Probing the tunnelling site of electrons in strong field enhanced ionization of molecules

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Molecules show a much increased multiple ionization rate in a strong laser field as compared with atoms of similar ionization energy. A widely accepted model attributes this to the action of the joint fields of the adjacent ionic core and the laser on its neighbour inside the same molecule. The underlying physical picture for the enhanced ionization is that it is the up-field atom that gets ionized. However, this is still debated and remains unproven. Here we report an experimental verification of this long-standing prediction. This is accomplished by probing the two-site double ionization of ArXe, where the instantaneous field direction at the moment of electron release and the emission direction of the correlated ionizing centre are measured by detecting the recoil sum- and relative-momenta of the fragment ions. Our results unambiguously prove the intuitive picture of the enhanced multielectron dissociative ionization of molecules and clarify a long-standing controversy.
Understanding and controlling the ionization dynamics of electrons from atoms and molecules in intense ultrashort laser pulses is a primary step towards the coherent control of chemical reactions and photobiological processes. In the strong-field long-wavelength regime, tunnelling is a highly successful concept used to understand the ionization process. In this picture, the effect of the intense light is described as a strong electric field suppressing the atomic binding potential, such that one or multiple electrons can tunnel into the continuum. In the case of multiple ionization of a diatomic molecule, the ionization rate is greatly enhanced if the molecular axis orients parallel to the field as compared with an atom of similar binding energy.

A widely accepted intuitive mechanism explaining this universal behaviour is referred to as 'enhanced ionization' of the molecular ion after the removal of one or more electrons from the neutral molecule. The two-centre potential binding the outmost remaining electron of a model molecular ion, superimposed with a strong electric field, is shown in Fig. 1a. The electron (green blob) is delocalized between the centres. It can tunnel into the continuum (green arrow) through the outer potential barrier. The width and height of the barrier relates to the probability of tunnelling. Figure 1b depicts the same model molecular ion with its internuclear distance stretched to a so-called 'critical' value. Here, an inner barrier has emerged between the centres, such that electron population now localizes at each of the cores. Owing to this localization, the laser-induced electric field raises the potential energy of the electron attached to the left centre (up-field core), while lowering it on the right (down-field core). To get free, the 'up-field' population only needs to tunnel through the inner barrier between the centres, which is considerably more likely than tunnelling through the outer barrier between the 'down-field' centre and the continuum. Hence, within this picture, preferential ionization of the 'up-field' ion as compared with the 'down-field' one is expected, dramatically increasing the overall ionization probability.

Such enhancement of the multiple ionization rate for molecules has been theoretically predicted and experimentally observed for many years. However, the cornerstone of the underlying physical picture, that is, it really is the 'up-field' atom that gets ionized, stands yet untested. In this respect, some works even contradict the predictions of the outlined scenario, which is still a puzzle.

The reason for the missing experimental test of this heart of the enhanced ionization model is twofold. First, in a symmetric single-colour laser field, the up-field and down-field directions alternate. It is usually unknown at which swing of the field an electron escapes and the corresponding ion is formed. Equal flux of any charge state to both sides is observed. Second, if the symmetry of the electric field is broken either by using a carrier-envelope phase locked few-cycle pulse or a two-colour pulse, then the asymmetric laser field can potentially redistribute the remaining electrons in the left behind molecular ion. This is a highly interesting phenomenon in itself. Nevertheless, it can overwrite the directionality of the initial enhanced ionization step, prohibiting a conclusive experimental test of the enhanced ionization scenario.

We overcome these hurdles by using a single-colour elliptically polarized femtosecond laser pulse. We do not break the symmetry of the laser field. Instead, we measure the instantaneous direction of the laser electric field at the moment when the electron escapes for each ionization event. This is achieved by detecting the direction to which the electron is streaked by the elliptically polarized light. We then measure in coincidence if the ion that has released the electron is up- or down-field with respect to this instantaneous field direction.

**Results**

**Experimental setup and target.** We choose the van der Waals force-bound ArXe dimer as a clean target to model the stretched diatomic molecule. It has several advantages for probing the enhanced ionization scenario. First, the equilibrium internuclear distance of the neutral dimer (R = 8.0 a.u.) is already in the 'critical' range, enabling the enhanced ionization process without beforehand stretching. Second, the influence of the molecular orbitals on the ionization is avoided as the electrons are localized at the atomic centres right from the beginning. Third, the ArXe dimer allows us to remove the first electron from the Xe site and then the second electron from the Ar site owing to their very different ionization energies (I_Xe = 11.9 eV for Xe and I_Ar = 15.9 eV for Ar). We hence can probe the enhanced ionization by studying the release of the second electron and the ejection direction of Ar⁺. Although here ArXe dimer is used as a clean system to test the enhanced ionization scenario, the mechanism of enhanced ionization is not special to this system but ubiquitous in molecular ions. The measurements were performed in a reaction microscope of cold target recoil ion momentum spectroscopy (see Methods).

