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Understanding the modulation mechanism in resonance enhanced multi-photon probing of molecular dynamics

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Abstract

Time resolved spectroscopy on isolated molecules gives fundamental insight into the conversion of light energy to other degrees of freedom. Probing of the photoinduced dynamics can be accomplished by ionization, via a single- or multi-photon transition. In this letter we directly contrast transient spectra on the molecule perylene obtained with multi-photon ionization (MPI) to single-photon ionization (SPI). The photoinduced nuclear geometry relaxation modulates the MPI transient with a decay time constant of 0.9 ± 0.2 ps. In contrast, the SPI transient completely lacks any indication for relaxation. We attribute this difference to a change in resonance enhancement of the MPI probe as the molecular geometry changes. Our results underline the importance of a detailed knowledge about these resonances for a proper interpretation of transient signals of molecular dynamics subject to nuclear and electronic relaxation effects. At the same time, the direct comparison to SPI directly demonstrates the higher sensitivity of resonance enhanced MPI as a probe in time-resolved dynamical studies.

The conversion of photon energy into other forms of energy is often happening with high selectivity and efficiency [1]. Time-resolved photoelectron (PE) and photoion (PI) spectroscopy have proven to be important tools for the investigation of photoinduced relaxation processes in isolated molecules [2]–[4], since their observables can be simulated from theoretical investigations with high accuracy and thus directly compared [5], [6]. The molecular relaxation generally leads to an increased ionization potential [7]. In order to detect the dynamics via a single photon ionization (SPI) scheme, increased probe photon energies in the vacuum ultraviolet (VUV) [8]-[13] have been recently applied. Experimentally less demanding and therefore widely used is multi photon ionization (MPI) [2], [4], [14], [15], as shown for metal carbonyls [16], [17] or nucleobases [18]–[21]. As pointed out before, the transient MPI signal can be enhanced by resonant excited molecular states [14]–[17]. However, the role that the resonances play in actively shaping the observable transient in a pump/MPI-probe experiment has not been investigated. In this paper, we fill this gap by comparing SPI and MPI for identical molecules and excitation conditions finding stark differences in the transient spectra obtained by the two processes. The comparison of the SPI and MPI transients together with laser intensity characteristics directly reveals the active influence of the resonances on the observable in MPI probing.

Fig. 1 shows a sketch of the important processes in time-resolved PE and PI spectroscopy. An excitation (pump) laser pulse promotes the molecule from the ground state (S₀) into an electronically excited state (S₁). A second ultrashort laser (probe) pulse interacts with the excited molecule after a controlled delay. It can ionize the excited molecules either via SPI or MPI. In order to ionize the molecule via SPI, the photon energy has to be larger than the excited state ionization potential [7]. For this case, the probe laser field projects the excited state nuclear wave packet onto one or more electronic continua. The resulting PI and PE spectra are measured as a function of pump-probe pulse delay. The ionization yield as well as the PE energy are modulated by the molecular dynamics if the SPI step is sensitive to nuclear and/or electronic relaxation. Alternatively to SPI, the ionization energy

can be overcome by several photons in MPI schemes [2], [14]–[21]. Here, relaxation processes are generally observed as transient changes of ion (fragment) yields. As shown in Fig. 1, the MPI signal can be enhanced by intermediate resonant states.

For our systematic study, we choose the polycyclic aromatic hydrocarbon perylene. It presents a conceptually simple case performing *only* nuclear dynamics after photoexcitation with negligible internal conversion and intersystem crossing [22], [23] (excited population thus stays in the S₁ state in Fig. 1). We will show that for the resonantly enhanced MPI, the transient ion signal exhibits a transient signal decay of 0.9 ± 0.2 ps. In contrast, the SPI shows absolutely no modulation in the PE energy. In the following, we will discuss the differences in the transient signals of the two schemes in combination with additional information about the MPI process from laser intensity scans and from time-dependent density functional theory (TDDFT) calculations.



