Ultrafast dynamics of polycyclic aromatic hydrocarbons initiated by highly intense laser fields


We present an investigation of the ultrafast dynamics of the polycyclic aromatic hydrocarbon fluorene initiated by an intense femtosecond infrared laser pulse and probed by a weak visible pulse. Using a multichannel detection scheme (mass spectra, electron and ion velocity-map imaging), we provide a full disentanglement of the complex dynamics of the vibronically excited parent molecule, its excited ionic states, and fragments. We observed channels resulting from both multiphoton- and tunnel-ionization regimes. In particular, we observed the formation of the unstable tetracation of fluorene, above-threshold ionization features in the photoelectron spectra, and evidence of ubiquitous secondary fragmentation. We produced a global fit of all observed time-dependent photoelectron and photoion channels. This global fit includes four parent ions extracted from the mass spectra, 15 kinetic-energy-resolved ionic fragments extracted from ion velocity map imaging, and five photoelectron channels obtained from electron velocity map imaging. The fit allowed the extraction of 60 transient lifetimes corresponding to different photoinduced intermediates.

1 Introduction

Polycyclic aromatic hydrocarbons (PAHs) are a large class of molecules that consist of carbon and hydrogen atoms. Each PAH contains at least two rings, of which at least one is aromatic. These molecules play an important role in astrochemistry since they are estimated to account for about 10% of all galactic carbon.

PAHs can also form in combustion reactions of fossil fuels, which lead to the presence of PAHs in the Earth's atmosphere. Here, they act as environmental pollutants, and they impact the health of the population.

In the atmosphere, PAHs are also subjected to high-energy particles. Therefore, investigating the photoinduced chemistry of PAHs is an interesting topic.

Previously, the photoinduced ionization and fragmentation of PAHs upon strong-field excitation-ionization were studied with tabletop laser setups. In particular, laser pulses with peak intensities of about ~10^14 W/cm^2 interacting with PAHs have produced ionic species up to tetracations. Investigations of the ultrafast ionization and fragmentation of PAHs induced by highly energetic XUV photons have recently gained attention in the scientific community for tracing down mechanisms of photochemical reactions. The ultrafast pump-probe studies with ultrashort XUV pulses produced either by high-harmonic generation (HHG) sources or free-electron lasers (FELs) revealed rich dynamics of charge state evolutions up to PAH triations. Femtosecond strong-field laser pulses, produced by Ti:Sa lasers, were also used to induce dynamics in benzene and toluene. These studies found interesting dynamics, such as oscillations between vibrational states in toluene or conical intersection relaxations in Jahn-Teller benzene cations.

In this work, we present an investigation of the dynamics of a three-ring PAH fluorene (C_{13}H_{16}, FLU). The XUV-induced dy-
namics of this molecule have recently been studied in detail.\textsuperscript{13,15} Therefore, it is interesting to investigate how the observed dynamics change in a different excitation regime. We used ultrashort strong-field infrared (IR) laser pulses to excite FLU and subsequently probed it using a weaker visible (VIS) pulse. The pump pulse peak intensity was estimated to be $3 \times 10^{13}$ W/cm$^2$, which corresponds to a Keldysh parameter value of $\gamma = 2.1$.\textsuperscript{20} for the single ionization of FLU. Based on the calculated $\gamma$ value and considering a non-uniform distribution of intensities in the laser pulse, we expect to observe a mixed strong-field regime between tunneling ionization (TL, $\gamma \ll 1$) and multiphoton ionization (MPI, $\gamma \gg 1$). The interaction of strong-field laser with FLU results in the production of electrons and multiple ions, which were measured using a double-sided velocity map imaging (VMI) spectrometer. VMI enabled the disentangling of the various photodissociation fragmentation pathways on the basis of momentum. In addition, the time-of-flight (TOF) mass spectra (MS) of the ions were measured. This multi-channel data accumulation allowed for a detailed and comprehensive decomposition of the strong-field-induced dynamics of FLU. In this manuscript, we describe the experimental details, data analysis, and theoretical calculations. This is followed by pump-probe independent results, dynamical observations, and conclusions.

Fig. 1 Experimental MS of FLU obtained with only the IR laser pulse for different intensities. The spectra were normalized to the maximum intensity peak of the FLU monocation ($C_{13}H_{10}^+$, $m/z = 166$) observed with the unattenuated data corresponding to a laser peak intensity of $3.1 \times 10^{14}$ W/cm$^2$. The absolute values of the ion yields ($|IY|$) were taken to allow for a logarithmic scale since the baseline oscillates around zero signal and can have negative values. Gray-shaded areas highlight the four peaks of interest: $C^+$ (m/z = 12), FLU$^{3+}$ (m/z = 55.3), FLU$^{2+}$ (m/z = 83), and FLU$^+$ (m/z = 166). The dependencies of the intensities of these peaks as a function of laser intensities are given in ESI.

