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Introduction

Recent developments of X-ray free-electron lasers (XFELs) delivering extremely short X-ray pulses of a few femtoseconds¹ and of a mega-electron-volt pulsed electron beam² at SLAC have paved new pathways to image structural changes of molecules in chemical reactions, for example, employing time-resolved X-ray diffraction³ and ultrafast electron diffraction.⁴ While

^a Institut f
ür Kernphysik, Goethe-Universit
ät, Max-von-Laue-Stra
ße 1, 60438 Frankfurt am Main, Germany

b CDL i UL i CTL

- ^c Institut für Physik und CINSaT, Universität Kassel, Heinrich-Plett-Straße 40, 34132 Kassel, Germany. E-mail: demekhin@physik.uni-kassel.de
- ^d Institute of Multidisciplinary Research for Advanced Materials, Tohoku University, Sendai 980-8577, Japan
- ^e Deutsches Elektronen-Synchrotron (DESY), Notkestraße 85, 22607 Hamburg, Germany
- ^f Molecular Physics, Fritz-Haber-Institut der Max-Planck-Gesellschaft,
- Faradayweg 4-6, 14195 Berlin, Germany. E-mail: trinter@fhi-berlin.mpg.de
- ^g Sorbonne Université CNRS, Laboratoire de Chimie Physique-Matiere et Rayonnement, LCPMR, F-75005, Paris, France
- ^h Department of Physics, University of Nevada, Reno, Nevada 89557, USA
- ⁱ RIKEN Center for Advanced Photonics, RIKEN, 2-1 Hirosawa, Wako, Saitama, 351-0198, Japan
- ^j Department of Chemistry, Tohoku University, 6-3 Aramaki Aza-Aoba, Aoba-ku, Sendai 980-8578, Japan
- ^k European XFEL, Holzkoppel 4, 22869 Schenefeld, Germany.

E-mail: till.jahnke@xfel.eu

High-energy molecular-frame photoelectron angular distributions: a molecular bond-length ruler

I. Vela-Peréz,^a F. Ota, ^b A. Mhamdi,^c Y. Tamura, ^b J. Rist,^a N. Melzer, ^b ^a S. Uerken,^a G. Nalin, ^b ^a N. Anders,^a D. You, ^b ^d M. Kircher, ^b ^a C. Janke,^a M. Waitz,^a F. Trinter, ^b *^{ef} R. Guillemin, ^b ^g M. N. Piancastelli, ^b ^g M. Simon, ^b ^g V. T. Davis, ^b ^h J. B. Williams,^h R. Dörner,^a K. Hatada, ^b ^b K. Yamazaki,ⁱ K. Fehre,^a Ph. V. Demekhin, ^b *^c K. Ueda,^{dj} M. S. Schöffler ^b ^a and T. Jahnke ^b *^k

We present a study on molecular-frame photoelectron angular distributions (MFPADs) of small molecules using circularly polarized synchrotron light. We find that the main forward-scattering peaks of the MFPADs are slightly tilted with respect to the molecular axis. This tilt angle is directly connected to the molecular bond length by a simple, universal formula. We apply the derived formula to several examples of MFPADs of C 1s and O 1s photoelectrons of CO, which have been measured experimentally or obtained by means of *ab initio* modeling. In addition, we discuss the influence of the back-scattering contribution that is superimposed over the analyzed forward-scattering peak in the case of homonuclear diatomic molecules such as N₂.

catching the motion of the individual atoms, including hydrogen atoms, of single molecules during chemical reactions is of fundamental interest, it still remains a challenge.

