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Velocity map photoelectron-photoion coincidence imaging on a single detector

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Here we report on a new simplified setup for velocity map photoelectron-photoion coincidence imaging using only a single particle detector. We show that both photoelectrons and photoions can be extracted toward the same micro-channel-plate delay line detector by fast switching of the high voltages on the ion optics. This single detector setup retains essentially all the features of a standard two-detector coincidence imaging setup, viz., the high spatial resolution for electron and ion imaging, while only slightly decreasing the ion time-of-flight mass resolution. The new setup paves the way to a significant cost reduction in building a coincidence imaging setup for experiments aiming to obtain the complete correlated three-dimensional momentum distribution of electrons and ions. © 2012 American Institute of Physics. [http://dx.doi.org/10.1063/1.4749843]

I. INTRODUCTION

Charged particle imaging spectrometers or “reaction microscopes” employing the velocity map imaging technique have become a proliferating tool in reaction dynamics studies today. The determination of the full three-dimensional momentum distribution of the fragment products (ions and electrons in an ionization event) in coincidence provides complete kinematic information of the event. Time-resolved coincidence measurements are particularly useful in studying the dynamical behavior in many body systems like polyatomic molecules. Many versions of photoelectron-photoion coincidence spectrometers have evolved over the last two decades to study molecular photoexcitation and ionization processes.1–13 The temporal and spatial resolution of electrons/ions imaged has also improved enormously, especially by incorporating the velocity map imaging technique.14, 15

In 2008, a photoelectron-photoion coincidence imaging apparatus to study femtosecond molecular dynamics was constructed in our laboratory in LaserLaB Amsterdam with temporal resolution down to tens of picoseconds (∼16 ps for electrons) and spatial resolution of ions down to few hundred microns (∼115 μm for Xe+ ions).11 Using time switched voltages on open lenses, electrons and ions were velocity focussed onto two micro-channel-plate delay line detectors in opposite directions. A common characteristic of almost all coincidence detection setups reported in literature is the usage of two opposite facing time-of-flight (ToF) detectors. In a significant development, we now show that it is possible to extract and detect both electrons and ions in coincidence onto a single detector by time switching the lens voltages appropriately. We have used one half of the existing coincidence detection apparatus, i.e., the electron detector of the two-detector setup, and have successfully measured electrons and ions in coincidence using a single velocity map imaging spectrometer with total flight length of 11 cm. Because we have used the electron detector of the two-detector coincidence setup as the detector for the single coincidence setup, the electron resolution is unaltered and will not be discussed here. The details are described in Ref. 11. The performance of the new one-detector coincidence setup is described in terms of ion mass resolution and coincidence data using femtosecond ionization data of Xe and CF3I at 400 nm, respectively.

The present paper is organized as follows. In Sec. II, we present the apparatus. In Sec. III, we report the experimental data and we compare with data obtained using the standard two-detector coincidence imaging setup. We summarize our conclusions in Sec. IV.

II. EXPERIMENTAL

A. Vacuum and molecular beam

The coincidence imaging apparatus has been described in detail before and will be discussed only briefly here.11, 16, 17 It consists of three differentially pumped UHV chambers, the source chamber, a buffer chamber and the imaging chamber. Improving upon the first series of experiments, which were done using a continuous molecular beam source,11 in the present setup we use a home-built high-repetition rate (up to 5 kHz) cantilever piezo valve producing a pulsed molecular beam.18, 19 A pulsed molecular beam source matches well the duty cycle of the experiment because of the pulsed femtosecond laser source which operates at a repetition rate that is selectable between 1–5 kHz. The pulsed cantilever valve makes it possible to produce stronger beams and to use higher backing pressures. The piezo valve can operate both in pulsed mode and continuous mode and the gas was expanded through a nozzle with a diameter of 200 μm. In the present experiments, we operate the laser and the pulsed piezo valve at 1 kHz. Low seeding ratios of about 1% of CF3I in neon and 5% Xe in neon are used, so as to avoid clusters in the molecular beam. The typical speed ratio with 3 bars backing pressure is about S = 25–30.16

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B. Laser system

The regen femtosecond laser (Spectra Physics Spitfire Pro) operates at 1 kHz and central wavelength of 800 nm. For multi-photon excitation of Xenon and CF$_3$I frequency doubled light of 400 nm with an energy of 15–20 μJ and a duration of about 170 fs is used. A half-wave plate is used to keep the polarization parallel to the plane of the imaging detectors. The laser beam is focused in the coincidence apparatus and crosses the molecular beam with a spot size of approximately 100 μm.

C. Single detector coincidence setup

In our new single detector coincidence setup presented here, we use only one position and time sensitive particle delay line detector (Roentdek large anode DLD40X$^{20}$) for both electron and ion coincidence detection, see Fig. 1(a). The total ToF length is optimized for electron imaging, and equals 11 cm from the interaction region to the detector with a field-free ToF region of 5.25 cm. The charged particle lenses are used in pulsed mode to extract the electrons and ions toward the detector. The schematic of the operation is shown in Fig. 1(b). The repeller (R), extractor (E), and extra lens (L) are first set with appropriate negative voltages for electron velocity map imaging, i.e., R = −525 V, E = −385 V, and L = −270 V. Subsequently, about 200 ns after the laser interaction the voltages on all three lenses are quickly switched to a geometry for ion extraction, i.e., R = 2000 V, E = 1320 V, and L = 550 V. The positive voltages for ion extraction are retained for about 20 μs, sufficiently long for all ions to reach the detector. Subsequently, the voltages are switched back again in about 10 μs to electron extraction mode, so that when the next laser pulse arrives (1 ms inter pulse duration) the setup is ready for detection of electrons. All voltages are switched at the repetition rate of the laser system (1 kHz).