2 Methods

2.1 Experimental details

The IR (810 nm) – VIS (405 nm) pump-probe experiments were carried out at the permanent CAMP end-station\textsuperscript{21} at beamline BL1, at the free-electron laser in Hamburg (FLASH)\textsuperscript{22} facility. A Ti:Sa laser system\textsuperscript{23} produced pulses at 810 nm. Most of each pulse was used as an IR pump, and a part of each pulse went to the second harmonic generation (SHG) to produce a weaker VIS probe pulse of 405 nm. The duration (full width at half maximum, FWHM) of the IR pulse was 50 fs, while the duration of the VIS pulse was estimated to be less than 150 fs. We measured the ion yield produced as a function of pump-probe delay time. The arrival time of the pulse was varied from -1 ps to +1 ps in 15 fs steps relative to the pump pulse using a mechanical delay stage.

The CAMP end-station houses a double-sided electron and ion VMI spectrometer,\textsuperscript{24} enabling the measurement of ion and electron kinetic energies simultaneously. An open port in the vacuum chamber at the CAMP end-station allowed for the installation of a source chamber with the Even-Lavie valve (ELV),\textsuperscript{26} two subsequent skimmers, and a differential pump for sample introduction into the experiment.

The sample of FLU was purchased from Sigma-Aldrich and Thermo Fisher with 98% purity and used without further purification in two different beamtimes performed in 2018 (F-20170540) and 2023 (F-20211752), respectively. In the experiment, fluorene powder was heated to 200$^\circ$C in an in-vacuum reservoir, implemented in the high-temperature ELV, in order to obtain sufficient vapor pressure. The gas-phase molecules were then introduced into the vacuum chamber by means of a helium carrier gas with a backing pressure of 1 bar. A pulsed ELV-produced molecular jet was doubly skimmed on the way to the interaction region to obtain a well-collimated molecular beam.

Three types of data obtained from the experiments have been used for the analysis, namely, the TOF-MS, the electron VMI, and the ion VMI. Each ion and electron dataset was acquired using a combination of microchannel plate (MCP) in chevron configuration and phosphor screens, P47 and P20, respectively. The TOF-MS were obtained directly from the MCP readout with the help of a 2 GHz analog-to-digital-converter (ADQ412AC-4G-MTCA). The velocity map images of the ionic species, together with lower-resolution TOF-MS, were obtained using a Pixel Imaging Mass Spectrometry (PlmMS2) sensor.\textsuperscript{27} Lastly, the electron data was obtained directly from a CCD (in 2018) or CMOS (in 2023) camera readout.

2.2 Data analysis

Data extraction and analysis were performed using home-written scripts. To retrieve and analyze the data from the TOF-MS and electron VMI data, a combination of the BeamtimeDAQAnalysis\textsuperscript{28} and CAMPFancyAnalysis\textsuperscript{20} Python libraries was used. For the analysis of the PlmMS data, a home-written set of LabView scripts and the CAMPFancyAnalysis library were employed.

A detailed description of the recoil-frame covariance analysis for the ion VMI data can be found elsewhere.\textsuperscript{16,17,19} The photoelectron-photoion covariance analysis (PEPCOV) was performed in the following way:\textsuperscript{26,27} First, each single-shot electron image was converted into the radial distribution function (RDF), and each single-shot TOF-MS was converted into the mass-overflow (m/z) MS. The centers of the images and the momentum calibration were determined based on the helium II photoelectron data, which was obtained in pre- and post-measurements for the same molecular beam with 41 eV XUV radiation from FLASH, pub-
lished in Refs. [13,15]. Then, the covariance map was obtained as

\[ \text{cov}(\text{IY, PEY}) = \langle \text{IY} \cdot \text{PEY} \rangle - \langle \text{IY} \rangle \cdot \langle \text{PEY} \rangle , \]

where “IY” denotes ion yield at a given m/z value, “PEY” denotes photoelectron yield at a given image radius value, and \( \langle \ldots \rangle \) denotes averaging over all shots. A partial covariance correction for fluctuations in the pressure in the chamber was also applied [35].

Then, along the image radius axis of the map, a direct Abel inversion transformation was performed, as implemented in the PyAbel package [36].