Figure 1 (color online): Sketch of the electronic states of perylene and excitation/ionization schemes. Shown are the ground state (S_0) , the first excited state (S_1) , resonant intermediate states and the lowest ionic state (D_0^+) for S_0 and S_1 minimum geometries. After 400 nm excitation the molecule is either ionized by SPI with a single 89 nm VUV photon or via 800 nm MPI. During vibrational relaxation in the excited state the ionization potential $(S_1-D_0^+)$ distance) remains nearly constant, while the energy distance to states that lead to resonance enhancement of the MPI probe increases.

The experiments are carried out with a recently developed time resolved PE and PI spectrometer [24]. A molecular beam of perylene (Sigma Aldrich, purity > 99.5%) is introduced into the interaction region by a resistively heated capillary-oven [25].

Following 400 nm pump excitation, the molecular dynamics is probed after a time delay either using SPI or MPI. For use in SPI, VUV probe pulses are obtained from high-order harmonic generation in Ar gas [26]. The 9th harmonic (89 nm, 14 eV photon energy) is isolated from the high harmonic continuum by using the bandpass characteristics of an indium metal filter [24]. The isolated 9th harmonic pulse duration is 35 ± 8 fs, its spectral width is 190 ± 60 meV. For MPI the 800 nm laser fundamental is used. The pump-probe time resolution of 65 ± 10 fs is deduced from the rise time in the pump-probe spectra and applies for MPI and SPI [24].

For VUV SPI we detect PE spectra with a magnetic bottle electron time of flight spectrometer and deduce molecular dynamics from a transient change of the PE kinetic energy. We choose a pump pulse intensity of $\sim 4.10^{10}$ W/cm² to excite about 17% of the perylene molecules to the S₁ state [24]. Although PEs are produced by the 400 nm pulse alone due to some small multi-photon ionization

probability, excitations to states higher than S_1 are negligible for the pump-probe signal, as no PE bands are detected above 10 eV (Fig. 2a).

For MPI we deduce the molecular dynamics from a transient change of the ion yield. Here we do not show MPI-PE spectra since their interpretation requires the decomposition of electrons emerging from several ionic continua. We therefore operate the time of flight spectrometer in ion mode. Sufficiently low laser intensities are chosen to obtain negligible signals from the pump or probe laser alone, as compared to the pump-probe signal.

In addition to our experimental results, we perform geometry optimizations and excitation energy calculations on the TDDFT level of theory as implemented in the GAMESS-US program package [27]. The ground state geometry is optimized using B3LYP/def2-SVP. Excited state energy and geometry optimization calculations are carried out with TDB3LYP/MCP/NOSeC-V-DZP. Since both S₀ and S₁ minima exhibited D_{2h} symmetry, all calculations are carried out in D_{2h}. Ionization potentials are calculated with OVGF/QZVP as implemented in Gaussian 09 [28].

A comparison of SPI and MPI probing for transient pump-probe spectroscopy of perylene is shown in Fig. 2. The SPI PE spectrum of unexcited perylene obtained with harmonic number nine (14 eV, 89 nm) consists of five distinct PE bands in the kinetic energy range of 0-7 eV and fits the literature data [24], [29]. Our resolution suffices to clearly separate the lowest lying ionic state continuum (7 eV kinetic energy) from all the other continua. The lowest ionic states (c.f., Fig. 1) give rise to the two photoelectron peaks centered at 5.0 and 6.8 eV [30] (Fig. 2a). A preceding 400 nm (3.1 eV) pump pulse excites a fraction of the molecules to the S_1 ($1B_{2u}$) state and its ionization leads to the appearance of two new PE peaks at 8.1 and 9.8 eV in the PE spectrum (Fig. 2a), reflecting the excess potential energy. The difference of pump-probe (400 nm-H9, gray) spectrum and the time-reversed probe-pump (H9-400 nm, black) signal, serving as the background correction, is depicted in Fig. 2b. It clearly shows the two excited state PE peaks. The strong modulations of the difference signal below 8 eV reflect a decreased ionization yield from S_0 overlapping with emerging PE signal due to ionization from S_1 [24]. We fit the PE signal corresponding to ionization from S_1 to the ionic ground state D_0^+ at 9.8 eV by a Gaussian (red curve in Fig. 2b). Figures 2c and 2d show the peak position and amplitude, respectively, of the Gaussian as function of pump-probe time delay. We fit the data by the product of an error function and an exponential term with variable amplitude (red) [31], [32]. The 9.8 eV PE peak position shows essentially no delay dependence. Considering the 95% confidence interval of the exponential decay amplitude (red dashed lines) we find a negligible PE signal shift of $-0.2 \pm 0.5\%$. Note that Fig. 2c shows the peak positions within a narrow range of 0.4 eV. From this time-delay independent character we conclude that either no relaxation takes place in the S₁ state or the ionization potential from S_1 to D_0^+ is constant within an experimental uncertainty of <1% during relaxation from S_0 to S_1 geometry.