The magnitude and direction of the electric field at the instant of ionization are encoded in the final, detectable momentum vector of the electron. The rotating electric field of an elliptically polarized pulse leads to a final electron momentum approximately perpendicular to the instantaneous field vector owing to 'angular
streaking \(^{24-30}\). Hence, the field direction at the instant of ionization is mapped to the momentum direction of the released electron. For illustration purposes, as sketched in Fig. 1c, if an electron is freed when the laser field vector points along \(+y\), its final momentum \((p_z)\) will be along \(−z\). Likewise, an instantaneous field along \(−y\) will lead to a final electron momentum along \(+z\). Therefore, any possible asymmetry in ionization rate of the ArXe oriented along the \(y\)-axis (up/down in Fig. 1c) will be turned into an asymmetry of the electron momentum distribution along the \(z\)-axis (left/right in Fig. 1c). Technically, instead of detecting electron momentum directly, we detect the momentum vector of the centre of mass of the Coulomb exploding Ar\(^+\)Xe\(^+\) ion. By momentum conservation, this centre-of-mass or sum momentum of the recoil ions \((p_{\text{sum}})\) is equal and opposite to the sum momentum of the two emitted electrons. Meanwhile, in addition to this centre-of-mass motion of the molecular ion, the ionic cores repel each other, yielding a much bigger relative momentum \((p_{\text{rel}})\) along the former molecular axis. Hence, we obtain the information of the direction of the emitted electrons in the molecular frame of reference by only detecting in coincidence the 3-dimensional momenta of both fragment ions.

**Ion momentum distributions of doubly ionized ArXe.** We focus on the Coulomb exploding double-ionization channel Ar\(^+\) + Xe\(^+\). The elliptical light suppresses recollision of the freed electrons, thus the two electrons escape sequentially\(^{27}\) during the laser pulse. As the laser field is elliptical, and the tunnelling probability depends steeply on the field strength, each ionization step will maximize when the field vector is along the major polarization axis, yielding two burst of ionization per cycle. As depicted in Fig. 2, we find that double ionization maximizes for molecules lined up with the major axis. We restrict our data analysis and the discussion to such molecules by selecting only fragmentations within a cone of 45\(^\circ\) around the major polarization axis.

The first ionization step of removing an electron from Xe will necessarily be symmetric, that is, ionization when the laser field points in the \(+y\) and \(−y\) directions has equal probability, yielding symmetric momentum distribution of the first electron along the \(z\)-axis. No field direction-dependent ionization rate of the first electron is expected as it starts from the neutral molecule. The second step of ejecting an electron from the neutral Ar with the neighbouring Xe\(^+\) now is the one for which we test the enhanced ionization scenario. A preferred directionality of this second electron will lead to an asymmetry in the ion sum-momentum \((p_{\text{sum}}^2 = p_{\text{Ar}}^2 + p_{\text{Xe}}^2)\) distribution. If we do not distinguish between emission of Ar\(^+\) to the \(+y\) or \(−y\) directions, the \(p_{\text{sum}}^2\) for Ar\(^+\) + Xe\(^+\) is symmetric (grey squares in Fig. 3a). It is the convolution of the momentum distributions of two sequentially removed electrons. As first and second electron can both be emitted either in the \(+y\) or \(−y\) direction, the combination yields four possible sum momenta\(^{27,28,31}\) as indicated with the orange bars in Fig. 3a (see Methods). The solid grey curve shows a fit of the convolution by assuming a Gaussian distribution of each electron momentum (see Methods), which returns \(p_{\text{sum}}^2 = 0.45 ± 0.07\) and \(p_{\text{sum}}^2 = 0.61 ± 0.02\) a.u. in agreement with the fact that the second electron from Ar is released by a higher laser intensity in the laser pulse than the first one from Xe.

**Discussion.**

We will now address the essential question whether or not the second ionization step happens at the up-field core, as predicted by the classical model of enhanced ionization. We break the symmetry of the problem by gating on the orientation of the Ar\(^+\) + Xe\(^+\) breakup. In Fig. 3a, the red squares show an enhancement in rate at positive \(p_{\text{sum}}^2\) when the Ar\(^+\) departs along \(+y\). This shows that the Ar\(^+\) ion is born on the up-field site of the molecule in our clockwise light polarization. The relative-momentum \((p_{\text{rel}} = p_{\text{Ar}} - p_{\text{Xe}})\) distribution in Fig. 3b very intuitively shows this behaviour when we select events where the field at the release of the second electron pointed to \(+y\) by gating on \(p_{\text{sum}}^2 > 1.0\) a.u. for clockwise elliptical light. It shows a strong orientation of the Ar\(^+\) + Xe\(^+\) breakup with the Ar\(^+\) being up-field. The same curves are recorded with the opposite light helicity, that is, anticlockwise polarization. The results shown in Fig. 3c,d are mirror images of the clockwise light polarization.
case, confirming the validity of our data. All these results conclusively show that the second electron is more favourable to be freed from the up-field potential well.

