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Excited Electronic States in Total Isotropic Scattering from Molecules

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Abstract

Ultrafast x-ray scattering experiments are routinely analyzed in terms of the isotropic scattering component. Here we present an analytical method for calculating total isotropic scattering for ground and excited electronic states directly from ab initio two-electron densities. The method is generalized to calculate isotropic elastic, inelastic, and coherent mixed scattering. The presented computational results focus on the potential for differentiating between electronic states and the decomposition of the total scattering in terms of elastic and inelastic scattering. For the specific example of the umbrella motion in the first excited state of ammonia, we show that redistribution of electron density along this coordinate leaves a comparably constant fingerprint in the total scattering that is similar in magnitude to the effect of changes in molecular geometry.

1 Introduction

The X-ray Free-Electron Laser (XFEL) facilities that have emerged around the world in the last decade provide coherent and ultrashort x-ray pulses, whose peak brightness is more than ten orders of magnitude larger than for synchrotron radiation. With a pulse duration that rivals that of optical lasers, XFELs have greatly enriched the palette of experimental techniques used to study the most fundamental aspects of chemistry – how atoms in a molecule move during a chemical reaction, how chemical bonds are made or broken, and how electrons rearrange after interaction with light. One such powerful technique is nonresonant ultrafast x-ray scattering from gas-phase samples. In a pump-probe fashion, an ensemble of molecules is pumped by an optical laser to an excited electronic state, and the resulting photodynamics is probed via hard x-rays with varying delay time. Due to the fast nature of the dynamics and the relatively small number of scattering molecules, these experiments require the ultrashort pulse duration and large photon numbers currently only provided by XFELs.

Gas-phase x-ray scattering has some fundamental differences compared to x-ray crystallography. On account of the large average separation between molecules and the absence of a regular lattice, the gas-phase scattering is free from intermolecular interferences for all but the smallest values of the scattering vector. The signal should thus be understood as an incoherent sum of scattering intensities from isolated molecules. It is on this single-molecule scale that quantum effects are most easily observed. In contrast, the periodicity of crystals means that the signal at the Bragg peaks is strongly dominated by elastic scattering, which, unlike the total scattering in gas phase, is a one-electron property. It is furthermore worth noting that x-ray crystallography is traditionally concerned with molecules in their thermal ground states, while the laser-induced dynamics in pump-probe experiments evolves on multiple electronic states, each characterized by
its own distinct electron distribution. It follows that the theoretical tools developed for and successfully applied in x-ray crystallography for more than a century are not always best suited for ultrafast x-ray scattering.\textsuperscript{16–26}

Indeed, gas-phase scattering from ground state molecules initially and ultrafast x-ray scattering more recently have prompted the development of a number of algorithms that aim to accurately predict x-ray scattering directly from the \textit{ab initio} electronic structure of molecules. As gas-phase samples in thermal equilibrium are isotropic, a central question in these methodologies is that of rotational averaging. Wang and Smith have first suggested a direct analytical method for evaluating isotropic scattering intensities.\textsuperscript{26} Different variants of this approach have been devised since, with the ultimate goal to reduce the computational cost.\textsuperscript{28–30} Alternative strategies that rely on numerical rotational averaging\textsuperscript{23,25,31} or grid-density methods\textsuperscript{26} have also been suggested. We note that although molecules may be initially aligned by the pump pulse in a pump-probe experiment, the isotropic component in the experimental scattering signal can be separated out by means of a Legendre decomposition,\textsuperscript{32–34} with the isotropic component equivalent to the signal of a fully isotropic ensemble.\textsuperscript{35,36}

Accurate, yet efficient calculations of isotropic total scattering are important in the context of geometry determination of molecules, where existing inversion techniques for retrieving molecular geometry from experimental data may require the evaluation of scattering from a very large number of molecular conformations.\textsuperscript{37} The same is true for the prediction of experimental signals from molecular simulations.\textsuperscript{21,26} Currently available algorithms are often computationally too demanding for such a high-throughput task.

In this article we extend the method developed by Crittenden and Bernard\textsuperscript{30} for calculating isotropic scattering as a sum of spherical Bessel functions. We demonstrate the existence of a recursive relationship between the expansion coefficients that allows for a significant speed-up of the calculation, in addition to generalising the approach to an arbitrary angular momentum. We illustrate how the method scales with the level of theory and the basis set used. We show that it is applicable not only to elastic and total scattering but also to inelastic and coherent mixed terms between different electronic states.\textsuperscript{24,38,39} This work constitutes a continuation of our development of methods to calculate elastic,\textsuperscript{19,40} inelastic,\textsuperscript{24} and total\textsuperscript{25} scattering from aligned molecules. In other work, we have examined the influence of specific rotational and vibrational states on elastic scattering.\textsuperscript{20,23} The importance of total scattering stems from the fact that in most ultrafast x-ray scattering experiments, it is the total (energy-integrated) scattering that is detected.\textsuperscript{1,5,10}

We demonstrate the computational methodology developed herein by investigating the different components of the total scattering in the ground and first excited states of ammonia along the umbrella normal mode coordinate. In this specific case we find that the change in the scattering signal upon optical excitation cannot be simply attributed to the changes in molecular geometry but is strongly influenced by the electron density redistribution in the excited state.

## 2 Theory

### 2.1 Scattering

For ultrafast x-ray scattering from photoexcited molecules, the differential scattering cross section per solid angle $\Omega$ takes the following form,\textsuperscript{39}

$$
\frac{d\sigma}{d\Omega} = \left(\frac{d\sigma}{d\Omega}\right)_{\text{Th}} W(\Delta\omega) \\
\times \int dt \ I(t) |\Psi(t)| \sum_{m,n} e^{i q \cdot (r_m - r_n)} |\Psi(t)|,
$$

where $\Psi(t)$ is the time-dependent wavefunction of the molecule, $q$ is the momentum transfer vector, and $r_m$ and $r_n$ are the position vectors of electrons $m$ and $n$, respectively. The detected signal is proportional to $(d\sigma/d\Omega)_{\text{Th}}$, which is the differential Thomson scattering cross-section for a free electron, which includes the polarization factor $|e \cdot e_2|$ of the incoming and scattered x-rays\textsuperscript{35,41} and the window function $W(\Delta\omega)$ with a detection window defined by $\Delta\omega$.\textsuperscript{39} The bra-ket notation implies integration over both the electronic and nuclear coordinates. Eq. (1) is valid in the limit of a large de-
tection window, \textit{i.e.} in the absence of energy resolution on the detector, as discussed in Ref. 39. This is currently the most common scenario in ultrafast x-ray scattering experiments,\textsuperscript{1,5,10} however different limits may also be considered.\textsuperscript{38,39} In the total scattering limit, all transitions are collectively detected with equal weight. Note that in deriving this equation the high mean photon energies of the x-rays compared to the energy spectrum of a typical molecule allow us to apply the Waller-Hartree approximation\textsuperscript{42} and disregard the comparatively small changes in the photon energy.