The observed dependencies of the ion and electron yields as a function of pump-probe delay time were fit with standard expressions [33,41] using the MC² Fitting software [32] with a previously described algorithm [14]. A simultaneous fitting was done for the multiple channel data for the parent ions (from the high-resolution TOF-MS), momentum-disentangled fragment ion yields (from the ion VMI), and photoelectron yields (from the electron VMI). Fitting was done for 40 datasets of ion yield versus pump-probe delay time, which comprise momentum-resolved fragment ion yields of ions having 5 to 13 carbon atoms, the parent monocation, dication and trication, dehydrogenated parent monocation, momentum-resolved dications of fragment ions having 5, 7, and 11 carbon atoms and electron yields corresponding to five photoelectron bands. Fitting multiple datasets together enabled an estimation of the overlapping time of the pump and probe pulses \((t_0)\). Using this \(t_0\), five momentum-resolved fragment ions with more than one transient feature were also fitted, giving out 15 datasets.

The results of the data analysis are given in the Electronic Supporting Information (ESI).

### 2.3 Computational details

To calculate the MS of FLU, augmented Born-Oppenheimer molecular dynamics (aBOMD) simulations of the fragmentation of FLU were done using the DissMD script from the PyRAMD software [33].

The algorithm was based on the QCEIMS idea [14,31] and the modifications are described in detail in Refs. [13,44,46]. The forces for aBOMD were computed with the xTB software [37] at the semi-empirical GFN2-xtB level of theory [25] that was shown to be suitable for the fragmentation pattern predictions [29]. To emulate the high-energy excitation regime, the excitation energy was represented as \(n \cdot h \nu\), where \(n\) is the number of photons, and \(h \nu\) is the photon energy, which was taken to be 1.5 eV (corresponding to a wavelength of 810 nm). The parameter \(n\) was sampled from the Poisson distribution with a mean number of photons \((n) = 40\), and the result was adjusted to fit in the interval of \(6 \leq n \leq 100\) by setting the outlier \(n\) value to the closest value of \(n = 6\) or \(n = 100\).

The energetics of the reaction \(\text{C}_2\text{O}^{2+} \rightarrow \text{C}_2^+\), discussed later in section 3.2.5, were computed at CCSD(T)/aug-cc-pVTZ [25,53] using the ORCA 5 software [53]. The geometries of \(\text{C}_2\text{O}^{2+}\) were optimized for the singlet, triplet, and quartet multiplicities, with the triplet being the lowest energy state. For the carbon monocation, doublet and quadruplet states were computed, with the doublet found to be the lowest in energy. All the geometries and energies are given in the ESI.

### 3 Results and discussion

#### 3.1 Roadmap of the analysis

In the following two sections, we will present the analysis of the strong-field-induced reaction dynamics of FLU. First, we will discuss the main time-independent features observed in the photoion and photoelectron signals without consideration of the pump-probe effects. There, we will focus on the individual spectra of the species and on the covariance techniques that allow to find relations between the different observables. Afterward, we will look into the pump-probe dependencies of the discussed signal, extracting the information on the ultrafast photochemical processes in FLU.

#### 3.2 Time-independent observations

##### 3.2.1 Mass spectra

The behavior of benzene, benzene derivatives [16,17,55], and PAHs [55] in strong laser fields have been investigated for decades. The general fragmentation patterns of PAHs are very similar: at low peak field intensities (∼10^13 W/cm^2) [5], only single ionization is visible in the MS. As the peak power is increased, a rise of the parent monocation formation is observed as well as the formation of the parent dication and monocationic and dicationic fragments. At high peak powers (∼10^14 W/cm^2), trications and even tetracations occur in the MS for PAHs [55].

In the first set of experiments, in which FLU was subjected to IR pulses of variable intensity, we also observed this ubiquitous behavior, which can be seen in Figure [1]. We see the production of the highly-charged parent species up to the trication (FLU^{3+}), as well as a subsequent rise in the production of the fragment ions upon increasing the IR pulse peak power. The first, second, and third sequential ionization potentials (IPs) of FLU are 7.9, 13.1, and 17.4 eV, respectively [18], therefore, to reach the tricationic state by MPI, 26 photons of 1.5 eV energy (810 nm) would be required, which is highly improbable. Thus, trications are probably formed mostly through sequential TI (\(\gamma = 2.1/2.7/3.1\) for first/second/third ionization), or in the mixed MPI+TI regime, which is agreed upon in the literature [53,58].

##### 3.2.2 Photoelectron spectra

The photoelectron spectrum (PES) of FLU averaged over all pump-probe delays (∼2 ps in total) is shown in Figure [2]. The experimental results clearly show above threshold ionization (ATI) features as distinct peaks with equal spacing corresponding to the photon energy of the IR pulse (1.5 eV) [29]. The ATI peaks were fitted using the expression

\[
E_k = (n_{\text{min}} + k) h \nu - W ,
\]

where \(E_k\) is the position of the \(k\)-th ATI peak \((k = 0, 1, 2, 3, 4)\), \(n_{\text{min}} = 6\) is the minimal number of IR photons with energy \(h \nu = 1.5\) eV to produce ionization of FLU, and \(W\) is the fitted work function. From the fit (see ESI for details), we obtain \(W = 8.4\) eV, which is comparable to the first IP of FLU (7.9 eV).