A couple of decades ago, it was pointed out that core-level photoelectron angular distributions include molecular structure information, which can be accessed if the molecule is either fixed in space or if the photoelectron angular distribution in the molecular frame (MFPAD) is retrieved from a coincidence measurement.⁵ Accordingly, measuring the MFPAD of an isolated molecule is the gas-phase analog of photoelectron diffraction imaging (PED) routinely used for studying the surface structure.⁶ Following the spirit of PED, a couple of groups^{7,8} proposed time-resolved MFPAD measurements using an XFEL light pulse as an ionizing source, as one of the future routes to image the motion of individual atoms in a molecule. Several attempts towards time-resolved MFPAD measurements with XFELs were reported, in which the sample molecules were aligned by employing an impulsive or adiabatic alignment method using optical laser fields.9-12 However, no timeresolved studies of molecular structural changes have been reported, so far, partly due to the complexity of the needed pulse sequence: a first laser pulse is needed to align the molecule, a second laser pulse triggers the photoreaction, and a final XFEL pulse ejects the photoelectron at a particular site in the molecule in order to perform PED.

The first high-repetition-rate XFEL, the European XFEL,¹³ opened the door to coincidence experiments using COLTRIMS



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^b Department of Physics, University of Toyama, Toyama 930-8555, Gofuku 3190, Japan

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(Cold Target Recoil Ion Momentum Spectroscopy) reaction microscopes (REMI)¹⁴ for MFPAD measurements. In a COL-TRIMS measurement, the spatial orientation of a molecule can be deduced after triggering a fragmentation of the molecule and detecting the momenta of the fragment ions in coincidence. As the momentum of the ejected photoelectron is recorded in coincidence with the fragment ions, the photoelectron's emission direction with respect to the molecule can be determined. Very recently, Kastirke *et al.* have reported a first successful implementation of this technique at the soft X-ray beamline of the European XFEL.^{15–17}

To establish MFPAD measurements as a routinely usable tool for molecular structure imaging, however, an important ingredient is still missing, which is a method of extracting the molecular structure directly from the measured MFPAD without the need for a full theoretical modeling of the photoemission process. So far, concepts proposed for achieving this goal focused on polarization-averaged MFPADs (see, e.g., ref. 18 and references therein), i.e., on MFPADs, which have been obtained after integrating over all laboratory-frame orientations of the molecule. The concept relies on two prerequisites: firstly, at sufficiently high photoelectron kinetic energies, for which a single-scattering approximation is valid (say, above 70 eV), forward-scattering peaks observable in polarization-averaged MFPADs coincide with the relative location (from the point of view of the emitter atom) of neighboring atoms.^{19,20} Secondly, the molecular-frame interference pattern caused by the direct and scattered photoelectron waves can be correlated with the distance between the emitter and the scatterers.^{21,22} In addition, in the case of homo-nuclear molecules, bond-length information can be obtained exploiting the analogy between Young's double-slit principle and MFPADs (see, e.g., ref. 23).

In the present work, we demonstrate a different approach building on the idea of stereo-atomscope diffraction spectroscopy known in the field of surface-structure determination, in which circularly polarized light is employed as an ionizing source for PED.^{24,25} If the emitter of a photoelectron and a neighboring atomic scatterer are located within the polarization plane of the ionizing light, the forward-scattering peak belonging to that atomic neighbor shows a distinct angular tilt. Thus, employing the hetero-nuclear diatomic molecule CO as a prototype system, we show that high-energy MFPADs of fixed-in-space molecules provide direct access to structural features such as the molecule's bond length, if circularly polarized light is used for the ionization and the MFPAD is confined to the polarization plane. Such MFPADs are hereafter referred to as CP-MFPADs. Earlier works²⁶⁻²⁸ on CP-MFPADs showed in principle such an angular tilt as well. However, these studies covered in their discussion other properties of the measured emission distributions and in many cases were performed at low electron kinetic energies. At such energies, the tilt angle is sometimes obscured by higherorder multiple-scattering effects and, accordingly, we focus in the present study on high-energy electrons.

While this paper demonstrates this concept for a simple diatomic molecule, it should be possible to extend it in future work to larger molecules as well. Whenever an atom of the molecule is located together with the emitter atom within the polarization plane, its corresponding forward-scattering peak will show a distance-dependent tilt. As detailed below, COL-TRIMS reaction microscopes are particularly suited for such studies, as they enable the experimenters to determine the laboratoryframe molecular orientation in addition to the molecular-frame angular emission distribution of the photoelectron.