Upon photoexcitation the molecular wave function $|\Psi(t)\rangle$ is expressed in terms of the Born-Huang expansion in the basis of the $N$ electronic eigenstates $|\psi_k(\vec{r};\vec{R})\rangle = \langle \vec{r}|\psi_k(\vec{R})\rangle$ accessed during the dynamics. The electronic eigenstates are functions of the electronic coordinates $\vec{r}$ of the $N_e$ electrons in the molecule and depend parametrically on the internal nuclear coordinates $\vec{R}$,

$$|\Psi(t)\rangle = \sum_{k=1}^{N} |\chi^R_k(t)\rangle |\psi_k(\vec{R})\rangle.$$  \hfill (2)

In this expansion, each electronic state $|\psi_k(\vec{R})\rangle$ is multiplied by the corresponding time-dependent rovibrational nuclear wave packet $|\chi^R_k(t)\rangle$ that depends on the internal nuclear coordinates $\vec{R}$ and on the three Euler angles ($\alpha, \beta, \gamma$), which relate the molecular and laboratory frames. Using Eq. (2), the differential scattering cross section becomes,\textsuperscript{39}

$$\frac{d\sigma}{d\Omega} = \left(\frac{d\sigma}{d\Omega}\right)_{\text{Th}} W(\Delta\omega) \times \sum_{i,j}^{N} \int dt \ I(t) \langle \chi^R_i(t) | \Lambda_{ij}(\vec{q},\vec{R}) | \chi^R_j(t) \rangle,$$  \hfill (3)

The key quantity in Eq. (3) is $\Lambda_{ij}(\vec{q},\vec{R})$, which is the two-electron scattering matrix element,

$$\Lambda_{ij}(\vec{q},\vec{R}) = \sum_{m,n}^{N_e} \langle \psi_j(\vec{R}) | e^{i \vec{q} \cdot (\vec{r}_m - \vec{r}_n)} | \psi_j(\vec{R}) \rangle.$$  \hfill (4)

Since terms with $m = n$ in Eq. (4) reduce to the Kronecker delta $\delta_{ij}$, the two-electron scattering matrix element can be written as,

$$\Lambda_{ij}(\vec{q},\vec{R}) = N_e \delta_{ij} + \Lambda'_{ij}(\vec{q},\vec{R}),$$  \hfill (5)

where $\Lambda'_{ij}(\vec{q},\vec{R})$ is the pure two-electron part of $\Lambda_{ij}(\vec{q},\vec{R})$ with $m \neq n$. Using the sifting property of the Dirac delta function, $\exp[i(\vec{q} \cdot (\vec{r}_m - \vec{r}_n))]$ can furthermore be expressed in an integral form:

$$e^{i \vec{q} \cdot (\vec{r}_m - \vec{r}_n)} = \int dr_1 dr_2 e^{i \vec{q} \cdot (\vec{r}_1 - \vec{r}_2)} \times \delta(\vec{r}_1 - \vec{r}_m)\delta(\vec{r}_2 - \vec{r}_n),$$  \hfill (6)

so that the integral over the electronic coordinates in Eq. (4) becomes,

$$\Lambda_{ij}(\vec{q},\vec{R}) = N_e \delta_{ij} + 2 \int dr_1 dr_2 \rho_{ij}^{(2)}(\vec{r}_1,\vec{r}_2,\vec{R}) e^{i \vec{q} \cdot (\vec{r}_1 - \vec{r}_2)},$$  \hfill (7)

where $\rho_{ij}^{(2)}(\vec{r}_1,\vec{r}_2,\vec{R})$ is the expectation value of the two-electron density operator $\hat{\rho}(\vec{r}_1,\vec{r}_2) = (1/2) \sum_{m}^{N_e} \sum_{n: \neq m}^{N_e} \delta(\vec{r}_1 - \vec{r}_m)\delta(\vec{r}_2 - \vec{r}_n)$.\textsuperscript{43} At that point, it is prudent to differentiate between the diagonal elements with respect to the electronic states with $i = j$ and off-diagonal (mixed) terms with $i \neq j$. In the former case, $\rho_{ij}^{(2)}(\vec{r}_1,\vec{r}_2,\vec{R})$ gives the probability of finding one of the electrons of the system in state $|\psi_j(\vec{R})\rangle$ at $\vec{r}_1$ while another electron is at $\vec{r}_2$. This term can be further separated into two contributions by expanding the two-electron density as a sum of products over one-electron density functions, $\rho_{ij}^{(1)}(\vec{r},\vec{R})$,

$$N_e + 2 \rho_{ij}^{(2)}(\vec{r}_1,\vec{r}_2,\vec{R}) = \rho_{ij}^{(1)}(\vec{r}_1,\vec{R})\rho_{ij}^{(1)}(\vec{r}_2,\vec{R}) + \sum_{f \neq i}^{N_e} \rho_{ij}^{(1)}(\vec{r}_1,\vec{R})\rho_{j\hat{f}}^{(1)}(\vec{r}_2,\vec{R}),$$  \hfill (8)

which follows from insertion of the resolution of the identity in the basis of the electronic states. The contribution to the total scattering from the first term on the right-hand side of Eq. (8) is electronically elastic, while the contribution from the second term is electronically inelastic. We note that this terminology becomes inappropriate in the presence of strong non-adiabatic coupling between the electronic states, when the individual elastic or inelastic contributions in Eq. (8) become dependent on the choice of electronic basis.\textsuperscript{39} Import-
tantly, the observable signal involves contributions from all non-adiabatically coupled states and is independent of the basis.

When \( i \neq j \), the quantity \( p_{ij}^{(2)}(r_1, r_2, \mathbf{R}) \) in Eq. (7) is referred to as the two-electron transition density function \((i.e.\ the\ diagonal\ part\ of\ the\ density\ matrix)\) in order to differentiate it from the case of \( i = j \), which is simply known as the two-electron density function. The \( i \neq j \) scattering components play a critical role in coherent mixed scattering, which appears when there is a coherence between two electronic states.\(^{38,44–46}\) Finally, we note that quite similar matrix elements to those considered here appear for electron scattering.\(^{24,31,47–49}\)

