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Enhanced ionization of an inner orbital of I_2 by strong laser fields

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Using pump-probe spectroscopy, strong field enhanced ionization is found in an inner orbital of I₂. A wavepacket is launched in the B state of I₂, whose valence orbitals are $\sigma_g^2 \pi_u^4 \pi_g^3 \sigma_u^1$, and singly ionized to the I+I⁺ dissociation channel. The ionization signal peaks at two different internuclear separations: ~ 7.3 a.u. and ~ 8.7 a.u.. The latter one shows enhanced ionization of the σ_u state which has been studied before with the I₂⁺ signal. However, the peak at smaller *R* corresponds to the enhanced ionization of the σ_g state. The peak at ~ 8.7 a.u. in the dissociating channel reveals that there could be strong mixing of different molecular orbitals when the two iodine atoms are pulled apart.

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For diatomic molecules driven by intense laser fields linearly polarized along the molecular axis, their tunneling ionization rate increases with the internuclear separation R and peaks at a critical separation R_c . This very general phenomenon, known as enhanced ionization (EI), has been found in many experiments [1–5].

Theoretically, EI has been rigorously studied in a few relatively simple cases: the one-electron molecules, such as H_2^+ , and the two-electron molecule H_2 . For the oneelectron system, the mechanism of EI is Electron Localization (EL) [6, 7] and Charge-resonance-enhancedionization (CREI) [8, 9]. It is predicted that $R_c \sim 3/I_p$ from EL and $R_c \sim 4/I_p$ from CREI, where I_p is the atomic ionization potential. Furthermore, EI is highly dependent on molecular symmetry and R_c only occurs for σ states, not π or δ [10]. For the two-electron system, it is believed that the charge transfer from the covalent state to the ionic state is the crucial step and the ionic state works as the ionization doorway state [11–14].

In our previous work [5], we investigated EI on the B state $(\sigma_g^2 \pi_u^4 \pi_g^3 \sigma_u^1)$ of I₂ by using a pump-probe technique. The pump pulse promoted one electron from the highest occupied molecular orbital (HOMO) π_q state to the lowest unoccupied molecular orbital (LUMO) σ_u state, and the probe pulse removed the electron in the σ_u state leaving the molecular ions in the deeply bound $I_2^+ X^2 \Pi_{3/2q}$ state. The vibrational wavepacket (VWP) motion in the B state provided a large range of R and R_c was found to be at ~ 8.7 a.u.. More recently [15], we produced a VWP moving between 4.4 a.u. and 6.2 a.u. in the ground electronic state (GES) $(\sigma_g^2 \pi_u^4 \pi_g^4 \sigma_u^0)$ of I₂, and discussed the *R*-dependent single ionization of different orbitals. We found that the HOMO-2 σ_g orbital showed a strong *R*-dependence, while the HOMO π_g and HOMO-1 π_u orbitals did not, and the HOMO-2 σ_g orbital provided the dominant single ionization pathway. We even predicted that the σ_g orbital would also show an R_c if the VWP moved to a sufficiently large R.

In this work, using the pump-probe technique described in Ref. [5], we obtain a large range of VWP motion in the B state of I₂ over which the *R*-dependent single ionization of the HOMO-2 σ_q orbital is observed by



FIG. 1. (color online) Schematic potential energy curves, showing the physical scenario of the pump-probe experiments.

analyzing the final state of the (1,0) dissociation channel ((1,0) refers to the dissociating channel $I_2^+ \longrightarrow I^+ + I)$, and we find an R_c at ~ 7.3 a.u. Furthermore, the HOMO-2 σ_g orbital again provides the dominant ionization pathway. However, another peak at ~ 8.7 a.u. in the (1,0) channel reveals that different molecular orbitals might be strongly mixed at large R since the (1,0) channel shows the ionization feature of the LUMO σ_u orbital [5].

