Orientation-dependent ionization yields from strong-field ionization of fixed-in-space linear and asymmetric top molecules

Jonas L Hansen¹, Lotte Holmegaard², Jens H Nielsen³, Henrik Stapelfeldt¹,², Darko Dimitrovski³,⁴ and Lars Bojer Madsen³,⁴

¹ Interdisciplinary Nanoscience Center (iNANO), Aarhus University, 8000 Aarhus C, Denmark
² Department of Chemistry, Aarhus University, 8000 Aarhus C, Denmark
³ Department of Physics and Astronomy, Aarhus University, 8000 Aarhus C, Denmark
⁴ Lundbeck Foundation Theoretical Center for Quantum System Research, Aarhus University, 8000 Aarhus C, Denmark

E-mail: henriks@chem.au.dk and bojer@phys.au.dk

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Abstract
The yield of strong-field ionization, by a linearly polarized probe pulse, is studied experimentally and theoretically as a function of the relative orientation between the laser field and the molecule. Experimentally, carbonyl sulphide (OCS), benzonitrile and naphthalene molecules are aligned in one or three dimensions before being singly ionized by a 30 fs laser pulse centred at 800 nm. Theoretically, we address the behaviour of these three molecules. We consider the degree of alignment and orientation and model the angular dependence of the total ionization yield by molecular tunnelling theory accounting for the Stark shift of the energy level of the ionizing orbital. For naphthalene and benzonitrile, the orientational dependence of the ionization yield agrees well with the calculated results, in particular, we observe that ionization is maximized when the probe laser is polarized along the most polarizable axis. For OCS the observation of the maximum ionization yield when the probe is perpendicular to the internuclear axis contrasts the theoretical results.

(Some figures in this article are in colour only in the electronic version)

1. Introduction
The ionization step leading to single ionization in the multiphoton or tunnel ionization regime is a fundamental process and a first step that triggers the subsequent dynamics in attoscience [1], both in the process of high-order harmonic generation and in streaking experiments [2]. This step is thought to be well understood for atoms; however, for larger molecules much less is known. Of particular importance is the understanding of the dependence of the initial ionization step on the molecular orientation with respect to the external field. Commonly used theories to explain strong-field phenomena extended to larger systems are the molecular tunnelling theory (MO-ADK) [3] and the strong-field approximation [4, 5]. Recent experiments, however, have shown disagreement with these theories [6], and the interpretation of the underlying physics has been and still is an area of debate [7–13]. To fully test existing theories and to guide the way for new theory development, experiments extended to larger and more complex molecular systems are needed.

Experimental orientation-dependent ionization rates have only been obtained for small molecular systems such as N2, O2, CO, CO2 and C2H2 [6, 14–16]. Many of these previous experiments have involved several steps when extracting orientation-dependent ionization rates from the measurements which complicates the interpretation. The direct measurement of the angular dependence of ionization was reported in [6], where the molecules were aligned
non-adiabatically, demonstrating the essential need for the creation of an ensemble of highly aligned molecules. An alternative to pre-aligning the molecules is to infer their spatial orientation after the ionization event. In favourable cases, this can be accomplished if the ionization process leads to the fragmentation of the molecule and the recoil directions of the ionic fragments uniquely define the molecular orientation at the time of ionization. Recently, this approach was used to determine the orientation-dependent ionization yields for HCl molecules ionized by intense circularly polarized (CP) infrared femtosecond laser pulses [17]. For larger molecular species, it is mostly impossible to access molecular frame information using fragmentation measurements. This fact is very important since it puts more than technical constraints, like low counting statistics, on the type of systems that can be investigated by recoil methods. In addition, the often encountered situation where ionization does not cause fragmentation also cannot give access to orientation-dependent measurements by recoil spectroscopy.

Hence, in general, molecular frame measurements require pre-alignment or pre-orientation of the molecules. The latter approach was employed recently in a series of joint experimental and theoretical studies on molecular frame photoelectron angular distributions (MFPADs) for three different adiabatically aligned and oriented molecules, carbonyl sulphide (OCS), benzonitrile (C₇H₅N, to be denoted by BN in the following) and naphthalene (C₁₀H₈, to be denoted by NPTH in the following) molecules, ionized by intense infrared CP femtosecond pulses [18–21]. The theoretical description of the MFPADs was accomplished by amending the existing theories of strong-field ionization to account for the Stark shift induced by the molecular dipole and polarizability [18, 20, 22]. Here, we employ linearly polarized (LP) pulses to perform measurements on the orientational dependence of the ionization yield on OCS, BN and NPTH molecules and use the same theories to describe the experimental results.