To obtain the relative emission probabilities of the second electron from the up-field and down-field potential wells, we fit the asymmetric $p_{\text{sum}}$ distributions with the same model used above, but now featuring two separate fitting factors $A_{+}$ and $A_{-}$ for the magnitudes of the $p_{\text{pz}}>0$ and $p_{\text{pz}}<0$ cases instead of a common one (see Methods). The fits (red and blue curves in Fig. 3a) reveal that the probability for the second electron being released from the up-field potential well is about $-2.41 \pm 0.16$ higher than that from the down-field potential well.

In conclusion, we give a direct proof that enhanced multielectron ionization of a diatomic molecule is indeed more likely to happen at the up-field than at the down-field centre. This result clarifies the long-time controversy on the electron localization-assisted multiple ionization of molecules.

Methods

Experimental technique. We performed experiments with femtosecond laser pulses (35 fs, 790 nm, 8 kHz) produced from a multipass amplifier Tsapphire laser system (KMLabs Dragon). The laser pulses were sent into a standard cold target attachment (KMLabs Dragon). The laser pulses were sent into a standard cold target adapter before the vacuum chamber. The fast axis of the quarter-wave plate was fixed to be along the major axis of the elliptically polarized light. The peak intensity and ellipticity of the pulse were measured to be $9.3 \times 10^{14}$ W cm$^{-2}$ and $e^{-0.68}$, respectively.

Two-particle correlation identifies ArXe isotopes. As shown in Fig. 4a, there are seven Xe isotopes that can form ArXe dimers in our gas jet. The sharp stripes displayed in Fig. 4b owing to the ion–ion time-of-flight correlation allow us to identify and distinguish the ArXe isotopes. All the isotopes show similar behaviour, we use the Ar$^{133}$Xe and Ar$^{135}$Xe for the discussion in this work as they are best separated from the others.

Convolution model of the ion sum-momentum distribution. As the recollision process is mostly suppressed in the elliptically polarized pulse, the two electrons are predominantly ejected sequentially$^{77}$. As the tunnelling probability strongly depends on the field strength, the ionization steps maximize when the laser field is orientated along the major axis (Fig. 2), yielding streaked electron momentum along the polarization minor axis (Fig. 1c). Owing to momentum conservation, the sum of all electron momentum vectors is the inverse of the sum of all ion momentum vectors. Therefore, by measuring the latter, we can determine the former.

The sum-momentum distribution of the sequentially released two electrons can be modelled as the convolution of them. As illustrated in Fig. 5, each individual electron freed when the laser field vector parallel to the $y$-axis results in two peaks centred at $\pm p_{\text{pz}}$ (Fig. 5a) or $\pm p_{\text{pz}}$ (Fig. 5b) along the $z$-axis. The convolution of them yields four possible momenta (Fig. 5c). The central peaks, $p_{\text{pz}}$, are filled with events that two electrons are freed when the laser field points to opposite directions (back-to-back). The side peaks, $p_{\text{pz}} + p_{\text{pz}}$ and $-p_{\text{pz}} - p_{\text{pz}}$, account for the cases that both electrons are released by the laser field along the same direction. In more detail, by assuming a Gaussian shape of the electron momentum distribution, that is, $A_{ij}/[\sigma_{y}(2\pi)^{1/2}]\exp\left[-0.5(p_{\text{pz}} - p_{\text{sum}})^{2}/\sigma_{y}^{2}\right]$, the recoil ion sum momentum (that is, the reverse of sum momentum of the sequentially freed two electrons) can be expressed as$^{78}

$$
p_{\text{sum}ij}(p_{\text{pz}}) = \frac{1}{\sqrt{2\pi}} \sum_{y} A_{12ij} \exp\left[-0.5(p_{\text{pz}} - p_{\text{sum}})^{2} / \sigma_{y}^{2}\right]
$$

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Author contributions

J.W. designed the experiments and performed the measurements and data analysis; M.M., L.Ph.H.S., M.K., S.V., H.S., H.K., T.J., A.C. and R.D. provided technique support during the measurements and data analysis; J.W., M.M. and R.D. wrote the paper.

Additional information

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