The pump-probe scheme is shown in Fig.1. A VWP is launched in the B state by a weak green pump pulse through one-photon resonant excitation, and then singly ionized by a delay-controllable 800 nm probe pulse. With a 513 nm or 500 nm pump pulse, the VWP reaches $R \sim 8.72$ a.u or ~ 9.38 a.u., respectively, and the VWP motion is well understood [5]. Both of the pulses are linearly polarized along the time of flight (TOF) axis. Since the X-B transition is a parallel transition [16], the molecules excited in the B state will be aligned with the pump laser polarization with a $\cos^2(\theta)$ distribution [5] in which the



FIG. 2. (color online) The ionization signals of the I₂⁺ and (1,0) channel as a function of τ (background subtracted) with a step size of 0.030 ps with two different pump wavelengths. The delay of 0 ps is the temporal overlap of the pump and probe pulses. In the experiments, the intensities of the pump and probe pulses are estimated to be ~ 1.4×10^{11} W/cm² and ~ 1.4×10^{13} W/cm², respectively, and the I₂ pressure is ~ 7.0×10^{-7} torr.

 σ orbitals will be preferentially ionized. From the molecular orbital picture, the molecules end up in the I₂⁺ X ${}^{2}\Pi_{g3/2}$ state by removing the electron from the LUMO σ_{u} orbital, but in the (1,0) channel by removing one electron in the HOMO-2 σ_{g} orbital [17].

The experiments are performed with our home-built ultrafast Ti:sapphire laser system and a TOPAS (Optical Parametric Amplifier) system. The ionization signals of I_2^+ and I^+ are recorded with a TOF spectrometer. The ultrafast laser and the TOF spectrometer have been described in Refs [5, 18]. Room-temperature (295 K) I₂ gas is leaked effusively into the chamber with a base pressure of 2×10^{-9} torr. The Ti:sapphire laser produces linearly polarized pulses with a central wavelength of 800 nm, a transform-limited pulse duration of 37 fs, and an output energy of up to 800 μ J at a repetition rate of 1 KHz. The output beam is split with $\sim 80\%$ of the energy sent into the TOPAS to generate a pump beam while the rest serves as a probe beam. The pump pulses have a central wavelength of 513 nm or 500 nm, a duration of 50 fs and an energy of up to 2 μ J. The pump and probe beams are focused by a 3-inch-focal-length silver spherical mirror inside the TOF chamber. An aperture is used in the pump beam in order to decrease ionization and also increase the focus spot in the chamber. This creates a more uniform focal volume for the probe beam. The pump-probe delay τ is adjusted by a computer controlled translational stages. In the experiments, the probe pulse just singly ionizes the B state to avoid double ionization [19].

The ionization signal of the I_2^+ and (1,0) channel as a function of τ with different pump wavelengths is shown



FIG. 3. (color online) The ionization signal (normalized) of the (1,0) channel as a function of R with two different pump wavelengths: 500 nm and 513 nm. There are two R_c 's: the inner one at ~ 7.3 a.u. and the outer one at ~ 8.7 a.u.. The peaks at ~ 5.9 a.u. (500 nm) and ~ 5.2 a.u. (513 nm) are due to the temporal overlap between the pump and probe pulses.

in Fig. 2. With either wavelength, the I_2^+ signal shows one peak, while the (1,0) channel shows two peaks. The peaks with 500 nm pump wavelength appear earlier than those with 513 nm, since the 500 nm pump wavelength starts the VWP higher up in the potential curve of the B state of I_2 and the WVP moves faster. The ionization signals of the (1,0) channel are much stronger than those of I_2^+ with either wavelength which is consistent with our previous study [15]. The corresponding signal from the I_2^+ or (1,0) channel with 513 nm is stronger than those with 500 nm, due to the stronger X-B coupling at 513 nm [20]. There is only one peak in either of the I_2^+ curves which shows the EI of the LUMO σ_u orbital [5], and there are two peaks in either of the (1,0) channel curves, with the inner one not seen in I_2^+ and the outer one occuring at the same delay as in the I_2^+ .