The paper is organized as follows. In the following section, we describe the experimental setup. Section 3, organized into subsections corresponding to the different molecular targets, contains a detailed comparison of the experiment and theory. Section 4 concludes the paper.

2. Experimental setup

The experimental setup was recently described in detail [19] so the discussion here is brief. A mixture of 90 bar helium and either 10 mbar OCS, 5 mbar BN or 5 mbar NPTH is expanded into vacuum through a pulsed valve. The resulting molecular beam is collimated by two skimmers before entering a 15 cm long electrostatic deflector. Traversing the deflector, polar molecules (OCS, BN) are dispersed along the gradient of the field according to their effective dipole moment which depends on the individual rotational quantum states of the molecules [23, 24]. This dispersion allows for the quantum state selection of the molecular ensemble in the interaction region downstream from the deflector where the molecular beam is crossed by one or more laser beams producing either adiabatically or non-adiabatically aligned targets. Alignment in the adiabatic regime is achieved using the output from an injection-seeded Nd:YAG laser (20 Hz, τFWHM = 10 ns, λ = 1064 nm), whereas field-free alignment, only implemented for OCS, is obtained using a part of the output from a 1 kHz amplified Ti:sapphire laser (800 nm) stretched to 0.5 ps and performing the experiment at the peak of the half-revival 40.5 ps after the non-adiabatic alignment pulse.

The degree of alignment is initially measured by Coulomb exploding the molecules using an intense femtosecond probe laser pulse (1 kHz, 30 fs, 800 nm) and detecting the axially recoiling S⁺ fragments from OCS and CN⁺ fragments from BN which is described elsewhere [18]. Unlike OCS and BN, NPTH does not contain good observables which allow for a precise measurement of the degree of alignment. Confinement of the molecular axes can, however, still be demonstrated through H⁺ ion imaging, providing information about the spatial direction of the molecular plane [21].

Using similar alignment laser parameters but inverting the voltage on the velocity-map-imaging (VMI) spectrometer, electrons rather than ions are focused onto the detector. In the experiment, the (major) polarization axis of the alignment pulse is kept fixed along the static-field axis of the VMI spectrometer, whereas the angle, Θ, of the LP probe pulse with respect to the static field is varied between 0° and 180°. Using this particular alignment geometry, the direction of the dipole moment for OCS and BN will be parallel to the static field breaking the head-for-tails symmetry of the ensemble to produce oriented molecules. The degree of orientation is estimated to ~80% for both OCS and BN. In the current experiment, the ionization yield only depends on the alignment and not on the orientation of the molecule since the probe pulse contains more than a few cycles and has an uncontrolled carrier-envelope phase. As such it would have been sufficient to record the yield only in the range 0°–90°. Our measurements in the 0°–180° interval, and the observation of identical behaviour of the yield in the 0°–90° and 90°–180° ranges, show precisely that only alignment and not orientation is important for the probe pulse used.

In all measurements, the intensity of the probe pulse has been adjusted to a regime where the molecules investigated primarily undergo single ionization without fragmentation. As a consequence, the electrons produced will be from the ionization step alone and it is, therefore, possible to obtain orientation-dependent ionization yields by integrating the measured electron signal at the various angles between the probe and alignment laser. To prevent fluctuations in the laser intensity from affecting the measurements, the total ionization yield has been normalized to that obtained from unaligned molecules in all of the experiments.

3. Experimental results and comparison with theory

3.1. Carbonyl sulphide (OCS)

In order to measure the angular dependence of the total ionization yield, a high degree of alignment is required. This is achieved by inducing adiabatic alignment of OCS molecules...
prepared in the lowest rotational quantum states [25]. Using a LP alignment pulse, polarized in the plane of the detector, with an intensity of $I_{\text{YAG}} = 8.4 \times 10^{11}$ W cm$^{-2}$ (10 ns, 1064 nm), a degree of alignment of $\langle \cos^2 \theta_{\text{ad}} \rangle = 0.90$ is obtained, with $\theta_{\text{ad}}$ being the angle between the YAG polarization and the molecular axis. Figure 1 displays the ionization yield, obtained by measuring the total electron signal, as a function of the angle between the most polarizable axis and the linear polarization, $\Theta$, for two different intensities of the probe pulse. Figure 1 shows that there is hardly any difference between the orientational dependence of high and low probe intensities, both showing a clear minimum of the signal when $\Theta = 0^\circ$, and a maximum in the ionization yield at $\Theta = 90^\circ$. The two degenerate highest occupied molecular orbitals (HOMOs) of OCS have nodal surfaces coinciding with the internuclear axis. For a highly aligned ensemble of OCS molecules, there will be no or little contribution from the orbital density of the HOMOs which can participate in the ionization process when the field is strictly parallel to the internuclear axis. Some suppression of the emission along this axis is therefore expected. However, such an argument does not explain why there is a maximum in the ionization yield at $\Theta = 90^\circ$.

