(^2\).

| Channels | \((\text{CO}_2 \rightarrow )\) | Linear | Circular |
|----------|-------------------------------|--------|---------|
| Ch1      | CO\(_2^+\) + e               | 24.52% | 46.38%  |
| Ch2      | CO\(_2^+\) + 2e              | 14.10% | 1.60%   |
| Ch3      | CO\(_2^+\) + O\(^+\) + e     | 8.01%  | 5.62%   |
| Ch4      | CO\(_2^+\) + O\(^+\) + 2e    | 0.10%  | –       |
| Ch5      | CO\(_2^+\) + O\(^+\) + e     | 0.002% | –       |
| Ch6      | CO + O\(^+\) + e             | 16.38% | 12.65%  |
| Ch7      | CO\(^+\) + O + e             | 36.87% | 33.75%  |
to a higher charged state, schematically shown in Fig. 2, where a high-lying Rydberg state of CO$_2^+$ lies very close to one PEC of an electronically excited state of CO$_2^{2+}$. The kinetic energy release (KER) of the dissociation from these two states is similar because of negligible influence of the Rydberg electron to the dissociation process.

Since the measured momentum/energy distributions of detected particles contain information on the nuclear dynamics during strong field interaction, we first checked measured momentum and KER distributions of dissociation channels. The momentum distributions of the fragments generated through Ch3 and Ch4 are plotted in Figs. 3(a) and 3(b). Except for different acceptance angles, the momentum distributions of both channels are similar and peak along the laser polarization direction. Such anisotropic distributions indicate electron removal from low-lying $\sigma$-type MOs during the strong field ionization [20,21,32]. It implies that Ch3 and Ch4 originate from electronically excited states formed by removal an electron from lower-lying MOs, as in Fig. 2.

The KER distributions of Ch3 and Ch4 are plotted in Fig. 3(c). One clear observation is that the KER distribution of Ch4 (red squares) is almost the same as that of Ch3 (blue areas) with a peak at 5.8 eV, which agrees well with the simulated KER of 5.75 eV. Since the KER distribution is determined by the involved electronic states (see Fig. 2), this observation indicates that the dissociative electronic wave packets of Ch3 and Ch4 evolve on PECs with similar shapes. Furthermore, the same KER distributions of Ch3 and Ch4 not only shows that the recombination of the ion and the electron occurs during the molecular dissociation, but also provides a clear evidence that the electron is recaptured into high-lying Rydberg states such that the nuclear charge is not fully shielded by the trapped electron and leading to similar KERs of the two channels.

To summarize the answer to the first question, the LIDR happens as laser-induced dissociative double ionization through the removal of electrons from low-lying MOs followed by the electron recombination to one of the ionic fragments during the dissociation of the molecule.

Now we turn to the second question: which of the released electrons recombines more favorably with the parent ion? To answer it, we analyze momentum distributions of electrons and ions along the laser polarized direction. The final momentum of photoelectrons from the strong field interaction is determined by the vector potential of the laser field at the electron birth time [5,34]. Therefore, electron momentum distributions contain information about the ionization dynamics during strong field interaction. Due to momentum conservation, the electron momenta can be derived from measured ionic momenta. Figure 4 shows the measured ion momentum distributions of Ch1 to Ch5. For the dissociative processes the momentum sums of the two fragments are analyzed. First, we compare the momentum distributions of direct ionization processes (Ch1, Ch2, and Ch3). At the laser intensity of our measurement, single ionization happens at the leading edge of the laser pulses due to saturation, which leads to a rather narrow momentum distribution. On the other hand, in case of double ionization, we notice that the momentum distribution of the dissociative process (Ch3) is broader than that of the nondissociative process (Ch2). At the laser intensity of 5×10$^{14}$ W/cm$^2$, the dominant double ionization process happens sequentially, which means the two electrons are removed one after the other [35]. As shown in Fig. 3(a), the width of the sum momentum distribution for double ionization is broader than that for single ionization. It is due to the broader momentum width of the second electron which is removed after the first ionization step. As this lower-lying MO exhibits a
larger ionization potential than the HOMO [20], dissociative double ionization (Ch3) happens at an effectively higher laser intensity. This leads to a broader momentum distribution than that of nondissociative double ionization. Now, we compare the momentum distributions of Ch3 and Ch4. It is clear that the momentum distribution of Ch4 is much narrower than that of Ch3. For Ch4 only one electron is released, with the recombination of either the first or the second ionized electron.