The remaining part of the manuscript is structured as follows: first, we present the experimental C 1s and O 1s CP-MFPADs of CO. We find that the high-energy CP-MFPAD exhibits petal-like features (*i.e.*, a flower shape) with a strong forward-scattering peak in the direction from the emitter to the scatterer. The latter peak is slightly tilted upward or downward, depending on the handedness of the circularly polarized photons. We then show that our ab initio calculations reproduce very well the experimental CP-MFPADs. Finally, we demonstrate that the features of high-energy CP-MFPADs described above can be well captured by a very simple analytic expression, and that the bond length can be directly read off from the tilt angle of the forward-scattering peak without any need for ab initio modeling. For comparison and further insight, we present also the results for the homo-nuclear diatomic molecule N2. It turns out that great care needs to be taken when measuring the MFPADs. As we will show, the resulting error bars for the extraction of the bond length from our routinely measured MFPADs are surprisingly large in some cases and, for example, measuring MFPADs for both light helicities can greatly help to minimize systematic errors.

Experiment

The experiment was performed at Synchrotron SOLEIL (Saint-Aubin, France) using the well-established COLTRIMS technique¹⁴ during the 8-bunch mode (pulsed operation) of the synchrotron. A supersonic expansion of CO or N_2 gas was skimmed to form a molecular beam that crossed the synchrotron radiation provided by the variable-polarization undulator-based beamline SEX-TANTS. Electrons and ions generated by photoionization and subsequent Auger decay were accelerated (in opposite directions) using a static electric field, onto two position- and time-sensitive microchannel-plate detectors (MCPs) using a multi-hit-capable delay-line position readout.²⁹ From the position of the impact on the detectors, the known distance between the ionization region and the MCPs, and the time-of-flight, the particle trajectories inside the spectrometer can be deduced, and from these, the initial momentum vector of each particle can be determined.

In more detail, the whole COLTRIMS analyzer consisted of only a single acceleration region (spanning the full region between the two detectors) using a homogeneous electric field with a strength of $E = 38.3 \text{ V cm}^{-1}$. The ion arm had a length of 20.6 cm and the ion detector had a diameter of 125 mm. This configuration allowed for the detection of the breakup of the molecules into singly charged ions for a kinetic energy release of up to 30 eV with a full 4π coverage, *i.e.*, independent of the emission direction of the ions in the laboratory frame. Detecting high-energy electrons requires typically the use of either very

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high electric extraction fields or very high magnetic confinement fields for the electrons. While focusing on the measurement of high-energy electrons, we can exploit the fact that we are mainly interested in their emission distributions within the polarization plane of the ionizing photons (and more specifically in the MFPADs of molecules oriented in parallel to the polarization plane). To achieve this goal, it is therefore not necessary to measure the electrons with full solid-angle coverage, *i.e.*, it is sufficient to detect electrons, which are emitted perpendicularly to the light propagation direction. All possible molecular-frame electron emission angles are then still covered in this measurement as the ions are detected with 4π solid angle (and due to the symmetry features of the circularly polarized light, all molecular orientations within the polarization plane are equivalent). The electron arm had a length of 46.4 cm. The active area of the electron detector had a diameter of 125 mm. With these spectrometer properties and an additional superimposed homogeneous magnetic field with a strength of B = 10 G, we were able to measure electrons of, *e.g.*, $E_{kin} = 633$ eV (which corresponds to C 1s electrons of CO at $h\nu$ = 930 eV) if they are emitted within an angle of 20° relative to the symmetry axis of the COLTRIMS spectrometer. For each ionization event, a core-level photoelectron was detected in coincidence with the two fragment ions resulting from the Coulomb explosion of the molecule after Auger decay. The detection rates throughout our experiment were in the range of 11 kHz on the electron detector and 4.6 kHz on the ion detector. Such rates have been found suitable with respect to coincidence conditions and measurement background in the past. The exact values for the degree of circular polarization of the beamline are not available. Previous studies performed at the same beamline³⁰ suggest, however, that it is very close to 100%.