The ATI features are clear signatures of MPI processes. The photoelectron–photoion covariance (PEPICOV) analysis revealed that the disentangled PES corresponding to all the ionic fragments...
look very similar and resemble the PES of the whole system, with the same set of ATI peaks (see dashed PEPICOV lines in Figure 2). This indicates that the MPI regime dominates TI in producing low-charge parent ions and fragments. Additionally, the general mechanism of fragmentation implies obtaining first a highly excited mono- or dicationic state of the parent, which later decays into the charged fragments, thereby providing similar PES.

### 3.2.3 Recoil-frame covariance

Recoil-frame covariance analysis was performed on the mass-resolved VMI data to distinguish the various measured fragmentation channels. This approach was shown to be very effective in disentangling the fragmentation pathways of complex PAH molecules. In brief, this method correlates the relative positions of two ions (the reference ion and the ion of interest) on the imaging detector. If the ion of interest is produced simultaneously with the reference ion, we see a recoil feature, a positive covariance in the opposite direction to that defined by the reference ion, which we choose to be upwards. More information on this technique can be found in Ref. 31.

When we look at the covariance images obtained from the strong-field excitation (Figure 3), we see a major difference from the results observed upon the ionization and fragmentation induced with the single-photon XUV [13]. In the case of XUV excitation-ionization, the recoil features appeared on the diagonals, corresponding to \(C_2H_x^-\)-fragment loss (shown as dashed boxes in Figure 3). However, in the strong-field regime, we see distinct features off these diagonals, e.g., the \(C_3H_x^+\) ion of interest recoiling against \(C_2H_y^+\), or \(C_4H_x^+\) against \(C_3H_y^+\). Such off-diagonal signals indicate oddcarbon loss fragmentation channels, which were not observed in the XUV regime.

We also see recoil relation between the smallest fragments: \(C_3H_x^+\) ion of interest against \(C_2H_y^+\). A significant broadening of the recoil feature (see enlarged insets in Figure 3) may indicate that they are not produced in the same dissociation event but sequentially. In this case, the additional fragments would carry additional momentum, blurring out the recoil feature of the correlated ions.

A significant asymmetry in the intensities of the observed covariances was noted. On the one hand, in Figure 3 we see clear signals for the small reference ions (\(C_3^+\)–\(C_5^+\)) of interest recoiling against larger monocations (\(C_6^+–C_{11}^+\)) in the pairwise dissociation of the parent and of the acetylene-loss fragment \(C_{11}H_2^+\). On the other hand, the inverted covariances, e.g., \(C_{10}H_y^+\) ion of interest against \(C_3H_x^+\), are significantly less intense (however, present). This asymmetry is also a signature of sequential fragmentation, in which the fraction of events producing stable large fragments is much lower than the fraction of metastable large fragments that undergo further fragmentation, producing smaller ions without larger counterparts.

### 3.2.4 Formation and fragmentation of tetracation

While the trication \(FLU^+\) is clearly visible in the MS data (Figure 1), we have not observed any traces of the intact FLU tetracation (\(FLU^{2+}\)) in the MS data. However, we see evidence of the parent tetracation fragmentation in the ion VMI of dicaticonic fragments, namely \(C_7H_x^{2+}\) and \(C_{11}H_x^{2+}\) recoiled from a doubly-charged counterion. In Figure 3 the pump-probe delay averaged VMI image for the \(C_7H_x^{2+}\) ion and the corresponding angularly-integrated ion momentum distribution are shown. In the VMI im-
the hydrogen abstraction acetylene addition (HACA) mechanism for PAH fragmentation. Such preference was observed in analogous XUV-induced mass spectra.

2.5 Feature at m/z = 12

A distinctive feature appears in the strong-field induced mass spectra of substituted benzenes and PAHs that is not observed in analogous XUV-induced mass spectra. This feature is a peak at m/z = 12, which can either correspond to C\(^{2+}\) or to C\(_2\)H\(^{2+}\) (see Figures 1 and 5). On the one hand, if this fragment is mainly C\(_2\)H\(^{2+}\), this would strongly indicate a preferred acetylene loss mechanism for PAH fragmentation. Such preference was observed in the XUV-induced fragmentation patterns, which can be related to the hydrogen abstraction acetylene addition (HACA) mechanism of PAH formation that can be important due to the presence of the C\(_2\) molecule in space, for instance in comets. On the other hand, if the m/z = 12 peak corresponds to C\(^{2+}\), that could indicate a high-energy stochastic fragmentation of the backbone.