Since the VWP motion in the B state of I₂ is fully understood [5], τ will determine the expectation value of R for the VWP. From this, we obtain the R-dependent ionization signal of the (1,0) channel, as shown in Fig. 3. The two different pump wavelengths give almost the same inner R_c : $R_c = 7.29 \pm 0.18$ a.u. (513 nm) and $R_c = 7.28 \pm 0.16$ a.u. (500 nm), and also the same outer R_c at ~ 8.7 a.u. (within error bar). The peaks at ~ 5.9 a.u. (500 nm) and ~ 5.2 a.u. (513 nm) are due to the temporal overlap between the pump and probe pulses.

Having established the inner R_c in the ionization signal of the (1,0) channel, we need to understand its origin. There are two possibilities: one is from ionization of an inner orbital, and the other is from resonant population transfer from the bound I_2^+ to the (1,0) channel. However, population transfer would simultaneously increase the (1,0) signal while decreasing the I_2^+ signal. As this is not seen in Fig. 2, population transfer can be ruled out. Therefore, the inner peak should be associated with ionization of an inner orbital.

The B state of I₂ has valence orbitals of $\sigma_a^2 \pi_u^4 \pi_a^3 \sigma_u^1$. The ionization of the LUMO σ_u electron will leave the molecule in the bound I₂⁺ X² $\Pi_{g3/2}$ ($\sigma_q^2 \pi_u^4 \pi_q^3 \sigma_u^0$) state, and this was fully discussed in our previous work [5] in which only one R_c at ~ 8.7 a.u. was found. On the one hand, the (1,0) channel must be from the ionization of an inner orbital, HOMO π_q , HOMO-1 π_u or HOMO-2 σ_q ; on the other hand, the R_c implies that the target orbital should show a strong R-dependent ionization. In this case, the HOMO π_g should be ruled out first, because the final state of $\sigma_g^2 \pi_u^4 \pi_g^2 \sigma_u^1$ is also bound [21]. The removal of one electron in the HOMO-1 π_u orbital could leave the molecule in a dissociating channel. Normally, removal of the HOMO-1 would leave the molecule in the bound A state, but since the HOMO is excited to the LUMO, the $\sigma_a^2 \pi_u^3 \pi_a^3 \sigma_u^1$ configuration may not be bound. However, our previous work shows that the HOMO-1 π_u orbital does not have a strong *R*-dependent ionization feature [15]. Moreover, it is predicted that π states do not show EI [10]. So, the HOMO-1 π_u orbital is also ruled out. Therefore, the $R_c \sim 7.3$ a.u. must be associated with the HOMO-2 σ_q orbital. In our previous work [15], we did see a strong R-dependent ionization of the σ_q orbital when the VWP vibrated between 4.4 a.u. and 6.2 a.u. in the ground electronic state, and we predicted that there would be an R_c if the VWP moved to larger R [15]. Now, we find this R_c .

In order to determine if this value of R_c makes physical sense, we note that $R_c \times I_p$ appears to be a useful way to characterize enhanced ionization, independent of the details of the system under study (see above and [6, 9]). Since I_p of the HOMO-2 σ_q orbital is ~ 15.07 eV (0.554 a.u.) [21, 22], we get the product of R_c and I_p to be 4.04. The product is different from that of the LUMO σ_u orbital in which $R_c \times I_p \sim 3$ [5]. The LUMO σ_u orbital only has one active electron, and it is more like a one-electron system with its R_c well predicted by EL. However, the HOMO-2 σ_g orbital has two equivalent electrons in which the ionic states should play an important role in the ionization. As we do not know of any predictions of this product for enhanced ionization in two-electron systems, we need to answer two questions: is the magnitude of the product similar to one-electron systems and do R_c and I_p scale inversely?