Figure 2 (color online): Comparison of H9 (89 nm) SPI probing with 800 nm MPI probing for perylene pump-probe photoionization.

a) PE spectra obtained with 400 nm-H9 pump-probe (gray), H9-400 nm probe-pump (black), 400 nm pump only (red) and H9 probe only (blue). Traces are additionally shown with a magnification of 10 for better visibility.

b) Difference spectrum obtained as pump-probe minus probe-pump (black) together with a Gaussian fit of the PE peaks at 9.8 eV (red).

c) Position of the 9.8 eV PE peak as obtained from the Gaussian fit as function of the pump-probe time delay. A fit with the product of an error function and an exponential term with variable amplitude (red curve) [31], [32] reveals a negligible PE signal shift of $-0.2 \pm 0.5\%$. The red dashed lines mark the 95% confidence interval of the exponential term amplitude.

d) Same as in c) for the PE peak amplitude. The fit gives a 15 \pm 10% increase with a 1.2 ps time constant.

e) Transient PI signal for 800 nm MPI probing observed at the perylene parent ion mass at 252 u (red curve) and for the total ion yield (blue curve, vertically offset by 0.5). Fits to the same function as in c) and d) reveal decays of $32 \pm 3\%$ (252 u) and $34 \pm 3\%$ (total ion yield), both with a 0.9 ± 0.2 ps time constant [32].

The transient position of the 9.8 eV PE peak obtained with SPI is now compared to transient 800 nm MPI signals, which are shown in Fig. 2e for detection of the unfragmented parent ion $(C_{20}H_{12}^{+} \text{ at } 252 \text{ u}, \text{ red curve})$ and the total ion yield (blue curve). Surprisingly, both transient PI signals show a strong

signature for molecular relaxation. After the fast increase around zero delay, which represents the pump-probe cross correlation [24], we observe exponential signal decays by $32 \pm 3\%$ (parent ion) and $34 \pm 3\%$ (total ion yield), both with a time constant of 0.9 ± 0.2 ps [32]. This is in good agreement with time scales for intramolecular vibrational redistribution reported in literature [33]. In addition to the parent ion, we observe the ion fragments $C_{20}H_{10}^+$ (250 u, 5% of the parent yield), $C_{20}H_8^+$ (248 u, 1%), $C_{19}H_8^+$ (236 u, 7%), and the doubly charged parent ion $C_{20}H_{12}^{++}$ (126 u, 2%). These ions result from ionization to higher ionic continua and exhibit an exponential decay comparable to that of the parent and total ion yields. From the equivalency of all ion signal decays we conclude that fragmentation in the ionic ground state with increasing pump-probe delay, due to energy gain during relaxation in the S_1 state, does not take place. Hence, the transient parent ion signal (Fig. 2e, red curve) corresponds to MPI to the ionic ground state D_0^+ and should therefore represent the same time dynamics as the SPI-PE signal. Obviously, the decaying parent ion yield in MPI for increasing pump-probe delay is in strong contrast to the constant character of the SPI-PE peak position (Fig. 2c). From the unshifted PE peak in the SPI transient one would conclude that no molecular relaxation occurs and thus expect that the MPI transient ion yield would also be not modulated.