Details of the calculations

For our *ab initio* modeling, we employed the following approach: the angular distributions of the C 1s and O 1s photoelectrons of the CO molecule and of the N 1s photoelectrons of the N₂ molecule were computed by using the single-center (SC) method and code,³¹ which provides an accurate description of the partial photoelectron continuum waves in molecules. The calculations were carried out at the equilibrium internuclear geometries of the ground electronic states of CO and N₂ in the frozen-core Hartree–Fock approximation. The SC expansions of the occupied orbitals were restricted by partial harmonics with $\ell_c \leq 99$ and for photoelectrons with $\ell_{\varepsilon} \leq 49$.

Results and discussion

We performed measurements using right circularly and left circularly polarized light at the photon energies of $h\nu = 690$ eV and $h\nu = 930$ eV for the studies on CO, and $h\nu = 880$ eV and $h\nu =$ 1190 eV for the investigation on N₂. The resulting photoelectron kinetic energies are 393 eV and 633 eV for C 1s photoemission of CO, 155 eV and 395 eV for O 1s photoemission of CO, and 470 eV and 780 eV for N 1s photoemission of N₂. Fig. 1 depicts



Fig. 1 Comparison between the measured [red dots with error bars (statistical errors)] and calculated (blue solid lines) CP-MFPADs for different photoelectron kinetic energies. (a) CO O 1s at 155 eV, (b) CO O 1s at 395 eV, (c) CO C 1s at 393 eV, (d) CO C 1s at 633 eV, (e) N₂ N 1s at 470 eV, and (f) N₂ N 1s at 780 eV. The propagation direction of the ionizing light is pointing out of the paper plane, the circular arrow in the middle of each panel indicates the sense of the rotation of the electric field, and the handedness of the circular polarization σ is indicated in the lower right corner of each panel. The experimental data set is restricted to cases, where the molecular orientation and the electron emission are in the polarization plane within ±15°.

the respective experimental CP-MFPADs and those obtained from our *ab initio* calculations. The experimental data are confined to cases, where the electron emission occurs within the polarization plane within an opening angle of $\pm 15^{\circ}$, and for cases, in which the molecular orientation was parallel to the polarization plane within the same angular range. It should be noted that the tilt-angle effect discussed in the following is not drastically affected by these opening angles, as it scales linearly with that angle. Our modeling nicely reproduces all features observed in the experiment. Looking at the CO results in Fig. 1(a)–(d), we find that the number of petals (resembling a flower shape), which appear as a result of interference between the direct and scattered photoelectron waves, increases as photoelectron energy increases. The same behavior has been seen in studies on polarization-averaged MFPADs.²⁰

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The most striking feature of all the CP-MFPADs is a very strong forward-scattering peak, which appears in the direction of the scatterer atom that is neighboring the emitter atom: the forward scattering is not located exactly along the molecular axis, but occurs slightly tilted from it in the direction of rotation of the electric-field vector. This upward or downward tilt thus flips if the handedness of the ionizing light is changed from the right ($\sigma = -1$) to the left ($\sigma = +1$). This tilt is a characteristic feature of the CP-MFPADs, as also of PED on surfaces.^{24,25} Furthermore, close inspections reveal that the tilt angle decreases with the increase of the photoelectron energy. The results of N₂ in Fig. 1(e) and (f) depict these features as well.

In order to gather more detailed information on the findings described above, we performed further *ab initio* calculations over a wide range of photoelectron energies from 70 eV up to 1000 eV. The results are summarized in Fig. 2. Fig. 2(a), (c), and (e) illustrate that, indeed, the number of petals in the flower shape of the CP-MFPAD increases while the tilt angle of the forward-scattering peak decreases with the increase of the photoelectron kinetic energy. Fig. 2(b), (d), and (f) depict the extracted tilt angle θ_{tilt} as a function of photoelectron energy. For the CO molecule [Fig. 2(b) and (d)], we observe monotonic decreases of the tilt angle with the increase of the photoelectron energy. The results for N₂ depicted in Fig. 1(f) show that the decrease in the tilt angle is additionally modulated by a damped oscillation.