### 2.2 Isotropic scattering

The main focus of this article is the development of an efficient methodology to calculate isotropic scattering signals at specific molecular geometries. This is motivated by the central role that the isotropic scattering signal plays in the interpretation of experiments, irrespective of the degree of alignment in the sample.\(^{35,36}\) To appreciate this, we must consider the standard approach in gas-phase scattering experiments to decompose the observed signal in the basis of orthogonal Legendre polynomials, \( P_{\alpha}(\cos \theta_q) \),\(^{32}\)

\[
\frac{d\sigma}{d\Omega}(q) = \sum_{\alpha} P_{\alpha}(\cos \theta_q) S_{\alpha}(q), \tag{9}
\]

where \( \cos \theta_q \) is the component of the unit scattering vector projected upon the polarization axis of the pump laser shown as the Z axis in Fig. 1. The angle \( \theta_q \) is related to the detector angles by \( \cos \theta_q = \sin(\theta_d/2) \cos \delta - \cos(\theta_d/2) \cos \phi_d \sin \delta \), where \( \delta \) is the angle between the polarization axis of the pump laser and the x-ray beam’s direction of propagation. In the case of a pump-probe arrangement where the laser polarisation axis is perpendicular to the x-ray propagation, \( i.e.\ the\ case\ discussed\ here,\ this\ reduces\ to\ \cos \theta_q = -\cos \theta_d/2 \cos \phi_d \). It should be emphasised that the Legendre polynomials in Eq. (9) are functions of \( \theta_q \) and not of the detector angles \( \theta_d \) and \( \phi_d \). The advantage of this approach is that the contributions from the internal and external molecular degrees of freedom can be separated out, as will be shown below. It is important to point out that in many pump-probe gas phase scattering experiments, the detected signal is rarely fully isotropic on account of the preferential excitation of the molecules whose transition dipole moments align with the polarization axis of the linearly polarized pump laser.\(^{8}\)

![Figure 1: Illustration of the geometrical relations in x-ray scattering.](image)

A remarkable property of the decomposition in Eq. (9) is that information on the internal degrees of freedom can be extracted from any of the components, \( S_{\alpha}(q) \), or a combination of them.\(^{50}\) While in terms of order greater than zero the internal dynamics is intermingled with information about the rotational wavepacket, the zeroth order term in the Legendre expansion can be analysed solely from the point of view of the internal molecular degrees of freedom. The quantity \( S_0(q) \) is equivalent to the scattering from a fully isotropic ensemble and consistent with the framework of Debye scattering (see SI for derivation). Using the orthogonality of the Legendre polynomials and that \( P_0(\cos \theta_q) = 1, \)
the zeroth-order term, which is henceforth referred to as isotropic scattering, is given by,

$$
S_0(q) = \frac{1}{4\pi} \int_0^{2\pi} d\phi_q \int_0^\pi d\theta_q \sin\theta_q \frac{d\sigma}{d\Omega}(q) = \left(\frac{d\sigma}{d\Omega}(q)\right)_{\theta_q \phi_q},
$$

(10)

where the average is taken over the angular coordinates of the momentum transfer vector $q$. With reference to the full expression for the differential scattering cross-section in Eq. (3), only the two-electron scattering matrix element $\Lambda_{ij}(q, \vec{R})$ has to be averaged, as it is the only quantity that depends on $q$. Furthermore, the average renders the scattering signal independent of the orientation of the molecule in space, meaning that $\Lambda_{ij}(q, \vec{R}) = \langle \Lambda_{ij}(q, \vec{R}) \rangle_{\theta_q \phi_q}$ does not depend on the Euler angles (detailed derivation included in the SI). That allows for a separate integration of the rotational wave packets, resulting in a simple scaling factor, which is equal to one for $i = j$.

$$
S_0(q) = \left(\frac{d\sigma}{d\Omega}\right)_{\text{Th}} W(\Delta\omega) \times \sum_{i,j} \int dt I(t) \langle \Theta_i^f \Theta_j^f | \chi_j^i \rangle | \Lambda_{ij}(q, \vec{R}) | \chi_j^i \rangle,
$$

(11)

where the vibrational and rotational components of the rovibrational wavepacket, $| \chi_j^i(t) \rangle$ and $| \Theta_j^f(t) \rangle$, respectively, have been made explicit and their dependence on time has been omitted for brevity. It follows that the isotropic component of the experimental scattering signal can be analyzed as if the entire ensemble had a fully isotropic rotational distribution.

3 Methodology

We now turn our attention to the derivation of a method that permits an efficient analytical evaluation of the isotropic differential scattering cross section. We will seek a solution for the rotationally averaged two-electron scattering matrix elements, $\Lambda_{ij}(q, \vec{R}) = \langle \Lambda_{ij}(q, \vec{R}) \rangle_{\theta_q \phi_q}$, which are a prerequisite for a more detailed description that involves the role of nuclear motion later on. From Eq. (7) we need to evaluate the expression,

$$
\Lambda_{ij}(q, \vec{R}) = N_e \delta_{ij} + 2 \left( \int \int dr_1 dr_2 \rho_{ij}^{(2)}(r_1, r_2, \vec{R}) e^{-i \phi \cdot (r_1 - r_2)} \right),
$$

(12)

In the absence of energy resolution, i.e. the standard set-up for current time-resolved scattering experiments, the separation of the total scattering into its elastic and inelastic components is not needed, but it is instructive to show that our methodology is applicable to all four cases: total, elastic, inelastic, and coherent mixed. We can split the expression for total scattering ($i = j$) using Eq. (8),

$$
\Lambda_{ii}(q, \vec{R}) = \sum_f \left( \int \int dr_1 dr_2 \times \rho_{ij}^{(1)}(r_1, \vec{R}) \rho_{ji}^{(1)}(r_2, \vec{R}) e^{-i \phi \cdot (r_1 - r_2)} \right),
$$

(13)

where terms with $f = i$ are elastic, while those with $f \neq i$ are inelastic. The key quantities in Eqs. (12) and (13) are the one- and two-electron density functions, $\rho_{ij}^{(1)}(r_1, \vec{R})$ and $\rho_{ij}^{(2)}(r_1, r_2, \vec{R})$, which can be expressed as weighted products of molecular orbitals (dropping the parametric dependence on the nuclear coordinates),

$$
\rho_{ij}^{(1)}(r_1) = \sum_{a,b} D_{ij}^{ab} \phi_a(r_1) \phi_b(r_1),
$$

(14)

and

$$
\rho_{ij}^{(2)}(r_1, r_2) = \frac{1}{2} \sum_{a,b,c,d} d_{abcd} \phi_a(r_1) \phi_b(r_1) \times \phi_c(r_2) \phi_d(r_2),
$$

(15)

where the indices run over all $N_{MO}$ occupied molecular orbitals. The coefficients $D_{ij}^{ab}$ and $d_{abcd}$ with $i = j$ are referred to as the elements of the one- and two-electron reduced density matrix (1- and 2-RDM), respectively. If $i \neq j$, they are known as one- and two-electron reduced transition density matrix elements.