Figure 3 (color online): a) Laser intensity scans for 800 nm (red, 252 u detection, slope = 3.60 ± 0.18), and 400-800 nm pump-probe MPI (blue, 252 u detection, time delay 0.3 ps, slope = 1.67 ± 0.13). Due to saturation of the process the data points (crosses) for the highest laser intensities deviate from the expected linear dependency. Only encircled data points are included in the fit. b) Slopes for 400-800 nm pump-probe MPI at different time delays.

To obtain a deeper insight into this discrepancy we determine the order of the MPI process. For MPI, the ion yield as function of the laser intensity shows a linear dependency in a log-log plot and its slope is related to the order of the ionization process. Figure 3a presents the unfragmented parent ion yield as function of the 800 nm laser intensity for MPI from the S₀ ground state (red) and for MPI from the S₁ excited state with preceding 400 nm excitation (blue). Due to the vertical ionization potential of 7 eV of ground state perylene [29] a process order of five is expected for 800 nm (1.55 eV) MPI. The observed order of 3.60 ± 0.18 demonstrates the presence of intermediate states, which lead to resonant enhancement of the ionization process. The non-integer value is due to intensity averaging over the focused laser beam [34], [35]. With preceding 400 nm (3.1 eV) pump excitation to the S₁ state we observe a process order of 1.67 ± 0.13 . This reduction by a factor of two reflects the

absorption of one 400 nm photon. It demonstrates the presence of resonant intermediate states also for MPI from S₁ which we will use in the following to explain the discrepancy of the transient SPI and MPI signals. To obtain insight into the transient behavior of these intermediate resonances, we measure the S₁ MPI process order for different pump-probe time delays (Fig. 3b). Although the error bars overlap, the graph indicates an increase of the observed slopes from 1.67 ± 0.13 at 0.3 ps delay to 1.83 ± 0.09 at 4.7 ps.

Based on the resonant character of the MPI probe process, we are able to explain the origin of the dynamic modulation *purely based on experimental data*. The constant PE energy in the SPI transients can only result from a constant energy difference of the cationic ground state D_0^+ and S_1 along the nuclear relaxation coordinate. Thus, the modulation in the transient MPI signal must be a consequence of a different energy change related to the intermediate resonant state(s). Indeed, the changing order as a function of delay also supports this assignment. The decrease of the transient results from an intermediate state(s) shift out of resonance with respect to the laser spectral center. The amplitude of the SPI-PE peak as a function of the delay is shown in Fig. 2d. Unlike the constant peak position, the amplitude shows an increase of $15 \pm 10\%$ with a 1.2 ps time constant. This points towards an increase of the transient (Franck-Condon factor and/or dipole moment) between S_1 and D_0^+ during molecular relaxation from S_0 to S_1 geometry. The temporal behavior of the 8.1 eV PE peak shows the same transient trends in position and amplitude, albeit with increased noise presumably because of the mentioned modulations of the difference signal due to overlapping PE bands.

Although the experimental results are self-conclusive, we perform geometry optimizations and excitation energy calculations on the TDDFT level of theory to support our findings. Despite the uncertainties of the TDDFT calculations especially regarding states with strong Rydberg character close to the first ionization potential (IP), our calculations predict the presence and relative shift of intermediate resonances. We find an almost parallel decrease in potential energy of S_1 and of the D_0^+ for relaxation from the Franck-Condon to the S₁ minimum nuclear geometry [36]. Over this path, the S_1 - D_0^+ potential energy difference increases by only 50 meV. This is in agreement with our SPI data and the observed shift of the 9.8 eV PE peak by -20 ± 50 meV ($-0.2 \pm 0.5\%$), which is negligible with respect to the spectral width. Our calculations also predict the presence of intermediate states that lead to resonant enhancement of MPI from both the S₀ and S₁ state, in agreement with the observed MPI orders (Fig. 3). For MPI from S_0 (gerade symmetry) these are an ungerade state at 4.98 eV ($2B_{1u}$) and a gerade state at 6.36 eV ($2B_{1g}$) [36]. For MPI from S_1 (ungerade) two gerade states ($3B_{3g}$, $4B_{3g}$) at 4.22 eV and 4.67 eV and five ungerade states (5B_{1u}, 5B_{2u}, 6B_{2u}, 6B_{1u}, 7B_{2u}) in the range of 6.1 eV to 6.3 eV can lead to resonant enhancement. For these intermediate states the potential energy distance to S_1 depends much more on the molecular geometry than for the cationic ground state D_0^+ . The energy gap to S_1 increases from S_0 to S_1 minimum geometry by 50 meV and 130 meV, respectively, for the two gerade states and by 130 meV to 330 meV for the five ungerade states. A comparison to the transform limited laser bandwidth of 70 meV shows that the magnitude of resonance enhancement decreases from S_0 to S_1 geometry. This is directly observable in our MPI measurements as decrease in ion yield (Fig. 2e) and increase of the process order (Fig. 3b) with progressing pump-probe delay.