Aiming at elucidating these characteristic behaviors of the measured and calculated CP-MFPADs, we derive analytic expressions for describing the CP-MFPAD of hetero-nuclear diatomic molecules AB in the high-energy regime. We choose atom A as the photo-absorbing atom that emits a photoelectron. We define the bond-length vector $\mathbf{R} = \mathbf{r}_{\rm B} - \mathbf{r}_{\rm A}$ (along the *x* axis), the photoelectron wave vector $\hat{\mathbf{k}}$, and angle θ between them. We assume that CP-MFPADs are measured in the polarization plane of the circular light (*x*-*y* plane) and employ the electric-dipole, single-channel, and single-scattering approximations using a site *T*-matrix expansion.^{32,33} According to the approach of ref. 18 and 22, the CP-MFPAD is given by the following superposition:

$$I^{\mathbf{A}}(\mathbf{k},\,\hat{\varepsilon}_{\sigma}) \propto |B_{x}^{\mathbf{A}}(\mathbf{k}) + i\sigma B_{y}^{\mathbf{A}}(\mathbf{k})|^{2}, \qquad (1)$$

where $B_x^A(\mathbf{k})$ and $B_y^A(\mathbf{k})$ are the photoionization amplitudes excited by the *x* and *y* linear components of the circularly polarized light, respectively, and polarization index $\sigma = -1$ and +1 corresponds to light with negative and positive helicity, respectively. Introducing the scattering amplitude of atom B *via* $f(k,\theta) = |f(k,\theta)|e^{i\phi(k,\theta)}$ with a phase function $\phi(k,\theta)$, the respective photoionization amplitudes can be written in the singlescattering plane-wave approximation as^{18,22}

$$B_{x}^{A}(\mathbf{k}) = -it_{1}^{A}\sqrt{\frac{k}{\pi}}\sqrt{\frac{3}{4\pi}} \times \left[e^{i\frac{1}{2}kR\cos\theta}\cos\theta + \frac{|f(k,\theta)|}{R}e^{i\phi(k,\theta)}e^{ikR\left(1-\frac{1}{2}\cos\theta\right)}\right],$$
(2)



Fig. 2 CP-MFPADs resulting from our *ab initio* calculations for (a) CO O 1s, (c) CO C 1s, and (e) N₂ N 1s as a function of photoelectron kinetic energy in a range from 70 eV to 1000 eV. The tilt angles of the forward peak, $\theta_{\rm tilt}$, of the CP-MFPADs are displayed as functions of the photoelectron kinetic energy for (b) CO O 1s, (d) CO C 1s, and (f) N₂ N 1s, respectively. The color encodes the photoelectron energy in all panels.

$$B_{y}^{A}(\mathbf{k}) = -it_{1}^{A}\sqrt{\frac{k}{\pi}}\sqrt{\frac{3}{4\pi}} \Big[e^{i\frac{1}{2}kR\cos\theta}\sin\theta} \Big], \qquad (3)$$

where t_1^A is a scattering *t*-matrix of the atom A.¹⁸ The first terms in both amplitudes correspond to the direct waves emitted from the absorbing atom A, and the second term of eqn (2) is the single-scattered wave by the neighboring atom B. In the plane-wave approximation, such a contribution of the scattered wave to the amplitude for the *y* component of the circularly polarized light (3) can be neglected, since the directly emitted wave ($\sim \sin \theta$) does not propagate towards the neighboring atom B at $\theta = 0$. Substituting eqn (2) and (3) in eqn (1) and performing necessary simplifications, we arrive at the following compact expression for the CP-MFPAD as a function of the photoelectron emission vector **k**:

$$I^{\mathcal{A}}(\mathbf{k},\hat{\varepsilon}_{\sigma}) \propto \left|t_{1}^{\mathcal{A}}\right|^{2} \left(\frac{k}{\pi}\right) \frac{3}{4\pi} \left\{1 + \frac{\left|f(k,\theta)\right|^{2}}{R^{2}} + 2\frac{\left|f(k,\theta)\right|}{R}G_{\sigma}(k,\theta)\right\}, \quad (4)$$

with the fringe function, $G_{\sigma}(k,\theta)$, defined as

$$G_{\sigma}(k,\theta) = \cos[kR(1-\cos\theta) + \phi(k,\theta) - \sigma\theta].$$
 (5)

The first and the second terms of eqn (4) are the direct wave from the emitter A and the scattering wave from the scatterer B, respectively. The third term represents the interference between the direct and single-scattered waves and depends on the bond length.