Insertion of Eqs. (14) and (15) into the expression for the isotropic scattering, Eqs. (12) and (13), shows that the efficient $ab\ initial$ solution to the
isotropic scattering problem requires the evaluation of integrals of the type,

\[
K_{ij}(q) = \left\langle \int d\mathbf{r}_1 d\mathbf{r}_2 \sum_{a,b,c,d} z_{abcd}^j \phi_a(\mathbf{r}_1) \phi_b(\mathbf{r}_1) \times \phi_c(\mathbf{r}_2) \phi_d(\mathbf{r}_2) e^{i\mathbf{q} \cdot (\mathbf{r}_1-\mathbf{r}_2)} \right\rangle_{\theta_\epsilon \phi_\epsilon},
\]

(16)

where

\[
z_{ij}^{abcd} = \begin{cases} d_{ii}^{abcd}, & \text{total (} i = j \text{)} \\ D_{ij}^{ab} D_{jk}^{cd}, & \text{elastic (} i = j \text{)} \\ D_{ij}^{ab} D_{kj}^{cd}, & \text{inelastic (} i \neq j \text{)} \\ d_{ij}^{abcd}, & \text{coherent mixed (} i \neq j \text{)} \end{cases}
\]

Note that the inelastic component of the scattering signal in Eq. (17) refers to an individual electronic transition \(|\psi_i(\mathbf{R})\rangle \leftarrow |\psi_j(\mathbf{R})\rangle\), not to the total inelastic signal that is measured in experiments without energy resolution. The latter corresponds to an infinite sum of transitions to all electronic states and can be obtained by subtracting the elastic from the total scattering signal. A common strategy in molecular electronic structure theory is to expand the orbitals as a weighted sum of \(N_{bf}\) primitive Cartesian Gaussian-type orbitals (GTOs),

\[
\phi_a(\mathbf{r}_1) = \sum_{\mu} N_{bf} M^{(a)}_{\mu} g_{\mu}(\mathbf{r})
\]

(18)

\[
= \sum_{\mu} N_{bf} M^{(a)}_{\mu} (x-A_{\mu x})^l (y-A_{\mu y})^m e^{-\gamma_{\mu} (r-A_{\mu})^2}
\]

where \(A_{\mu}\) is the centre of the \(\mu\)th primitive Gaussian and \(M^{(a)}_{\mu}\) the molecular orbital expansion coefficient. In the case of contracted Gaussian functions, \(M^{(a)}_{\mu}\) is premultiplied by a contraction coefficient. The sums of \(l_{\mu}, m_{\mu}\), and \(n_{\mu}\) specify the orbital angular momentum. The integral \(K_{ij}(q)\) in Eq. (16) then takes the form,

\[
K_{ij}(q) = \sum_{\mu,\nu,\xi,\zeta} N_{bf} z_{\mu \nu \zeta \xi} \langle J_{\mu \nu}(q) J^*_{\xi \zeta}(q) \rangle_{\theta_\epsilon \phi_\epsilon},
\]

(19)

with the two-electron charge density \(Z_{\mu \nu \zeta \xi}^{ij} = \sum_{a,b,c,d} z_{ij}^{abcd} M^{(a)}_{\mu} M^{(b)}_{\nu} M^{(c)}_{\xi} M^{(d)}_{\zeta}\) and where we have labeled the Fourier integrals over \(\mathbf{r}_1\) and \(\mathbf{r}_2\) as \(J_{\mu \nu}(q)\) and \(J^*_{\xi \zeta}(q)\), respectively,

\[
J_{\mu \nu}(q) = \int d\mathbf{r}_1 g_{\mu}(\mathbf{r}_1) g_{\nu}(\mathbf{r}_1) e^{i\mathbf{q} \cdot \mathbf{r}_1}
\]

(20)

and

\[
J^*_{\xi \zeta}(q) = \int d\mathbf{r}_2 g_{\xi}(\mathbf{r}_2) g_{\zeta}(\mathbf{r}_2) e^{-i\mathbf{q} \cdot \mathbf{r}_2}.
\]

(21)

Eq. (19) reveals that the calculation of the \textit{ab initio} isotropic scattering requires the evaluation of \(N_{bf}^4\) angular integrals. This is a formidable computational challenge, even for the smallest molecules with an adequate basis set. However, schemes for the efficient evaluation of such integrals have been proposed \cite{27,30} and make use of the properties of the Gaussian functions and their analytical Fourier transforms. The first step is to express the angular momentum properties of the product of two primitive Gaussians, \(g_{\mu}(\mathbf{r})\) and \(g_{\nu}(\mathbf{r})\), in a derivative form, \cite{51}

\[
\Pi_{\mu \nu}(\mathbf{r}) = g_{\mu}(\mathbf{r}) g_{\nu}(\mathbf{r})
\]

\[
= \mathcal{G}_{\mu \nu} \sum_{L_1=0}^{l_{\mu}+l_{\nu}} \sum_{M_1=0}^{m_{\mu}+m_{\nu}} \sum_{N_1=0}^{n_{\mu}+n_{\nu}} E_{L_1 M_1 N_1}^{l_{\mu} m_{\mu} l_{\nu} m_{\nu}} E_{L_1 M_1 N_1}^{n_{\mu} n_{\mu} n_{\nu} n_{\nu}}
\]

\[
\times \left\langle \frac{\partial}{\partial P_x} \right\rangle_{L_1} \left\langle \frac{\partial}{\partial P_y} \right\rangle_{M_1} \left\langle \frac{\partial}{\partial P_z} \right\rangle_{N_1} e^{-\gamma_{\mu}(r-A_{\mu})^2},
\]

(22)

where we do not explicitly show the dependence of the McMurchie-Davidson expansion coefficients \(E_{L_1 M_1 N_1}^{l_{\mu} m_{\mu} l_{\nu} m_{\nu}}\) and \(E_{L_1 M_1 N_1}^{n_{\mu} n_{\mu} n_{\nu} n_{\nu}}\) on the Cartesian components of the centres of the Gaussian functions \(x_\mu\) and \(x_\nu\) as well as on the Gaussian exponents \(\gamma_{\mu}\) and \(\gamma_{\nu}\). The expression also exploits that the product of two Gaussians functions is another Gaussian function, leading to the following definitions that appear in Eq. (22),

\[
\gamma_P = \gamma_{\mu} + \gamma_{\nu},
\]

\[
\mathcal{G}_{\mu \nu} = e^{-\gamma_{\mu} \gamma_{\nu}(A-B)^2},
\]

(23)

\[
P = (\gamma_{\mu} A + \gamma_{\nu} B)/\gamma_{P}.
\]

The McMurchie-Davidson expansion is widely used in computational chemistry programme pack-
ages to express integrals of Gaussian functions with high angular momentum as derivatives of s-type Gaussian integrals. Their utility stems from the existence of a recursive relationship between the coefficients, which enables their rapid evaluation. Substituting $\Pi_{\mu\nu}(r)$ into the expression for $J_{\mu\nu}(q)$ in Eq. (20) and moving the derivative in front of the Fourier integral result in,

$$J_{\mu\nu}(q) = \int dr \Pi_{\mu\nu}(r)e^{iqr}$$

$$= G_{\mu\nu} \sum_{L_1=0}^{l_1} \sum_{M_1=0}^{m_1} \sum_{N_1=0}^{n_1} E_{l_1}^{m_1} E_{N_1}^{n_1}$$

$$\times \left( \frac{\partial}{\partial P_x} \right)^{L_1} \left( \frac{\partial}{\partial P_y} \right)^{M_1} \left( \frac{\partial}{\partial P_z} \right)^{N_1}$$

$$\times \int d\mathbf{r} e^{-\gamma_{r-P}^2} e^{iq\mathbf{r}}.$$  \hspace{1cm} (24)