We do not observe any conclusive recoil-frame covariance features for this fragment. However, this might also be due to it displaying the strongest distortion of the ion VMI among all fragments, which makes the center determination for this ion complicated. The momentum distribution map for this ion has three dissociation channels that can be interpreted as (1,0), (1,1), and (1,2) or as (2,0), (2,1), and (2,2) (see Equation 2), which does not favor either of the interpretations. However, the TOF-MS data is more in favor of the C\(^{2+}\) interpretation of this peak: we see two peaks that are positioned at m/z = 12 and m/z = 13 (Figure 5). This pattern suggests that this feature corresponds to C\(^{2+}\) because, in case of the dication, we would expect m/z = 12 for C\(_2\)H\(^{2+}\), m/z = 12.5 for C\(_2\)H\(^{2+}\), and m/z = 13.0 for C\(_2\)H\(^{2+}\). The isotope 13C in this case does not explain the observed pattern since, in the natural abundance, it is expected to be around 1% of the most intense peak, while the ratio of the peaks at m/z = 12 and m/z = 13 is about 2:1, respectively (Figure 5).

To support this assignment from the mass spectrum, we have calculated the equilibrium dissociation energy \(D_e\) for the reaction C\(_2\)H\(^{2+}\) \(\rightarrow\) 2C\(^+\) at the CCSD(T)/aug-cc-pVTZ level of theory. The obtained value of \(D_e = -5.3\) eV shows that this reaction is strongly exothermic, which means that C\(^{+}\) is a more energetically favorable ion. The estimated lifetime of the metastable triplet C\(_2\)H\(^{2+}\) into two C\(^+\) ions is 0.9 ns (see ESI for details), which makes the detection of this species with MS impossible. Using abOMD simulations of the fragmentation dynamics, we also observe peaks at the m/z region of interest (12 \(\leq m/z \leq 14\)), which all correspond to CH\(_x^{+}\) species (Figure 5). The further fragmentation of CH\(_x^{+}\) up to C\(^+\) could not be traced with the semi-empirical method used for abOMD. Nevertheless, from a combination of experimental data and simulations, we conclude that the features around m/z = 12 in the high laser power-induced MS of PAHs indeed correspond to C\(^+\).

3.3 Pump-probe-dependent results

Now, we present the analysis of the time-resolved pump-probe measurements for our system. The pump pulse was a strong-field IR (810 nm) laser pulse with an estimated peak power of \(3 \times 10^{13}\) W/cm\(^2\). This pulse initiated the ionization and fragmentation dynamics in FLU, which was described in the previous section. The probe pulse was a weaker VIS pulse (405 nm) produced from the same laser system. The comparison of the pump-only, probe-only, and pump-probe averaged MS is given in the ESI. We monitor var-

---

**Fig. 4** (a) Ion VMI of the C\(_7\)H\(^{2+}\) fragment from FLU. The left side of the image is the raw experimental data, while the right part is the result of filtering and Abel inversion with the BASEX method. (b) Angularly integrated momentum distribution function of the C\(_7\)H\(^{2+}\) ion with a three-Gaussian profile fit. The shaded areas indicate the individual fit components that represent (2,0), (2,1), and (2,2) channels.
ious observables (various ion and electron yields) as a function of the delay between pump and probe pulses ($t$). We chose the temporal overlap of both pulses ($t_0$) to be at $t = 0$. The region in which the probe pulse comes before the pump pulse is $t < 0$, and the region when the probe comes after the pump is $t > 0$.

The time-dependent changes in the MS, PES, and a typical ion momentum distribution map of the C$_{13}$H$_{y}$ + ion are shown in Figure 5. The momentum map illustrates the evolution of the momentum of the ion extracted from the ion VMI (as in Figure 4(b)) as a function of pump-probe delay $t$. In the pump-probe-dependent MS, we see similar trends for the fragmentation dynamics of PAHs as were observed with the XUV-pumped experiments. In the $t < 0$ region, we see more of the parent species than in the $t > 0$ region. For the fragments, this trend is inverse: for $t < 0$, we see fewer ions than for $t > 0$. Such behavior indicates that the pump pulse produces an excited state ensemble, and the probe pulse destroys it by pumping even more energy to overcome fragmentation thresholds. This is observed for the PAH in both XUV-excited regimes probed with IR and VIS pulses as well as for benzene and toluene excited by a strong-field laser.