A reasonable concern of the experiment is if the presence of the YAG laser field perturbs the ionization pathway and, in particular, the orientational dependence measured. To experimentally test this, the orientational dependence of the ionization yields was also measured from field-free-aligned OCS molecules. Using similar experimental conditions, the measurements were repeated, replacing the adiabatic pulse with a 0.5 ps long non-adiabatic alignment pulse of intensity $I_{\text{non-ad.}} = 8.3 \times 10^{12}$ W cm$^{-2}$ (800 nm). At the peak of the half-revival of OCS, occurring 40.5 ps after the alignment pulse, an alignment degree of $\langle \cos^2 \theta_{\text{ad}} \rangle = 0.84$ was obtained. The ionization yield was extracted from the total electron signal measured at an intensity of the probe beam corresponding to $I_{\text{probe,low}} = 1.5 \times 10^{14}$ W cm$^{-2}$ (30 fs, 800 nm).

Figure 2 presents a comparison of the orientation-dependent ionization yield of single ionization of field-free-aligned and adiabatically aligned OCS molecules. The ionization signal from field-free-aligned molecules exhibits an orientation dependence similar to that observed for adiabatically aligned molecules, i.e. a clear suppression at $\Theta = 0^\circ$ and a clear enhancement at $90^\circ$. The only discrepancy lies in the ratio ($R$) between the yield, $Y$, at $\Theta = 0^\circ$ and at $90^\circ$,

$$R = \frac{Y(\Theta = 0^\circ)}{Y(\Theta = 90^\circ)}. \quad (1)$$

For adiabatically aligned molecules $R = 0.65$ and for field-free-aligned molecules $R = 0.74$. We ascribe this difference to the fact that the degree of alignment is stronger in the adiabatic case.

The permanent dipole moment of OCS results in large Stark shifts due to the probe pulse that should be considered when calculating the ionization yields [18, 20]. Also, the alignment distribution of the molecules should be taken into account. We use a Gaussian approximation for the alignment distribution that can be used in cases of adiabatic alignment [26]. This alignment distribution is used for all molecules analysed here, with the only input being $\langle \cos^2 \theta_{\text{ad}} \rangle$. Calculations are performed using the Stark-shift-modified MO-ADK [3] and taking into account the HOMO, the dipole and the polarizability of OCS, see [22]. The ionization potential of the HOMO is $I_p = 11.4$ eV and for the probe intensity
in the experiment with the adiabatic alignment the Keldysh parameter \(^27\gamma \simeq 0.7\).

From figure 2, it is apparent that the theory cannot reproduce the maximum for the ionization yields perpendicular to the molecular axis. Instead, the ionization yields peak at approximately 45° with respect to the molecular axis. Compared to the conventional MO-ADK theory \(^3\), the Stark-shifted MO-ADK theory represents an improvement of around 10° when comparing to the experiment, see figure 2.

Previously, the ionization of OCS with CP fields have been studied \(^18, 20\), and it was found that ionization occurs efficiently when the field points along the permanent dipole moment, i.e. along the molecular axis. In this connection, we note two important differences between strong-field ionization induced by CP and LP fields.

First, there is a fundamental difference relating to recombination and rescattering of the ejected electron. In CP fields, the wave packet of the tunnelled electron does not revisit the parent ion, recombination and rescattering are minimized and the instantaneous ionization rate is mapped to the final momentum distributions. In LP fields, this one-to-one mapping is lost. Our results point to the fact that the total ionization yields by LP laser pulses should not be calculated solely based on ionization rates, although such calculations have reproduced the experimental results for small molecules such as O\(_2\) and N\(_2\) \(^6\).