In order to learn more about $R_c \times I_p$ in a two-electron system, we consider a one-dimensional model of a generic diatomic molecule A_2 with two equivalent electrons moving in a double-well potential [23, 24]:

$$H_s(x_1, x_2, t) = \frac{-Z}{\sqrt{(x_1 - d)^2 + a^2}} + \frac{-Z}{\sqrt{(x_1 + d)^2 + a^2}} + \frac{-Z}{\sqrt{(x_2 - d)^2 + a^2}} + \frac{-Z}{\sqrt{(x_2 - d)^2 + a^2}} + \frac{1}{\sqrt{(x_2 - d)^2 + a^2}} - (x_1 + x_2)F(t)$$



FIG. 4. (color online) The calculation of the ionization probability as a function of internuclear separation R with two electrons initially in the ground state. The two different values of a correspond to different ionization potentials.

where Z = 1 is the charge on each well, x_1 and x_2 are the coordinates of the two electrons, R = 2d is the internuclear separation, a is the smoothing parameter of the Coulomb potential and F(t) is the electric field strength. Thoughout the calculation, atomic units are used.

In this 1D model, the parameter a affects the ionization potential I_p : the smaller the a, the deeper the ground state. a=0.700 or 0.742 corresponds to $I_p=0.90$ a.u. or 0.86 a.u., respectively. The ionization probability at different R is obtained after solving the Schrödinger equation. As it turns out (Fig. 4), R_c becomes smaller as the initial target state becomes more deeply bound. Moreover, this model gives $R_c \times I_p \sim 3.15$ (a=0.700), and $R_c \times I_p \sim 3.44$ (a=0.742), verifying the inverse relationship between R_c and I_p and that the magnitude of this quantity is between 3 and 4, as is true for one-electron systems. Thus, this corroborates our identification of the peak at 7.3 a.u. with enhanced ionization of an inner orbital.

The outer R_c in the (1,0) channel is at ~ 8.7 a.u., and this is the same as that in the I_2^+ signal since the peaks occur at the same delay. In Ref. [5], we fully studied the EI of the LUMO σ_u state in which the bound I₂⁺ X² $\Pi_{q3/2}$ state was the final state and the R_c was found at ~ 8.7 a.u.. The peak at ~ 8.7 a.u. can be considered as a signature of ionizing from the LUMO σ_u state. However, from the molecular orbital picture, the ionization of the LUMO only leaves the molecule in the bound $I_2^+ X^2 \Pi_{a3/2}$ state, not the (1,0) channel. It is a question why the (1,0) channel shows the ionization characteristic of the LUMO σ_u state. One possibility is resonant population transfer between molecular ions. However, population transfer should be weak, if there is any, since the resonant crossing is far away from 8.7 a.u., although this cannot be completely ruled out.

Another possible explanation is the mixture of differ-

ent molecular orbitals [25] in I₂ when the two I atoms are pulled apart. The molecular orbital picture works well for light molecules, like N₂, O₂ [26], CO₂ [27] and HCl [28], however, whether it still works properly for the heavy molecules, like I₂, especially when the two atoms are far away from each other, is still an open question. The heavy molecule I₂ shows different ionization characteristics from the light molecules, seen from the ionization branching ratio of different orbitals. For light molecules, the branching ratio of inner orbitals is usually rather small [28], compared with the HOMO. However, for I₂, the HOMO-2 provides the dominant single ionization pathway [15]. At large R, the σ_u and σ_g orbitals are probably mixed and the electron transfer can occur

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In conclusion, with a pump-probe technique, we generate a VWP in the B state of I₂ and study the single ionization of the HOMO-2 σ_g orbital. As predicted by previous work, an R_c is found at ~ 7.3 a.u.. This implies that the enhanced ionization also occurs in lower lying orbitals, not just the HOMO or LUMO. Another R_c at ~ 8.7 a.u. implies that there might be a mixture of molecular orbitals in I₂ at large R, since the (1,0) channel shows the ionization feature of the LUMO σ_u orbital.

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