Near the temporal overlap of the two laser pulses ($t = 0$), there is a strong increase in the ion production of all kinds of ions, as well as an increase in the yields of the ATI peaks. The PES (Figure 6b) shows qualitatively the same ATI spectrum given by Equation 1 being present in the whole range of the pump-probe delays. The new photoelectron features, different from the ATI pattern, do not appear in the PES because probe photons are just twice the energy of the IR pump photons. The transient increase in each ATI peak’s intensity arises from transferring each $k$-th ATI peak into

![Fig. 5](image_url) Experimental and aBOMD-calculated theoretical MS of FLU (upper trace) and disentangled theoretical MS (lower trace). Top: full m/z range, bottom: zoom for a region between m/z = 10 and m/z = 17.

![Fig. 6](image_url) Experimental parts of the pump-probe dependent MS (a) and photoelectron spectrum (b) of FLU, and (c) ion momentum distribution map of the C$_{13}$H$_{y}$ + fragment. Horizontal lines in (b) indicate the positions of the ATI peaks from Figure 2. Purple arrows in (b) indicate the probe photon conversion of the ATI electrons. Horizontal lines in (c) show the approximate delimiters of the (1,0), (1,1), and (1,2) channels of the C$_{13}$H$_{y}$ + ion. The full m/z-range version of the (a) is available in ESI.

### Table 1

<table>
<thead>
<tr>
<th>ATI band order $k$</th>
<th>Lifetime, fs</th>
<th>Parent species</th>
<th>Lifetime, fs</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>102 ± 4</td>
<td>C$<em>{13}$H$</em>{10}$ +</td>
<td>62 ± 1</td>
</tr>
<tr>
<td>1</td>
<td>93 ± 3</td>
<td>C$<em>{13}$H$</em>{9}$ +</td>
<td>91 ± 1</td>
</tr>
<tr>
<td>2</td>
<td>87 ± 3</td>
<td>C$<em>{13}$H$</em>{10}$ +</td>
<td>79 ± 1</td>
</tr>
<tr>
<td>3</td>
<td>90 ± 4</td>
<td>C$<em>{13}$H$</em>{10}$ +</td>
<td>42 ± 1</td>
</tr>
<tr>
<td>4</td>
<td>83 ± 3</td>
<td>C$<em>{13}$H$</em>{10}$ +</td>
<td>42 ± 1</td>
</tr>
</tbody>
</table>

The transient lifetimes observed in various photoelectron and photon yields. The left part of the table contains lifetimes of transients in ATI bands of various orders with photoelectron energies given by Equation 1. The right part shows ion yields of parent FLU ions (C$_{13}$H$_{10}$ +, C$_{13}$H$_{10}$ +, C$_{13}$H$_{10}$ +) and monodehydrogenated FLU monocation (C$_{13}$H$_{10}$ +). The uncertainties of the lifetimes are the 1σ values obtained from the Monte-Carlo sampling.
the \((k+2)\)-th peak by a probe VIS photon with 3 eV of energy (see Equation 1 and vertical arrows in Figure C for details). The fitted lifetimes of the transients in the ATI bands seem to decrease with increasing order of the band, going from 102 ± 4 fs to 83 ± 3 fs (Table 1). However, they are all centered around 90 fs, which might indicate that the source of these ATI bands is the same set of excited states of the neutral FLU parent molecule.

Based on the channels observed in XUV-pumped investigations and the time-independent results observed in the IR-only case from the previous section, the general scheme for the observed strong-field induced dynamics of FLU in the IR-VIS regime can be described as summarized in Figure 7. Here, we will briefly describe it. Upon interaction with the IR pump pulse, FLU can produce multiple parent species, from the excited neutral molecules (FLU\(^0\)) to tetracations (FLU\(^{4+}\)). The probe VIS pulse can ionize the near-threshold species, providing the observed transient increase in the production of the higher-charge parent ions, which we directly detect from the MS. The excited parent species (FLU\(^0\), FLU\(^+\), FLU\(^2+\), FLU\(^3+\), and FLU\(^4+\)) can fragment, producing the fragments in different momentum channels \((n,m)\), i.e., \((0,0)\), \((1,0)\), \((1,1)\), and so forth. The probe photon can increase the fragmentation charge state by one, switching from a \((n,m)\) dissociation channel to a \((n+1,m)\) channel. The fragments can spontaneously fragment further into smaller species. However, the probe pulse can also enhance this fragmentation in the smaller fragments of the \((1,0)\) channel, leading eventually to the C\(^+\) fragment, for which we see a transient increase in production. All the fitted lifetimes are shown in Figure 8 and Table 1 and all the fitted pump-probe dependencies of the ion yield and numerical values are given in ESI.