Constructive and destructive interference in the differential photoemission intensity (*i.e.*, the fringes in the CP-MFPAD) correspond to the maxima and minima of the fringe function (5). In order to find the angles, at which the maxima and minima appear in $G_{\sigma}(k,\theta)$, we set $dG_{\sigma}(k,\theta)/d\theta = 0$:

$$\frac{\mathrm{d}G_{\sigma}(k,\theta)}{\mathrm{d}\theta} = -\sin[kR(1-\cos\theta)+\phi(k,\theta)-\sigma\theta] \\ \times \left[kR\sin\theta+\frac{\mathrm{d}\phi(k,\theta)}{\mathrm{d}\theta}-\sigma\right] = 0.$$
(6)

This condition is satisfied either when the sine function becomes zero, *i.e.*, $\sin[kR(1 - \cos \theta) + \phi(k,\theta) - \sigma\theta] = 0$ or when the second term in braces vanishes, *i.e.*, $kR \sin \theta + d\phi(k,\theta)/d\theta - \sigma = 0$. The first condition describes the angles, at which maxima and minima form the flower shape in the CP-MFPAD shown in Fig. 1(a)–(d) and Fig. 2(a), (b). The second condition yields a relationship between kR and the tilt angle θ_{tilt} . This relation confirms that the tilt angle decreases with the increase of the photoelectron momentum k (and thus, of the photoelectron kinetic energy), as depicted in Fig. 2(b) and (d) for CO. If $|f(k,\theta)|_{\theta=0,\pi} \neq 0$, as in this study, $d\phi(k,\theta)/d\theta|_{\theta=0,\pi} = 0$ holds.³⁴ We can thus assume that $d\phi(k,\theta)/d\theta$ is negligible for small tilt angles from the forward and backward directions. From the solution of the second part of eqn (6), we finally obtain the following relationship for the case of hetero-nuclear molecules:

$$R_{\text{Hetero}}^{\text{tilt}} = \frac{\sigma}{k \sin(\theta_{\text{tilt}})} = \frac{1}{k \sin(|\theta_{\text{tilt}}|)}.$$
 (7)

Eqn (7) connects directly the bond length of a molecule and the tilt angle $|\theta_{tilt}|$ of the forward-scattering peak of the CP-MFPAD, and thus can be thought of as a simple, universal bond-length ruler. As pointed out in an insightful argument by Daimon and coworkers, 35 the physics underlying eqn (7) can be understood very intuitively. Upon birth the photoelectron carries one unit of angular momentum, which it inherited from the photon. Classically, an electron moving with a momentum kalong a straight line thus must follow a path of impact parameter \vec{b} with respect to the nucleus such that $\vec{k} \cdot \vec{b} = 1\hbar$. This straight line defines the tilted symmetry axis of the photoemission. For $R \gg b$ this results in $R = 1/k \sin(\theta)$ and a symmetry axis of the problem, which is tilted by θ with respect to the molecular axis. This simplified classical explanation suggests that the tilt angle should scale linearly with the ellipticity of the ionizing light. Fig. 3 depicts the bond length R_{Hetero} estimated from the computed and measured CP-MFPADs using eqn (7). The resulting values are plotted as functions of kR_{eq} , where $R_{eq} = 1.1283$ Å and 1.0977 Å for CO and N_2 , respectively. Note that eqn (7) is derived for hetero-nuclear diatomic molecules, and its application to the homo-nuclear N₂ molecule is a rather crude approximation, as we will discuss below. From Fig. 3(a), it is evident that the bond lengths,