In the following, we will quantify the contributions of the first and second electrons to the overall recombination yield from the measured momentum widths. Since double ionization happens mainly sequentially in our experiment, it is reasonable to assume that the first ionization step is saturated. We now can use the momentum width for single ionization (Ch1) as that of the first ionization step ($w_{1i}$). Then the momentum width of the second ionization step ($w_{2i}$) can be derived from the sum of measured electron momentum width ($w_{1i}+w_{2i}$) with the relation of $w_{2i} = \sqrt{w_{1i}^2 + w_{2i}^2}$. From the measured widths of the ion momentum distribution of Ch1 (1.24 a.u.) and Ch2 (2.42 a.u.), we obtain the momentum width of the second ionization step of Ch2 to be 2.08 a.u. With the knowledge that the second ionization of Ch3 is similar to nondissociative double ionization (Ch2) [20], we used the obtained momentum width of the second ionization step of Ch2 to get the momentum width of the first ionization step by removing an electron from low-lying MOs, which yields 1.92 a.u.. Since only one of these two electrons recombines during dissociative double ionization, the measured electron momentum distribution is determined by the recombination probabilities of the two electrons [$S_{Ch4} = \alpha_{e1} \exp(-4 \log 2 P_{e1}^2/w_{e1}^2) + \alpha_{e2} \exp(-4 \log 2 P_{e2}^2/w_{e2}^2)$]. With the obtained widths of $w_{e1} = 1.92$ a.u. and $w_{e2} = 2.08$ a.u., we performed a fitting of the electron momentum distribution with $S_{Ch4}$ and got the recombination probabilities of the first and second electron: $\alpha_{e1} = 23\%$ and $\alpha_{e2} = 77\%$. The result shows that the recombination probability of the second electron is about three times as high as that of the first electron. This observation agrees with previous measurements on D$_2$ [14] and argon atoms [36]. A straightforward explanation could be that the diffusion of the second electron wave packet is much less than the first electron at the conclusion of the laser field and thus it has a higher recapture probability.

In the end, we compare two LIDR processes (Ch4 and Ch5). Both the KER spectrum [Fig. 3(c)] and the ion sum momentum distribution [Fig. 4(b)] along the laser polarization direction of Ch5 is similar to that of Ch4, which strongly implies that the two channels undergo similar nuclear and electron dynamics. To compare with Ch4, the only difference in Ch5 is that during the dissociation one electron recombines to the CO$^+$ fragment other than the O$^+$. One clear observation is that the yield of Ch4 is much higher than that of Ch5, with a yield ratio of about 50 between electron localized to O$^+$ and CO$^+$, as shown in Table I. A straightforward explanation of this observation can be that the electronegativity of O$^+$ is higher than that of CO$^+$ which leads to a higher electron recombination probability to O$^+$. We would like to note that the dissociative recombination process observed in our experiments is different from those performed in storage rings with CO$_2^+$ in the electronic ground state [2].

In summary, we experimentally investigated the laser-induced dissociative recombination processes of CO$_2$ with coincidence detection of all involved particles. The measured KER and momentum distributions allow for understanding of both nuclear and electron dynamics during the LIDR of CO$_2$. Our measurements clearly show that LIDR of CO$_2$ originates from the recombination of one electron to one of the two ionic fragments during laser-induced dissociative double ionization. An analysis of electron momentum distributions shows that the second emitted electron has a much higher probability to recombine with the ionic fragment during the laser-induced dissociative processes. Based on the findings in this work and previous works on diatomic molecules, we believe that LIDR is a general process in strong field interaction with molecules.

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