In general, the strong-field induced dynamics are in many aspects similar to the dynamics observed by the XUV pulses. This is direct evidence of general photochemical properties of the PAHs, dictated by their molecular and electronic structures, such as the closely connected carbon framework and a highly conjugated nature of \(\pi\)-bonds. However, there are some noticeable differences with the XUV regime, which are signs of the strong-field excitation character. Here, we will focus the discussion on the channels that were not observed with XUV excitation.

First, let us discuss the dynamics of the parent ions, given in Table 1. Unlike before, with the IR pump, we see short-lived transients in the monocations, corresponding to the lifetime of excited neutral FLU. We observed the transient peaks in both the intact parent FLU monocation \((C_{13}H_{10}^+)\) and in its monodehydrogenated version \((C_{13}H_{9}^+)\). The lifetime extracted from the \((1,0)\) channel yield is 30 fs longer than the lifetime from \((1,1)\), which is most likely a sign of the unresolved delay for the loss of hydrogen. The lifetime of the near-threshold FLU\(^+\) observed in the yield of the parent dication \(C_{13}H_{10}^{2+}\) is longer than for the excited neutral. This could be due to a larger density of the excited states with the increase of the charge state. The fitted lifetimes of the near-ionization-threshold FLU dication \(FLU^{2+}\), observed through the pump-probe dependent Y of \(C_{13}H_{10}^{3+}\), re-

---

**Fig. 7** Reaction scheme of the observed pump-probe dynamics in FLU under the influence of an intense IR pump pulse (indicated by a red arrow) and probed with a VIS pulse (indicated as curved violet arrows). The gray dashed arrows indicate possible evolution pathways of the excited-ionized ensemble of molecules. Species in bold font were observed in the experiment, whilst the normal font indicates unobserved or uncharged (and thus invisible in the MS) species.

**Fig. 8** All fitted relaxation lifetimes of the different fragmentation-related processes, extracted through the global fitting of the pump-probe dependent observables. The \(x\)-axis indicates the size of the fragment in which the corresponding lifetime was observed, and the \(y\)-axis shows lifetimes on a logarithmic scale. The bold font in the legend indicates in which channel of the fragment ion the transient was observed. The \((1,\ldots)\) and \((2,\ldots)\) bold channels' notation indicates that the lifetime was observed either in the monocations or dications fragment, respectively.

**Fig. 9** An example of the pump-probe fit for the \((1,0)\) channel of the \(C_{13}H_{9}^+\) fragment. All the fits are available in the ESI.
revealed a significantly shortened lifetime (42 ± 6 fs), compared to those obtained with the XUV pump (126 ± 16 in the XUV-IR experiment and 184 ± 44 fs in XVUVVIS).[15] This drastic difference arises from a difference in the distribution of the population of the FLU2+ excited states obtained in the strong-field and single-XUV-photon ionization regimes.

In the kinetic-energy resolved IV of the monocationic fragments (Figure 5 and “Peak 1” in Figure 9), we see the same channels as in the XUV experiments,[13][15] namely (1,0)→(1,1) and (1,1)→(2,1) (Figure 2 and ESI). However, because the low-energy IR pump photons do not necessarily ionize the molecule, a new channel was observed, namely (0,0)→(1,0), similar to that in the parent monocations (see Figure 7). This channel has a much shorter lifetime on the order of a few fs to 10 fs than the analogous process in parent ions (FLU0→FLU+) with a lifetime of 62 ± 1 fs (see Table 1 and Figure 8).

Another previously unseen channel corresponds to a slow dynamics with lifetimes of 0.1 – 1 ps observed in the (1,0) channels of small ions (CHx+ to C8Hx+). An example of such a channel can be seen in Figure 9 denoted as “Peak 2”. This dynamic feature is a transient decrease for the fragments from C2Hx+ to C8Hx+, but for the CH4+, it is a transient increase. Therefore, we attribute this channel to the secondary dissociation of the larger ions, eventually ending up with the smallest CHx+ fragment. The slow transient lifetimes could be attributed to the relaxation lifetimes of the vibronically excited fragment ions that eventually fragment further upon the interaction with the probe pulse, leading to transient decay of its own ion yield and corresponding transient increase in the ion yield of the CHx+ species. Such fragmentation is consistent with observations from the static recoil-frame covariance analysis.

The last new set of channels that have not been observed before for FLU in the XUV-pump regime[15] are the (2,1) and (2,2) channels of the dicationic fragments (C7Hx2+ and C11Hx2+), as shown in Figure 6. These channels indicate the lifetimes of the excited FLU trication (FLU3+) and tetracation (FLU4+). The lifetimes of these channels have similar values and show similar trends, as those observed in monocationic fragments for the (1,0)→(1,1) and (1,1)→(1,2) channels (Figure 5). This might indicate that all these fragments are formed through a set of similar excited states of the di- and trications.