Distinct resonance-enhanced ionization channels can be formulated for the two ionization schemes (800 nm and 400/800 nm) based on state symmetry considerations and Koopmans correlations. In the case of 800 nm ionization from the ground state (S_0 , $1A_g$), the energy of 5 photons (each 1.55 eV) is needed to overcome the first IP of 7 eV to the lowest cationic state [29]. The first step is a three-photon transition, since states of suitable energy and symmetry are missing for a single- or two-photon transition. The most probable candidate is the $2B_{1u}$ state at a calculated potential energy

of 4.98 eV [36]. It exhibits the character of a HOMO-LUMO+4 excitation. Absorption of a further photon leads to an energy region just below the IP where a plethora of states with suitable symmetry and character can be expected. From there the molecule can be ionized with an additional photon to the lowest cationic state (D_0^+) , which is characterized by removal of an electron from the HOMO. For 400 nm-800 nm two color ionization, a possible ionization channel leads in a first step to the lowest excited state ($1B_{2u}$, HOMO-LUMO excitation) upon absorption of a 400 nm photon. By absorption of one 800 nm photon each, the molecule is most likely further excited to the $4B_{3g}$ state (4.67 eV), which exhibits at least partly the character of a HOMO-LUMO+1 excitation, and to the energy region just below the IP. From there single-photon ionization is possible.

In summary, we present a systematic comparison of SPI and resonant MPI to study photoinduced dynamics in perylene, a conceptually simple molecule only exhibiting ultrafast intermolecular vibrational relaxation after photoexcitation into the S₁ state. The two probing schemes, which are widely used in time resolved experiments [2], [8]–[21], exhibit strong differences in their sensitivity to this relaxation processes. While photoelectron spectroscopy based on SPI does not show any modulation with pump-probe delay, the MPI shows a clear signal decay within one picosecond. We establish the presence of intermediate resonances for the MPI probe step via laser intensity scans. The time dependent SPI photoelectron spectroscopy indicates a constant energy difference of the S_1 and D_0 state over the relaxation path and in addition a negligible decay of population into other product states, as anticipated due to the high fluorescence yield of the molecule [33]. We therefore conclude that the decay observed in MPI must necessarily result from the resonances shifting as the molecule relaxes. We qualitatively support this experimental argument by TDDFT simulations. This finding is generally important for the interpretation of ultrafast MPI transients in molecules with more complex relaxation pathways, since signatures arising from IVR cannot be distinguished from signatures of non-Born-Oppenheimer processes within the experimental method. The distinction can only be made with the help of quantitatively correct results from quantum chemistry. High level guantum chemical calculations are needed even to interpret results from SPI investigation of non-Born-Oppenheimer dynamics. The requirements for theory in the MPI case has to be even higher, since the calculation of highly excited neutral states necessarily requires a higher level treatment of electron correlation to predict acceptable energy differences. We therefore regard single photon probing a better method for a systematic investigation of underlying molecular dynamics. Although high harmonic sources have only been used on conceptually simple molecules [8]–[13] so far, application of SPI on more complex systems such as nucleobases, base pairs and base pair models [37] will be possible.

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