Fig. 3 Bond lengths (refer to the left-handed vertical axis), estimated from the tilt angles of the *ab initio* computed (solid lines with small symbols) and experimentally measured (large symbols with error bars) CP-MFPADs *via* the derived relation (7) as functions of kR_{eq} , and the respective relative errors (broken lines, refer to the right-handed vertical axis). The equilibrium bond lengths $R_{eq}^{eq} = 1.1283$ Å and $R_{eq}^{N_2} = 1.0977$ Å are indicated by the horizontal dashed lines.

estimated from both the *ab initio* modeled and measured CP-MFPADs, agree well with the equilibrium bond length of the neutral ground state of CO (the horizontal dashed line), with a deviation of less than 10%. Please note that the counter-intuitive increase of the relative error with respect to the *ab initio* modeling for smaller kR_{eq} is due to the fact that at lower electron energies higher-order multiple-scattering effects provide a non-negligible contribution to the transition amplitude. On the other hand, the bond lengths of N₂ in Fig. 3(b), estimated from the *ab initio* CP-MFPADs, oscillate as a function of kR_{eq} around R_{eq} . Those oscillations are damped and drop below 10% for sufficiently high photoelectron energies. The bond length of N₂ estimated from the experimental CP-MFPADs is, however, in good agreement with the expected value of R_{eq} within the experimental uncertainties.

The large oscillation in $R_{\rm Hetero}^{\rm tilt}$ in Fig. 3(b) can be interpreted as follows. In the experiment, one is unable to distinguish photoelectrons originating from the N 1s gerade and ungerade molecular orbitals of N₂, which are separated by ~ 105 meV³⁶ (or from the degenerate 1s orbitals of the left and right nitrogen atoms). The resultant CP-MFPADs are therefore the sum of the two contributions:

$$\bar{I}(\mathbf{k},\hat{\varepsilon}_{\sigma}) \propto I_{g}(\mathbf{k},\hat{\varepsilon}_{\sigma}) + I_{u}(\mathbf{k},\,\hat{\varepsilon}_{\sigma}) = I^{A}(\mathbf{k},\hat{\varepsilon}_{\sigma}) + I^{A}(-\mathbf{k},\,\hat{\varepsilon}_{\sigma}).$$
(8)

Here, we used the fact that $f^{A}(k,\theta) = f^{B}(k,\pi - \theta)$ for homonuclear diatomic molecules. One can see that eqn (8) has a C_{2} point symmetry, as is evident from Fig. 2(e). The fringe function of eqn (8) can thus be reduced to a superposition of fringe functions of two emitting nitrogen atoms

$$H_{\sigma}(k,\theta) \propto G_{\sigma}(k,\theta) + G_{\sigma}(k,\pi-\theta),$$
 (9)

with $G_{\sigma}(k,\theta)$ given by eqn (5). We see that it has also C_2 symmetry, namely $H_{\sigma}(k,\theta) = H_{\sigma}(k,\pi - \theta)$. Because of the superposition of $G_{\sigma}(k,\theta)$ and $G_{\sigma}(k,\pi - \theta)$, the forward-tilted peak of $G_{\sigma}(k,\theta)$ overlaps with the backward-scattering peak of $G_{\sigma}(k,\pi - \theta)$. Since the backward scattering exhibits EXAFS-type oscillations, the tilt angle of the homo-nuclear diatomic molecules oscillates with *k*. These oscillations are damped with the energy as $\sim k^{-2}$, which corresponds to the asymptotic behavior of the backward-scattering amplitude.³⁷