4 Conclusions
We investigated the behavior of fluorene (C13H10), a three-ring polycyclic aromatic hydrocarbon, under the influence of strong-field laser pulses in the mixed multiphoton-ionization and tunneling regime. We have identified the photoion and photoelectron channels that can be ascribed to either of these strong-field ionization or excitation regimes. For instance, the above threshold ionization peaks in the photoelectron spectra are clear signs of the multiphoton regime, while the triply charged parent cations are probably appearing through tunneling ionization.

In this work, we provided a complete disentanglement and characterization of the photochemical dynamics of fluorene in the strong-field regime. Excitation by intense IR pulses tends to induce more secondary fragmentation and double and triple ionization than single-photon XUV ionization, which is seen from the recoil-frame covariance analysis and pump-probe dynamics. The strong-field excitation also opens new channels that were not observed in the XUV ionization. First of all, this is the dynamics of the near-threshold neutral molecule, which produces previously unobserved features appearing from the presence of the excited neutral molecule. The second new observation is the tetracation formation, which was observed via the (2,2) channel of the dicationic fragments C7Hx2+ and C11Hx2+.

All the observed time-dependent channels in photoion and photoelectron spectra were included in a global fit. This fitting dataset consisted of four parent ions extracted from the mass spectra, 15 kinetic-energy resolved ionic fragments extracted from ion-velocity-map imaging, and five photoelectron channels obtained from electron-velocity-map imaging. From this analysis, we extracted 60 transient lifetimes corresponding to different photoinduced intermediates. This extensive dataset of transient species lifetimes of a prototypical polycyclic aromatic hydrocarbon provides new insights into the rich photochemistry of PAHs under the influence of intense electric fields.

Author Contributions
BM, JWLL, and MS conceived and designed the experiments. DG, PC, JWLL, SK, OA, FA, RB, AAB, BE, EG, SPG, LH, DH, EJ, MMK, JL, AKL, ZL, DL, SM, RM, JM, EM, TM, CCP, CP, JP, MP, DRA, PR, DROM, AS, ALS, AT, SHT, ATN, NV, ISV, ST, JK, AMR, DRO, MBT, SB, PEJ, CV, MBU, BM, and MS performed the experiments. DG, JWLL, PC, and SK analyzed the data. DST and PC performed theoretical simulations. DG, JWLL, DT, PC, BE, and MS performed detailed discussions of the results. DG, PC, DT, and MS wrote the manuscript with input from all authors.

Open data statement
Electronic supporting information (ESI) consists of 1) a PDF file with the fitting results and detailed complementary pictures to those presented in the manuscript, 2) a ZIP archive with the pre-processed experimental and theoretical data (including data analysis and plotting scripts) that can be used to reproduce the results and figures in the paper. All additional data is available by request from the corresponding author.

Conflicts of interest
There are no conflicts to declare.

Acknowledgements
This work was supported by the ERC Starting Grant ASTROROT, grant number 638027, and the project CALIPSOplus under grant agreement 730872 from the EU Framework Programme for Research and Innovation HORIZON 2020.

We acknowledge DESY (Hamburg, Germany), a member of the Helmholtz Association HGF, for providing experimental facilities. Parts of this research were carried out at FLASH. Beamtime was allocated for proposals F-20170540 and F-20211752. This research was also supported through the Maxwell computational resources operated at Deutsches Elektronen-Synchrotron DESY, Hamburg, Germany.

We acknowledge the Max Planck Society for funding the de-
velopment and the initial operation of the CAMP end-station within the Max Planck Advanced Study Group at CFEL and for providing this equipment for CAMP@FLASH. The installation of CAMP@FLASH was partially funded by the BMBF grants 05K10KT2, 05K13KT2, 05K16KT3, and 05K10KTB from FSP-302.

MBu and CV gratefully acknowledge support from the UK EPSRC EP/T021675/1, EP/V026690/1. MBu, EJ, ZL, and JM gratefully acknowledge support from the UK EPSRC (EP/S028617/1). EJ, ZL, and JM additionally thank Lincoln College, Oriel College, and University College, Oxford, for their respective travel funding.

The research was further supported by the federal cluster of excellence “Advanced Imaging of Matter” (AIM, EXC 2056, ID 390715994) of the Deutsche Forschungsgemeinschaft (DFG). L.H. acknowledges a fellowship within the framework of the Helmholtz-OCP postdoctoral exchange program.

PEJ, JP, JL, and MP acknowledge the support of the Swedish Research Council and the Knut and Alice Wallenberg Foundation.

SB acknowledges funding by the Helmholtz Initiative and Networking Fund.

DRol acknowledges the National Science Foundation grant PHYS-1753324.

References