The Fourier Transform of the s-type Gaussian function in the last line of Eq. (24) can be evaluated analytically using the Fourier shift property,

$$\int d\mathbf{r} e^{-\gamma_{r-P}^2} e^{iq\mathbf{r}} = \left( \frac{\pi}{\gamma_P} \right)^{3/2} e^{-q^2/4\gamma_P} e^{iq\mathbf{P}}.$$  \hspace{1cm} (25)

Having performed the Fourier Transform of the product of two GTOs, the full expression for $J_{\mu\nu}(q)J_{\xi\zeta}(q)$ prior to integrating out the angular dependence becomes,

$$J_{\mu\nu}(q)J_{\xi\zeta}^*(q) = \int dr_1 \Pi_{\mu\nu}(r_1)e^{iqr_1} \int dr_2 \Pi_{\xi\zeta}(r_2)e^{-iqr_2}$$

$$= \frac{\pi^3 G_{\mu\nu} G_{\xi\zeta}}{(\gamma_{r-P}^2)^{3/2}} e^{-q^2(1/1\gamma+1/\gamma_P)/4}$$

$$\times \sum_{L_1=0}^{l_1} \sum_{M_1=0}^{m_1} \sum_{N_1=0}^{n_1} \sum_{L_2=0}^{l_2} \sum_{M_2=0}^{m_2} \sum_{N_2=0}^{n_2}$$

$$\times E_{l_1}^{m_1} E_{N_1}^{n_1} E_{l_2}^{m_2} E_{N_2}^{n_2}$$

$$\times F_{L_1M_1N_1}^{L_2M_2N_2}(q,P,Q).$$  \hspace{1cm} (26)

where

$$F_{L_1M_1N_1}^{L_2M_2N_2}(q,P,Q) = \left( \frac{\partial}{\partial P_x} \right)^{L_1} \left( \frac{\partial}{\partial P_y} \right)^{M_1} \left( \frac{\partial}{\partial P_z} \right)^{N_1}$$

$$\times \left( \frac{\partial}{\partial Q_x} \right)^{L_2} \left( \frac{\partial}{\partial Q_y} \right)^{M_2} \left( \frac{\partial}{\partial Q_z} \right)^{N_2} e^{i(q-P)\cdot Q}.$$  \hspace{1cm} (27)

Introducing $H = P - Q$ and the combined angular momentum quantum numbers $L = L_1 + L_2$, $M = M_1 + M_2$, and $N = N_1 + N_2$, Eq. (27) can be written as,

$$F_{L_1M_1N_1}^{L_2M_2N_2}(q,P,Q) = (-1)^{L_2+M_2+N_2}$$

$$\times \left( \frac{\partial}{\partial H_x} \right)^L \left( \frac{\partial}{\partial H_y} \right)^M \left( \frac{\partial}{\partial H_z} \right)^N e^{iH}.$$  \hspace{1cm} (28)

It should be recognized that, if there was no need to perform the rotational average calculation, \textit{i.e.} for scattering of aligned molecules, the derivatives in Eq. (28) trivially evaluate to $(iH)^L(iH)^M(iH)^N\exp[iH\cdot H]$. However, even in the case of isotropic scattering the quantity $\langle J_{\mu\nu}(q)J_{\xi\zeta}^*(q) \rangle_{\theta,\phi,q}$ expressed in its current form has a relatively simple analytic solution. Resolving the angular integrals, which only affects $\exp[iH\cdot H]$, results in,

$$\langle F_{LMN}(q,H) \rangle_{\theta,\phi,q} = \left( \frac{\partial}{\partial H_x} \right)^L \left( \frac{\partial}{\partial H_y} \right)^M \left( \frac{\partial}{\partial H_z} \right)^N \sin qH.$$  \hspace{1cm} (29)

Solutions to the the equation above are discussed by Wang \textit{et al.},\textsuperscript{27} where it is given as a four dimensional sum over trigonometric functions scaled by precalculated numerical factors. Here, we follow more closely the approach suggested by Critenden \textit{et al.},\textsuperscript{30} who calculated the result analytically for a limited number of angular momenta as a sum of spherical Bessel functions. In contrast to their approach, we recognize the existence of a recursive relationship between the expansion coefficients, which allows for a fast calculation and handling of arbitrarily large angular momenta. For that we will distinguish two cases defined by a cutoff value $\epsilon_{\text{cut}}$. When $H < \epsilon_{\text{cut}}$, the exponential in Eq. (28) is approximately unity and Eq. (29) takes
the form,

\[ \langle F_{LMN}(q, H) \rangle_{\theta \phi q} = i^{L+M+N} \langle q_x^L q_y^M q_z^N \rangle_{\theta \phi q} = B_{LMN}(iq)^{L+M+N}, \tag{30} \]

where the average involves only the Cartesian components of \( q \) that stem from the derivatives of \( \exp[iq \cdot H] \) and where

\[ B_{LMN} = \left( \sin^{L+M} \theta_q \cos^N \phi_q \cos^L \phi_q \sin^M \theta_q \right)_{\theta \phi q} \tag{31} \]

is a numerical constant. When \( H \geq \epsilon_{\text{cut}} \), the evaluation of the derivatives relies on the properties of the spherical Bessel functions, \( j_\beta(qH) \),

\[ \langle F_{LMN}(q, H) \rangle_{\theta \phi q} = \left( \frac{\partial}{\partial H_x} \right)^L \left( \frac{\partial}{\partial H_y} \right)^M \left( \frac{\partial}{\partial H_z} \right)^N \sin qH \left( \frac{q}{H} \right)^\beta j_\beta(qH), \]

\[ = \sum_{p=0}^{L} \sum_{s=0}^{M} \sum_{t=0}^{N} a^p_L(H_x)b^s_M(H_y)c^t_N(H_z) \left( \frac{q}{H} \right)^\beta j_\beta(qH), \tag{32} \]

where \( \beta = \left[ (L+M+N-p-s-t)/2 \right] + p + s + t \) with the ceiling function \( \lceil \ldots \rceil \) that gives the least integer greater than or equal to its argument. The coefficients \( a^p_L, b^s_M, \) and \( c^t_N \) are related to the Hermite polynomials and obey the following recursive relations (here given for \( a^p_L \))

\[ a^p_L(H_x) = \begin{cases} 1, & L = 0, \ p = 0 \\ 0, & L = 1, \ p = 0 \\ -H_x, & L = 1, \ p = 1 \\ -a^{p-1}_{L-2}(L-1), & L > 1, \ p = 0 \\ -a^p_{L-1}H_x - a^{p-1}_{L-2}(L-1), & L > 1, \ p > 0 \end{cases} \tag{33} \]