The HV switching causes ringing on the delay line and may lead to false events in the data. However, these false events due to switching are easily detected and rejected in the recorded TDC (time-to-digital converter) events of ions, as its time of occurrence (i.e., between 0.2 and 1.5 μs) is after the electron hit and before the arrival of most ions. The electrons are detected within 200 ns while the fastest ions arrive typically after about 1.5 μs, sufficiently long for the imaging conditions. Therefore, all events recorded in between are rejected during data processing. Thus, the pick-up due to switching does not interfere with our correlated (e,ion) events. A typical TDC signal accumulated for 100 laser shots is shown in the bottom panel of Fig. 1(c). The electron, ringing, and ion signals are indicated.

III. RESULTS AND DISCUSSION

A. Ion mass resolution

To characterize the ion mass resolution we have recorded the ion ToF spectrum obtained from multiphoton excitation of Xe with the frequency doubled laser output centered at 400 nm, see Fig. 2. The strongest five isotopes are clearly resolvable, the two weakest isotopes (128Xe and 130Xe) are discernible. The total ToF time is 3.8 μs with a FWHM $\Delta t = 2$ ns (for the strongest five isotopes). This corresponds to a mass resolution $\Delta m_{\text{FWHM}}/m \approx 1/950$.

The shorter length of the ToF tube in the case of the single detector setup essentially reduces the arrival time resolution and thereby the achievable mass resolution of the
ions. This can be compared to the ion ToF-spectrum acquired using the two detector setup as reported before. In the two detector case, the ion ToF tube had a length of 29.5 cm and the ions experience a total flight length of 35.2 cm from the interaction region to the detector. The reported ToF for the Xe isotopes was 8.3 μs with a FWHM Δt = 2 ns. This corresponds to a mass resolution ΔmFWHM/m ≈ 1/2000 for masses near 130 amu. The shorter length of the flight tube in our new single detector coincidence setup reduces the ion mass resolution by a factor of two (ΔmFWHM/m ≈ 1/950) while retaining the spatial resolution. However, for experiments with masses of m ≥ 14, the mass resolution is still adequate enough to discriminate isotopes and mass peaks with Δm ≥ 2 will be totally separated. One of the problems of the shorter ToF tube in the present single detector setup is detection of ion masses m ≤ 14, as they coincide with the switching pickup on the detector. The ringing time cannot be easily reduced, and therefore the extraction field would have to be reduced for low masses. For a longer ToF tube, the ringing is never a problem since the first ions will arrive long after the ringing. The short length ToF was optimized for electron detection as part of the two-detector coincidence setup. It is certainly possible to design a better version with slightly longer tube clearly separating electrons, ions of all masses, and the intermediate pickup in time. Also, the short ToF length is not ideal for high-resolution three-dimensional slice imaging detection of ions. Cases where the ionic fragment have low kinetic energy (or without any resolvable structure) are seen to have roughly the same spatial resolution (spread) irrespectively of the ToF length.

B. Single detector coincidence measurements

Coincidence measurements on CF3I following single color (400 nm) multiphoton excitation have been previously reported using the two-detector coincidence setup. To demonstrate that our novel single detector coincidence setup can be used to study reaction dynamics without the loss of information, we have repeated the previous measurement using only the single detector setup. The results are depicted in Fig. 3. Panel (a) shows the total ion ToF spectrum (left) and the photoelectron spectrum (right) of all events measured in coincidence. (b) Time sliced electron image, photoelectron spectrum, and ion image (insert) measured in coincidence with CF3I+ ions. (c) Time sliced electron image, photoelectron spectrum, and ion image (insert) measured in coincidence with CF3+ ions. (d) Time sliced electron image, photoelectron spectrum, and ion image (insert) measured in coincidence with iodine ions. The photoelectron kinetic energy values are listed in Table I. The ion images are shown for the same detector area.
TABLE I. Appearance energy/ionization potential for the production of the different ions after multiphoton ionization of CF$_3$I at 400 nm. The excess energy is given with respect to four-photon ionization ($4 \times h\nu = 12.4$ eV), excess energy = $12.4 - AE/IP$.

<table>
<thead>
<tr>
<th>Ion/State</th>
<th>AE/IP (eV)</th>
<th>Excess energy (eV)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CF$_3$I$^+$, e$^-$</td>
<td>11.384 (Ref. 23)</td>
<td>1.03</td>
</tr>
<tr>
<td>I$^+$, e$^-$</td>
<td>12.78 (Ref. 24)</td>
<td></td>
</tr>
<tr>
<td>CF$<em>3$I$^+$ (X$^2E</em>{3/2}$), e$^-$</td>
<td>10.37 (Ref. 25)</td>
<td>1.31</td>
</tr>
<tr>
<td>CF$<em>3$I$^+$ (X$^2E</em>{1/2}$), e$^-$</td>
<td>11.10 (Ref. 26)</td>
<td>2.04</td>
</tr>
</tbody>
</table>

By fast switching of the high voltages on ion optics we demonstrate that it is possible to obtain the complete three-dimensional momentum distribution of electrons and ions in coincidence using only a single detector. This new approach would reduce the building costs of a new coincidence imaging setup significantly while not compromising much on the experimental sensitivity and mass resolution.

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