Second, there is a difference relating to the influence of electronically excited states on the ionization process. For example in \(^28\) the comparison between theory and experiment on the strong-field ionization of laser-aligned CS\(_2\) indicated that excited states play a role. For an arbitrary orientation between the external field and the molecular axis, the molecular orbital is no longer an \(L_z\) eigenstate in the laboratory fixed frame. Circular polarization, however, still changes the \(\Lambda\) quantum number monotonously towards higher values. For linear polarization, lower \(\Lambda\) values will be preferred due to selection rules. The fact that within the excited state spectrum only states with relatively low \(\Lambda\) are within reach by few-photon absorption means that excited states are expected to play a larger role for linear polarization than for circular polarization. Along these lines, we also note that the discrepancy between the experiment and the theory, observed for OCS, is similar to recent studies on CO\(_2\) \(^6\). In the case of CO\(_2\), different theoretical methods and possible explanations for the discrepancy included the influence of excited states \(^7\), the influence of HOMO-1 and HOMO-2 \(^10\), and orbital modification and interference \(^12\). The debate is ongoing, see also \(^8, 9, 11, 13\). In the case of OCS, all candidates proposed in the theories for CO\(_2\) are candidates for OCS as well. For OCS, we note that by now experimental data for the orientation-dependent ionization yields exist for both LP probe pulses (present work) and for CP probe pulses \(^18, 20\). The two sets of results for OCS will stimulate further development of theory. In this connection, it is interesting that very recent time-dependent density-functional theory (TDDFT) calculations on OCS \(^29\), performed for LP laser pulses at approximately twice the intensity considered here, also do not agree with the present experimental trends.

In \(^29\) the calculated yield is smaller at \(\Theta = 90°\) than that at \(\Theta = 0°\) and is larger, again, than that at \(\Theta = 45°\) in agreement with the theoretical model predictions shown in figure 2, but in disagreement with the experimental findings.

### 3.2. Benzonitrile, BN

In the case of BN, we are able to record the orientation-dependent ionization yields from 1D- and 3D-oriented molecules. First, the orientational dependence of the ionization yield is studied for 1D adiabatically aligned molecules. As for OCS, the 1D alignment is induced by a LP YAG pulse resulting in the confinement of the C-CN axis (the most polarizable axis, see figure 3) along the YAG polarization with \(\langle\cos^2\theta_{2d}\rangle = 0.89, \theta_{2d}\) being the angle between the YAG polarization and the C-CN axis. The experimental results, obtained from the total electron signal as a function of \(\Theta\), are displayed in figure 3(a). As opposed to OCS, a clear maximum in the ionization yield is observed when the polarizations of the YAG and probe pulses are parallel (probe pulse polarized along the C-CN axis), and a clear minimum is observed when they are orthogonal.

Next, the orientational dependence of the ion yield is measured for 3D-aligned molecules. BN is an asymmetric top molecule and the LP YAG pulse leaves the molecule free to rotate about the arrested C-CN axis. If an elliptically polarized YAG pulse is employed (here with an ellipticity of 3:1, the value referring to the ratio of the intensities measured along the major and the minor polarization axis), then the C-CN axis remains restricted along the major polarization axis, while the molecular plane aligns with the polarization plane, i.e. the molecule is 3D aligned \(^19, 30, 31\). The experimental results are shown in figure 3(b). As for 1D-aligned molecules, a clear enhancement of the signal is observed at \(\Theta = 0°\), whereas the yield is almost constant when 60° < \(\Theta < 120°\).

Compared to OCS, BN is a larger molecule and has a larger dipole moment and larger polarizabilities, see \(^19\). The ionization potential of the HOMO is \(I_p = 9.79\) eV and at the intensity of the experiment the Keldysh parameter \(^27\gamma \simeq 0.82\). Using Stark-shifted MO-ADK theory \(^18–20, 22\), it is possible to explain the orientation-dependent ionization yields for LP pulses. This theory models the HOMO of BN with the simplest orbital that has a nodal plane in the molecular plane of BN, the 2p orbital (the \(x\)-axis points towards the reader in figure 3(b)) and includes the Stark shift of the ionization potential \(^19\). These results are then convoluted with the alignment distribution, assuming that the most polarizable axis is restricted to deviate from its ideal position only in the directions of the confined molecular plane. This procedure is followed for naphthalene as well. The agreement between the theory and experiment is very good.