The implementation of a sensible cut-off value, \( \epsilon_{\text{cut}} \), is essential for the numerical stability of the algorithm, which might otherwise be affected by prohibitively large values of \( (q/H)^\beta \) factor in Eq. (32). The efficient procedure for evaluating Eq. (29) makes use of the recursive relationship for the coefficients \( a^p_L, b^s_M, \) and \( c^t_N \) as well as of the recursive formula for the spherical Bessel functions. In addition to that, it is of paramount im-
portance for the computational efficiency to take into account the symmetries \( (J_{\mu \nu}(q))_{\xi \xi}(q)_{\theta \phi q} = (J_{\xi \xi}(q))_{\mu \nu}(q)_{\theta \phi q} \) and \( J_{\mu \nu}(q) = J_{\nu \mu}(q) \), which can be utilized to yield a speed-up factor of approximately 8. Another important simplification stems from careful consideration of the contraction scheme of the basis set used. If a given primitive GTO is a part of multiple contractions, the corresponding integral should be performed only once and the constant \( Z_{\mu \nu \xi \xi}^{ij} \) needs to be modified to reflect the combined contribution of this primitive to the molecular orbitals. The final trick for improving the computational performance is to treat together all GTOs whose centres and exponents are the same. Careful examination of these cases shows that they ultimately lead to the same values of \( H \) and can only differ in their angular momentum numbers \( l, m, \) and \( n \). Treating them together results in a family of \( \langle F_{LMN}(q, H) \rangle_{\theta \phi q} \) integrals, for which most of the terms in Eq. (32) are shared. In addition, a global cut-off linked to the relative size of \( z_{abcd}^{ij} \) could significantly speed up the calculation, at the expense of an effective decrease of the total number of electrons integrated. We have observed that allowing for 0.1% electron density loss can lead to an approximate speed-up factor of around two in most molecules explored without significant effect on the results.

## 4 Results and Discussion

### 4.1 Benchmarking and scaling

Our method is tested and validated in a series of \( \textit{ab initio} \) calculations for the ammonia molecule, NH\(_3\), varying the basis set and the size of the active space. All \( \textit{ab initio} \) calculations are performed using the MOLPRO electronic structure software package.\(^{52,53}\) We chose ammonia because it has previously been used by Hoffmeyer et al.\(^{51}\) to illustrate the importance of multiconfigurational wavefunctions in total x-ray and electron scattering. Although, as shown above, our methodology also encompasses individual inelastic transitions and coherent mixed scattering, we focus on the total and elastic signals here. The discussion of inelastic scattering is limited to the cumulative sum of all inelastic transitions, which is given by
the difference between total and elastic scattering. The richness of the information encoded in the inelastic and coherent mixed terms will be the target of a follow-up publication.

Fig. 2 compares total and elastic scattering at different levels of theory, namely Hartree-Fock (HF) and Complete Active Space Self-Consistent Field (CASSCF) with 6 and 8 active orbitals and all electrons active, i.e. CASSCF(10,6) and CASSCF(10,8), respectively. The basis sets include Pople’s and Dunning’s correlation-consistent basis sets with double-zeta, double-zeta plus diffuse functions, and triple-zeta plus diffuse functions. Calculations with the STO-3G minimal basis set are also performed and are included in Tables S1 and S2 in the SI, which summarise the benchmark results. The geometry used in all calculations is optimized at the CASSCF(10,8)/aug-cc-pVTZ level of theory. The results are first calculated as the fractional signal change,

$$\Delta S(q) = \frac{I(q) - I_{\text{ref}}(q)}{I_{\text{ref}}(q)}, \quad (34)$$

where the reference, $I_{\text{ref}}(q)$, is either the total or the elastic scattering cross section computed at the CASSCF(10,8)/aug-cc-pVTZ level. We refer to the fractional signal change to ensure that the difference at each value of momentum transfer $q$ is relative to the absolute value of the intensity at this point. The results presented in Fig. 2 are given as the integral of the absolute values in the range $[q_{\text{min}}, q_{\text{max}}] = [0, 11.34] \, \text{Å}^{-1}$,

$$E = \int_{q_{\text{min}}}^{q_{\text{max}}} dq \, |\% \Delta S(q)|. \quad (35)$$

where $\% \Delta S(q)$ is the fractional signal change in percent, i.e. $\% \Delta S(q) = \Delta S(q) \times 100\%$. It is worth noting that the total and elastic scattering signals in the limit of large $q$ differ by the number of the electrons in the molecule, which results in a larger value in the denominator in Eq. (34) and a smaller integral in Eq. (35) for total scattering.

It is clear that the two families of basis functions are comparable at a given level of theory. Interestingly, Pople’s basis sets with the smaller active space and Hartree-Fock seem to be closer to the reference CASSCF(10,8)/aug-cc-pVTZ reference calculation compared to the correlation-consistent basis sets. It is possible that the split-valence basis sets provide a better description in cases where the active space is not sufficient to capture the static electron correlation adequately. In addition and not surprisingly, we found that STO-3G is largely unsuitable for scattering calculations (see Tables S1 and S2 in the SI). In fact, STO-3G performs similarly to the Independent Atom Model (IAM), which gives $E = 17.2 \, \text{Å}^{-1}$ and $E = 32.8 \, \text{Å}^{-1}$ for total and elastic scattering. Overall, the convergence increases smoothly with the number of basis functions, both with Pople’s and Dunning’s basis sets. The most significant effect is seen when going from the smaller CASSCF(10,6) to the larger CASSCF(10,8) active space, which can be attributed to the effects of electron correlation on the electron density of the system. We note that the total scattering cross section is largely a two-electron property and can be expected to be significantly affected by electron correlation. Moreover, it is clear that the convergence of the elastic scattering is equally affected, suggesting that the electron density relaxation associated with static correlation, implicit in the CASSCF calculations, is a major factor in that case. For accurate total scattering calculations, *ab initio* methods that capture dynamic correlation are desirable.

Fig. 3 compares the computational time required for the calculations discussed above. Generally, the calculations scale with the fourth power of the number of basis functions, $N_{\text{bf}}^4$, as all distinctive Fourier transforms over four basis functions need to be considered as long as they are not related by permutational symmetry. As discussed above, grouping together integrals with equal origin and exponents partially offsets this scaling. Practically, the number of unique centres increases with the number of atoms, so that the speed up for large molecules can be small. In the case of elastic scattering, the two-electron charge density $Z_{\mu \nu \xi}^{ij}$ can be expressed as two independent pairs $Z_{\mu \psi}^{ij}$ and $Z_{\xi \xi}^{ij}$. After precomputing them, they can easily be accessed in the calculation with no additional computational cost. Hence, the computational time required for the calculation of elastic scattering is largely independent of the level of theory or active space used. For the total scattering, the two-electron charge density remains a function of all
Figure 2: Convergence of the total and elastic x-ray scattering signals of NH$_3$ computed with HF, CASSCF(10,6) and CASSCF(10,8) using various basis sets. The height of the bars represent the integral of the percentage intensity change with respect to CASSCF(10,8)/aug-cc-pVTZ total and elastic scattering for the range 0 \leq q \leq 11.34 \text{ Å}^{-1}.