In particular, for high electron energies, θ_{tilt} becomes small and, accordingly, its extraction from the experimental data becomes very challenging. Already small systematic errors can prevent the determination of the bond length in such cases. It turned out that performing a set of two measurements using both light helicities as references in the calibration can circumvent these problems to some extent, as the measured MFPADs have to be by definition mirror-symmetric when switching between left- and right-handed polarization. Checking for this mirror symmetry when fine-tuning the calibration parameters of the experiment helped to decrease systematic errors on the required level of detail. While in Fig. 1 all six measured MFPADs agree well with the computed ones, we were not able to extract the offset angle with the required precision in all cases. For N2 at 780 eV, we recorded only one data set with one helicity, and its corresponding data point has a considerably larger error bar. Furthermore, the two measurements performed at the CO C 1s edge are compromised by a small systematic error due to overlapping flight times of the two ionic fragments, which directly (and almost solely) affected the part of the CP-MFPADs, from which θ_{tilt} is extracted. We included the lower-energy data point, with an estimate of the error, which is caused by the overlapping flight times. Such experimental difficulties should be considered in future measurements employing the θ_{tilt} approach. In the end, we evaluated several methods to extract the tilt angle from the experimental data. For example, the MFPADs can be fitted by a sum over spherical harmonics $Y_{l,m}$ with contributions of up to l = 9. By examining the fitting results, the tilt angle can be directly obtained from the corresponding maximum in the fitting function. However, estimating an error bar for this resulting angle is not straightforward. Instead, we finally determined the angular position of the forward-scattering peak by employing a Gaussian fit to the corresponding region of the measured angular distribution. Unless stated differently, the error bars shown in Fig. 3 are the errors reported by this fitting procedure. Within these error bars both approaches provide the same results.

Conclusions

In summary, we have investigated experimentally and theoretically the CP-MFPADs of core-level photoelectrons emitted from CO and N_2 molecules. We showed that (at high photoelectron energies) the tilt angle of the forward-scattering peak appearing in the CP-MFPAD of hetero-nuclear diatomic molecules can be directly connected to the bond length by a simple, universal equation. We applied the derived bond-length ruler formula to several examples of measured and computed CP-MFPADs. The extracted bond lengths agree well with the known bond lengths of CO and N2 in their neutral ground states. The present study illustrates that employing high-energy circularly polarized light, which is available not only at most synchrotron-radiation facilities, but also at XFEL facilities, has many advantages for the extraction of the molecular structure. It allows extraction of molecular bond lengths not only on a qualitative, but also on a quantitative level. We assume that our findings hold not only for the static case, but also for timeresolved MFPAD measurements aiming at catching the motion of individual atoms of molecules undergoing photoreactions, as, e.g., the stretching of chemical bonds. Further studies are required to target the extension of this method to larger molecules. The concept for such an extension is in principle straightforward (as indicated above), but there are still many aspects that need to be addressed when targeting real-life measurement results. Most prominently, the validity of the axial-recoil approximation (*i.e.*, a rapid fragmentation of the molecule as compared to possible rotation or geometrical changes) may not be given in all cases and needs to be confirmed by the measurement, as, e.g., done in ref. 38. However, previous work by Fehre et al.³⁹ demonstrated that it is possible to obtain three-dimensional MFPADs for a molecule as large as methyloxirane. This might be a first indication that the tilt-angle technique will be applicable to larger molecules in the future.

As compared to other experimental techniques for molecular structure determination, our bond-length ruler approach provides very easy access to molecular bond lengths. For example, Coulomb explosion imaging requires complex modeling of the charge-up process of the inspected molecule for the extraction of geometrical structures. Despite addressing single molecules in the gas phase, our bond-length ruler avoids the problems caused by randomly oriented samples (or even the need for crystalline samples), which other diffraction methods face. However, its explicit extension to larger molecules remains an intriguing subject for further studies, and our first prototypical demonstration shows that great care needs to be taken when measuring the MFPADs for the bond-length extraction in order to reach a precision in the range of a few percent.

Author contributions

M. S. S., T. J., R. D., P. V. D., and K. U. conceived the present work. I. V.-P., F. O., and A. M. contributed to it equally. The experiment was prepared and carried out by I. V.-P., J. R., N. M., S. U., G. N., N. A., D. Y., M. K., C. J., M. W., F. T., R. G., M. N. P., M. S., V. T. D., J. B. W., R. D., K. U., M. S. S., and T. J. Experimental data analysis was performed by I. V.-P., K. F., and T. J. *Ab initio* calculations were performed by A. M. and P. V. D. Analytical formulas were derived and employed for the analysis by F. O., Y. T., and K. H. F. O., Y. T., K. H., K. Y., K. U., P. V. D., R. D., M. S. S., and T. J. wrote the paper. All authors discussed the results and commented on the manuscript.

Conflicts of interest

There are no conflicts to declare.

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