### 3.3. Naphthalene, NPTH

Naphthalene is also an asymmetric top molecule, and so the orientational dependence of the ionization yield can be studied for both 1D and 3D adiabatically aligned molecules, using linearly and elliptically polarized YAG pulses, respectively.
Figure 3. Total ionization yield of (a) 1D and (b) 3D adiabatically aligned BN molecules (red squares, $\langle \cos^2 \theta_{2d} \rangle = 0.89$) as a function of the angle between the major polarization axis of the alignment and the probe pulse. The filled symbols represent the measurements and the dashed curves, to guide the eye, are Legendre polynomial fits. For the case of 3D-aligned molecules the theoretically calculated ionization yields (solid green/light grey curve) for confined BN molecules ($\langle \cos^2 \theta_{2d} \rangle = 0.89$) is included (see text). The intensities of the laser pulses used for the experiments and theoretical calculations are $I_{YAG,1D} = I_{YAG,3D} = 7.7 \times 10^{11}$ W cm$^{-2}$ (10 ns, 1064 nm) and $I_{probe} = 1.2 \times 10^{14}$ W cm$^{-2}$.

Similar to BN, NPTH has very large polarizabilities, but zero dipole moment. We note that the ionization of NPTH has previously been studied in [32, 33], where the experiment focused on the role of dissociative ionization channels and their signatures in the energy distributions, using longer laser pulses. Here we focus on the orientation-dependent ionization yields.

For NPTH, the only experimental information about the molecular alignment stems from H$^+$ ions created in the Coulomb explosion. In the presence of the YAG pulse, LP perpendicular to the detector plane, the H$^+$ ion image has the shape of a doughnut. This shows that the long axis of the molecule is confined along the YAG polarization and that the molecular plane is randomly oriented around this axis [34]. This is in accord with the expectations since the long molecular axis possesses the largest polarizability and should, therefore, be the one that is confined along the YAG polarization. The H$^+$ ion images, do, however, not immediately provide quantitative information about the degree of alignment. Due to the high polarizability (anisotropy) of NPTH, the alignment is expected to be high [21] and of the same order as that for BN. Nevertheless, we use a conservative estimate for the degree of alignment, $\langle \cos^2 \theta_{2d} \rangle = 0.85$, in the theoretical calculations. The experimental results of the ionization yield, obtained from the total electron signal as a function of $\Theta$, is displayed in figure 4(a).

If the YAG polarization is changed from linear to elliptical (again with an ellipticity ratio of 3:1), 3D alignment is induced with the long molecular axis along the major component of the elliptically polarized field and the short molecular axis (still in the molecular plane) along the minor axis. The experimental results are shown in figure 4(b). Comparing figures 4(a) and (b), it can be seen that the measured ionization yields for 1D- and 3D-aligned molecules show similar features. In particular, the ionization yield peaks when the polarizations of the alignment and probe pulses are parallel, and reaches a minimum when they are orthogonal. For 1D-aligned molecules, the modulation ratio, $R = 2.1$, is slightly larger than for 3D-aligned molecules, $R = 2.0$.

To confirm that the angular dependence of the experimental ionization yields displayed in figure 4 indeed results from an orientation dependence rather than from some combined effects of the two laser fields, we repeated the measurements for identical laser intensities but for a backing pressure of the He carrier gas in the supersonic expansion of only 15 bar. When the He pressure is lowered from 90 to 15 bar the rotational temperature increases and the degree of alignment deteriorates significantly [35]. We observed that the pronounced angular dependence of the ion yield, apparent in figure 4 essentially vanishes. For the measurements on 3D-aligned molecules $R$ dropped from 2.0 to 1.3, which is close to the ratio observed for purely collisionally aligned molecules [36, 37], that is, with only the probe pulse present, $R = 1.1$.

We have performed quantum chemistry calculations [38] of NPTH at the Hartree–Fock level of theory using a TZV basis, including a few diffuse orbitals. We obtained the following ionization potentials of interest. $I_p$(HOMO) = 8.05 eV, $I_p$(HOMO-1) = 8.84 eV. The other orbitals are more...
Figure 4. Total ionization yield of (a) 1D and (b) 3D adiabatically aligned NPTH molecules (red squares) as a function of the angle between the major polarization axis of the alignment and the probe pulse. The filled symbols represent the measurements and the dashed curves, to guide the eye, are Legendre polynomial fits. For the case of 1D-aligned molecules, the theoretically calculated ionization yields have been modelled using the HOMO (solid orange/light grey line), the HOMO-1 (solid cyan/black line) and the HOMO including Stark shifts (solid dark yellow/dark grey line) and the true HOMO (magenta/dark grey line). All calculations assume $\langle \cos^2 \theta \rangle = 0.85$. The intensities of the laser pulses used for the experiments and theoretical calculations are $I_{\text{YAG},1D} = I_{\text{YAG},3D} = 7.7 \times 10^{11}$ W cm$^{-2}$ (10 ns, 1064 nm) and $I_{\text{probe}} = 8.2 \times 10^{13}$ W cm$^{-2}$.