Four GTOs, so that its on-the-fly calculation increases the computational time compared to the elastic scattering in a manner that scales with the number of active orbitals. The scalings for specific inelastic and coherent mixed terms are equivalent to elastic and total scattering, respectively. We note that the scattering calculation with the largest basis set used in this work is three orders of magnitude more expensive than a simple calculation with a minimal basis set. This scaling is of great practical importance for choosing an optimal method for scattering from a wide range of molecular geometries sampled along a reaction coordinate or for the purpose of iterative inversion of experimental data. Given the comparatively small overhead cost of total scattering, it is sensible to adopt this type of calculation in lieu of the common approach of calculating total scattering as a sum of elastic scattering and tabulated inelastic corrections.$^{1,4,5}$

In Fig. 4 we show the total, elastic, and inelastic scattering curves of NH$_3$ calculated at the reference CASSCF(10,8)/aug-cc-pVTZ level. The comparison with previous MR-SDCI (Multi-Reference Single and Doubles Configuration Interaction) calculations by Hoffmeyer et al.\textsuperscript{31} shows rather good agreement given the differences in the methods, levels of theory, and basis sets. Their approach relies on numerical integration, whereas our result is strictly analytical. Our best calculation employs aug-cc-pVTZ, while their work reports a smaller double-zeta basis set with polarization and diffuse functions, [5s3p2d/3s2p].

Figure 3: Comparison of the computational time required for the calculation of the isotropic elastic and total x-ray scattering signals in NH$_3$. Different ab initio methods are considered as well as basis sets. Note that the computational scaling is expressed as a logarithm of the CPU time in milliseconds. The solid part of the bars represent the elastic scattering, whereas the shaded area at the top shows the extra time required to compute the total scattering with the same level of theory and basis set.
Unlike MR-SDCI, CASSCF calculations account only for static electron correlation. The dynamic electron correlation has a smaller effect on the elastic component of the scattering signal as seen by comparing to the MR-SDCI results. Its influence increases when total and inelastic scattering are considered, demonstrating the importance of electron correlation for these quantities. A systematic study of the effect of electron correlation is critically important for fully understanding gas-phase x-ray scattering experiments and will be addressed in subsequent work.

4.2 Excited state total scattering

Table 1: The ground state equilibrium geometry and geometries along the umbrella normal mode in the first excited state. The geometries are presented in terms of the N–H bond distance and the pyramidization angle defined as the angle between the plane of the H atoms and any of the N–H bonds. The equilibrium geometries in both states are displayed in bold font.

| State    | Pyramidization angle | N–H bond / Å |
|----------|----------------------|--------------|
| Ground   | 23.5°                | 1.022        |
|          | 0.0°                 | 1.032        |
|          | 3.7°                 | 1.034        |
|          | 7.4°                 | 1.041        |
|          | 11.0°                | 1.051        |
|          | 14.6°                | 1.066        |
| Excited  | 18.0°                | 1.085        |
|          | 21.3°                | 1.108        |
|          | 24.5°                | 1.133        |
|          | 27.5°                | 1.163        |
|          | 30.3°                | 1.196        |
|          | 33.0°                | 1.231        |

Here, we consider a simple model that illustrates both the utility of our methodology and the nature of the signal detected in ultrafast x-ray scattering. For our purpose, a suitable candidate is the photoexcitation of ammonia to the first electronically excited singlet state, whose initial dynamics follows an umbrella motion. The goal is to track the changes and dominant contributions to the total scattering as the geometry changes. In order to achieve that, we optimize the ground state ion geometry at the CASSCF(9,8)/6-31+G* level as an approximation for the first excited state, which has Rydberg 3s character. The normal modes are calculated. The molecular geometry is then displaced in a series of steps along the umbrella mode and at each geometry the ground and the first excited states are calculated in a state-average fashion at SA2-CASSCF(10,8)/aug-cc-pVTZ level of theory. The geometrical parameters at the equilibria and along the displacement are presented in Table 1. The total and elastic scattering signals are computed for each state. The signals are then expressed as a fractional intensity change as it is commonly done in experiments (see e.g. Refs. 1,5,8,9),

\[
\Delta S^\text{tot}(q, \mathbf{R}) = \frac{I^\text{tot}(q, \mathbf{R})_\text{exc} - I^\text{tot}(q, \mathbf{R})_\text{gs}}{I^\text{tot}(q, \mathbf{R})_\text{gs}},
\]

where \(I^\text{tot}(q, \mathbf{R})_\text{exc}\) and \(I^\text{tot}(q, \mathbf{R})_\text{gs}\) are the total scattering intensities for the excited state and the ground state, respectively. This expression gives the change of the signal for the excited state at a specific geometry, \(\mathbf{R}\), relative to the scattering from the ground state at its equilibrium geometry, \(\mathbf{R}_0\), under the assumption that there is no geometry change in the ground state upon excitation. In order to investigate the nature of fractional intensity change in further detail, it is conceptually useful to rewrite the \(\Delta S^\text{tot}(q, \mathbf{R})\) in Eq. (36) as sum of two contributions,

\[
\Delta S^\text{tot}(q, \mathbf{R}) = \Delta S^\text{tot}\text{elec}(q, \mathbf{R}) + \Delta S^\text{tot}\text{nucl}(q, \mathbf{R})
\]

\[
= \frac{I^\text{tot}(q, \mathbf{R})_\text{exc} - I^\text{tot}(q, \mathbf{R})_\text{gs}}{I^\text{tot}(q, \mathbf{R})_\text{gs}} + \frac{I^\text{tot}(q, \mathbf{R})_\text{gs} - I^\text{tot}(q, \mathbf{R})_\text{gs}}{I^\text{tot}(q, \mathbf{R})_\text{gs}}.
\]

The first term, henceforth called electronic, shows the difference solely due to the redistribution of the electrons at any given geometry. The second, nuclear term indicates the contribution due to structural changes and is defined with respect to the electronic structure of the ground state only. Furthermore, making use of the fact that the total scattering is a sum of elastic and inelastic scattering, each of these terms can be split into two contribu-
Figure 4: Total, elastic and inelastic ground-state x-ray scattering curves of NH₃. The scattering curves are calculated with CASSCF(10,8)/aug-cc-pVTZ. The results are compared with previous calculations by Hoffmeyer et al., who used an MR-SDCI wavefunction ([5s3p2d/3s2p]) that was numerically integrated on a grid. Subfigure 4a shows the total intensity, while subfigure 4b shows the difference.

\[
\Delta S_{\text{tot}}(q, \mathbf{R}) = \Delta S_{\text{elec}}(q, \mathbf{R}) + \Delta S_{\text{in}}(q, \mathbf{R}) \\
\Delta S_{\text{nucl}}(q, \mathbf{R}) = \Delta S_{\text{elec}}(q, \mathbf{R}) + \Delta S_{\text{in}}(q, \mathbf{R}).
\]  