tightly bound, for example, $I_p($HOMO-2$) = 10.61$ eV. The HOMO of NPTH possesses three nodal planes, one coincides with the molecular plane and the other two are orthogonal to each other and also orthogonal to the molecular plane. The HOMO-1 also has a nodal plane in the molecular plane. The small energy difference between the HOMO and the HOMO-1 means that they are both expected to contribute to the ionization signal calculated by MO-ADK theory at experimental Keldysh parameters [27] of a typical value of $\gamma \simeq 0.9$. From figure 4 we see, as in the case of BN, that the theory reproduces the overall trend of the orientation-dependent ionization yields. Specifically, in figure 4(a) we consider the case of 1D-aligned molecules. The theoretical calculations corresponding to this case were obtained by averaging over the set of results obtained for different angles of orbital rotation around the molecular axis (angles with steps of $10^\circ$ are considered), and then convoluting with the alignment distribution. Because of the small difference of the ionization potential of HOMO and HOMO-1 for NPTH, and due to the symmetry of the HOMO-1, the angle-dependent ionization rates from both HOMO and HOMO-1 are comparable if not equal, see figure 4(a). Also, because of the large polarizabilities of NPTH and its cation, we have included the Stark shift of the ionization potential. Including the Stark shift does not change much in the orientation-dependent ionization yields, and on the scale of the figure it is difficult to distinguish the three theory curves. All three curves reproduce the overall trend in the experiment, but not the correct ratio $R$ between the experimental ionization yield along and perpendicular to the molecular axis. As is clear from the figure, the theory predicts a larger $R$ than the experiment. This may be due to the influence from electronically excited states in the ionization process or possibly due to the distortion of the HOMO and HOMO-1 by the strong ionizing field. The same discrepancy between theory and experiment persists for 3D-oriented targets as well. In figure 4(b), for the 3D-oriented target, we show the theoretical curves from Stark-shifted MO-ADK theory for a HOMO modelled by a $f_{xyz}$ orbital as in [21], and by taking the true HOMO of NPTH from our quantum chemistry calculation. In both cases, the Stark shifts are included. The curve corresponding to the true HOMO of NPTH is very similar to the theoretical curves for the 1D-oriented target. On the other hand, in the calculation using the model $f_{xyz}$ orbital, two nodal planes are visible in the ionization yield. Interestingly, this curve seems to fit very well with the experiment for ionization perpendicular to the molecular axis, pointing to a possible orbital modification by the field, especially for the geometry with the perpendicular orientation of the molecular axis and the field.
4. Conclusion

We have presented a set of experimental data for the orientation-dependent strong-field ionization yields for aligned OCS, BN and NPTH (one linear and two asymmetric top molecules) using LP probe pulses. The orientation-dependent yields were compared with simple theoretical models. For OCS, the maximum (minimum) ionization yield occurs when the polarization of the ionizing field is perpendicular (parallel) to the internuclear axis. Our theory, as well as numerical TDDFT approaches [29], predicts a maximum elsewhere—in our case at an angle of 45° between the OCS axis and the polarization of field. We discussed the possibility of the discrepancy being caused by the influence of electronically excited states in the multiphoton ionization process. It is an open question whether this is correct (indicating a shortcoming of the TDDFT to account for excited states), or whether the discrepancy is due to some other dynamics not accounted for in approaches used so far.

For BN and NPTH, the ionization yield is maximized (minimized) when the ionizing field is parallel (perpendicular) to the molecular axis with the largest polarizability. In both cases, the present modelling is accounted for that behaviour. In the case of BN, a very good agreement between experimental data and theory was obtained. In the case of NPTH, the theory overestimates the ratio between the ionization yields parallel and perpendicular to the most polarizable axis.

Our results suggest that the angular behaviour of the total ionization yields in LP fields cannot, in general, be described by ionization rates only. In this sense, our analysis suggests that to map out the instantaneous ionization rate of the molecule, it is better to use CP laser pulses because both rescattering with the parent ion and the influence of excited molecular states are minimized.

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