The breakdown of the total signal in terms of these four components is given in Fig. 5. The umbrella motion is tracked from 0 to 33.0 degrees in the pyramidization angle formed between the plane of the hydrogen atoms and one of the N–H bonds. The displacement along the normal mode is accompanied by a N–H bond elongation from 1.03 Å to 1.23 Å, which is seen to be the dominant factor for the changes in the nuclear part of the signal. The ground state equilibrium geometry used in this work has a pyramidization angle of 23.5 degrees and an N–H bond length of 1.02 Å. With respect to the N–H distance that dominates the scattering, the planar equilibrium structure of the excited state is thus more similar to the geometry of the ground state than to the bent geometry of the excited state. This is reflected by the small magnitude of the nuclear scattering in Fig. 5a for this geometry. As the pyramidization angle increases, so does the bond length, which ultimately results in a maximum amplitude of \%\Delta S(q) of about 14%.

Meanwhile, the elastic part of the electronic contribution in Fig. 5b shows much less variability with the change of the geometry. The overall shape seems to change very little and the maximum amplitude of the central peak ranges only from 7% to 13%. It is of paramount importance to point out that the magnitude of the electronic component is comparable to that attributed to the nuclear motion. While at large pyramidization angles, the nuclear contribution is approximately two times larger, at near-planar geometries the elastic signal is almost exclusively linked to the redistribution of the electron density in the Rydberg state. Given the range of momentum transfer for which the dip in Fig. 5b occurs, the observation can be explained by the effective loss of electron density in the molecular core associated with the delocalization of the Rydberg electron in the excited state. The small changes in the electronic component along the umbrella mode align with the fact that Rydberg electrons are not strongly affected by structural evolution of the ion-like core.

Inspection of Figs. 5c and 5d reveals that the magnitude of the inelastic fractional difference scattering signal is about 10 times smaller than the corresponding elastic contribution. Nonetheless, the size of the inelastic contribution is clearly large enough to have a tangible effect on the interpretation of experimental data. Here, the patterns parallel those seen in the elastic scattering. The electronic contribution is relatively constant with a magnitude comparable to the nuclear effect. The
largest nuclear and the smallest electronic components are observed at the largest pyramidization angle, with about twofold difference between the two. The planar $\text{NH}_3$ is dominated by the electronic scattering. Interestingly, the geometry dependence of the inelastic scattering is usually not accounted for in experiments. Specifically, the inelastic scattering is often approximated as an incoherent sum of inelastic Compton factors for individual atoms. This is clearly a poor approximation in the case at hand and most likely in general when small effects in the fractional signal change in time-resolved gas-phase experiments are considered.\textsuperscript{25,38,39} Given that the experimental observable is the total scattering that stems from the Fourier transform of the two-electron electron density, it seems natural to consider the inelastic effect on an equal footing with elastic scattering. As seen here, inelastic scattering can account for up to 10% of the fractional signal change partially attributed to the difference between the two states considered and exhibits a geometry-dependence similar to elastic scattering.

When pumped into the excited state, ammonia undergoes fast umbrella motion. The picture that emerges from this work is that the observed signal will alternate between two extrema driven by the elongation and contraction of the N–H distance. However, as shown in Fig. 5e, the baseline for this oscillation is set by the shape of the relatively constant difference in the scattering signals between the Rydberg state and the ground state. The latter is attributed to the electron-density hole brought about by the promotion of an electron from the HOMO to the diffuse 3s Rydberg orbital. The inelastic scattering has smaller but far from negligible effect on the scattering signal.

### 5 Conclusions

The mathematical framework and computational approach presented in this article allows for an efficient calculation of the isotropic total, elastic, inelastic, and coherent mixed scattering signals. In order to perform the integration over the Euler angles needed to achieve spherical averaging, we consider \textit{ab initio} wavefunctions expressed in a basis of Gaussian-type orbitals. Analytic solutions to the Fourier transform from real to reciprocal space results in a series of products of Gaussian and spherical Bessel functions, which are relatively easy to evaluate computationally. The approach is benchmarked against previous numerical calculations in the case of ammonia. We demonstrate the scaling of the algorithms with the basis sets and levels of theory used.

The methodology described in the paper is utilized to investigate a simplified model of the photoexcitation of ammonia to its 3s Rydberg state. The observed elastic fractional difference scattering signal shows a strong signature of the shift in the electron density associated with the promotion of an electron from the HOMO to the 3s Rydberg orbital. The magnitude of this purely electronic effect is comparable to the geometry dependent part of the signal. In addition, ammonia shows strong change in the inelastic scattering upon excitation, which is driven by changes in the electronic structure. It is furthermore shown that changes in geometry also play a role in inelastic scattering.

As ultrafast gas-phase x-ray scattering experiments are becoming more and more successful in obtaining high-quality data, it is of paramount importance to have the right tools to analyze the results. The internal dynamics, both nuclear and electronic, is encoded in the isotropic part of the signal, which can be extracted by means of a Legendre decomposition of the detector signal. The isotropic signal should be understood as the spherical average of the Fourier transformed two-electron density of the molecule. The approach presented here allows this signal to be calculated efficiently for ground and excited states and can be used to aid the interpretation of pump-probe ultrafast x-ray scattering experiments.

Going further, a similar mathematical apparatus can be applied to the case of static molecules or any higher-order terms in the Legendre decomposition of the signal. The former can be achieved readily by forgoing the Spherical Bessel function expansion, while the latter can be achieved, albeit in a less straightforward manner, by using higher-order spherical Bessel functions.

Equally important is the question of the \textit{ab initio} level of theory used to calculate the molecular wavefunctions. Given that the signal is related to the two-electron density, it can be expected that
the total scattering shows high sensitivity to electron correlation. Hence, a thorough investigation of the impact of post-Hartree-Fock methods is urgently needed.

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6 Supporting Information

Detailed derivation that shows the origin of the Legendre decomposition, tabulated raw benchmarking data used to generate Fig. 2 and Fig. 3 in the main text.

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Figure 5: Breakdown of the total percentage fractional intensity change along the umbrella mode in the 3s Rydberg state of ammonia relative to the ground state equilibrium geometry. The displacement along the normal mode is labeled in terms of the pyramidalization angle between an N–H bond and the plane of the three hydrogen atoms. The nuclear elastic (a) and nuclear inelastic (c) terms indicate the changes due to the geometry evolution in the elastic and inelastic scattering, respectively. Similarly, the electronic elastic (b) and inelastic (d) components show the changes in scattering as a result of the difference between the electronic structure of the ground and the excited state at each geometry. The total scattering, i.e. the sum of (a)–(d), is given in (e).
Graphical TOC Entry

![Graphical representation of a scientific diagram with labeled axes and a legend indicating an XFEL pulse.](image)