

### **PERSPECTIVE**

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# Quantum coherence in molecular photoionization

**Cite this:** *Phys. Chem. Chem. Phys.* 2022, **24**, 19673

Received 4th April 2022,

Accepted 1st August 2022 DOI: 10.1039/d2cp01562e

rsc.li/pccp

Marco Ruberti, 🕩 \*a Serguei Patchkovskiib and Vitali Averbukh 🕩 °

The study of onset and decay, as well as control of ultrafast quantum coherence in many-electron systems is in the focus of interest of attosecond physics. Interpretation of attosecond experiments detecting the ultrafast quantum coherence requires application of advanced theoretical and computational tools combining many-electron theory, description of the electronic continuum, including in the strong laser field scenario, as well as nuclear dynamics theory. This perspective reviews the recent theoretical advances in understanding the attosecond dynamics of quantum coherence in photoionized molecular systems and outlines possible future directions of theoretical and experimental study of coherence and entanglement in the attosecond regime.

The process of photoionization is a perfect example of breakup of a quantum system into a series of distinct sub-systems, namely the photoelectron drifting away and the parent ion that is left behind. In the case of atomic photoionization, the remaining parent ion can in general consist of a bound or highly-excited many-electron system, while in the molecular case additional breakup of the nuclear quantum system, such as fragmentation, can also be triggered during the process.

# 1 Ionic coherence: formal theory

Two interrelated, fundamental quantum-mechanical concepts characterize the many-body dynamics that is triggered by photoionization: quantum coherence and quantum entanglement. From a quantum mechanical point of view, maximal quantum electronic coherence within each of the photoelectron and parention sub-systems that are formed upon photoionization implies that, at large separations between the two, the total, timedependent, N-electron wavefunction of the photoionized system  $\Psi^N(x_1,\ldots,x_i,\ldots,x_N;R,t)$  is separable in the degrees of freedom of the two formed sub-systems. Note that here the parameter R stands for the nuclear coordinates. (When nuclear dynamics are relevant, R becomes a dynamical variable as well, on equal footing with  $x_i$ . This case will be considered below, in Section 4.) Therefore, the wavefunction of the composite N-electron system can be written, at least to a good approximation, as the anti-symmetrized product of a (N - 1)-electron wavefunction (in principle fully correlated)

$$\Psi^{N}(x_{1},..,x_{i},..,x_{N};R,t)$$

$$\approx \hat{A}\left[\chi^{N-1}(x_{1},..,x_{i},..,x_{N-1};R,t)\times\psi^{e^{-}}(x_{N};t)\right].$$
(1)

However, the product Ansatz of eqn (1) is valid only if the photoelectron and the parent-ion systems are not entangled and, as a result, the quantum coherence is at maximum within each subsystem. In fact, the total bipartite system is in general described by a time-dependent density matrix  $\hat{\rho}(t)$ . Each of the separate sub-systems is itself described by a reduced density matrix that is obtained by tracing out the extra degrees of freedom pertaining to the other, unobserved, sub-system. In the case of the parent ion, the so-called reduced ionic density matrix (R-IDM) reads as

$$\hat{\rho}^{\text{R-IDM}}(t) = \text{Tr}^{e^{-}}(\hat{\rho}(t)), \tag{2}$$

where the partial trace has been performed on the degrees of freedom representing the photoelectron. This is meaningful only for a total density matrix  $\hat{\rho}(t)$  that actually describes a photoionized system, *i.e.* where, at least at times t when the photoelectron is sufficiently far away from the parent-ion, one of the electronic degrees of freedom (the one that will be identified as pertaining to the photoelectron) is associated to single-electron states that are physically distinguishable (e.g. for energy/momentum and spatial separation) from the ones occupied by the remaining (N-1) electrons.

Even in the case where the total density matrix  $\hat{\rho}(t)$  corresponds to a pure state, *i.e.*  $\hat{\rho}(t) = |\Psi^N\rangle\langle\Psi^N|$ , the partial trace operation gives rise in general to a mixed quantum state for the resulting sub-system. The operator of eqn (2) can be projected onto a basis of many-electron states of the parent-ion. For

describing the parent-ion, *i.e.* only dependent on ionic degrees of freedom, and a one-electron wavefunction describing the photo-electron

<sup>&</sup>lt;sup>a</sup> Imperial College London, Department of Physics, South Kensington Campus, London SW7 2AZ, UK. E-mail: m.ruberti11@imperial.ac.uk

b Max-Born Institute, Max-Born-Straße 2A, 12489 Berlin, Germany. E-mail: Serguei.Patchkovskii@mbi-berlin.de

<sup>&</sup>lt;sup>c</sup> Imperial College London, Department of Physics, South Kensington Campus, London SW7 2AZ, UK. E-mail: v.averbukh@imperial.ac.uk

example, in the basis of eigenstates of the ionic Hamiltonian  $|E_I^{N-1}\rangle$  we obtain:

$$\hat{\rho}^{\text{R-IDM}}(t) = \sum_{I,J} \rho_{I,J}^{\text{R-IDM}}(R,t) |E_I^{N-1}(R)\rangle \langle E_J^{N-1}(R)|. \tag{3}$$

In the rest of this section the parameter R will be omitted. (We will return to it in Section 4.) The time-dependent populations  $P_I$  of each ionic eigenstate  $|E_I^{N-1}\rangle$  are given by the corresponding diagonal matrix elements,  $P_I(t) = \rho_{I,I}^{\text{R-IDM}}(t)$ , while the time-dependent degrees of coherence  $G_{I,J}$  and relative phases  $\phi_{IJ}$  between any pair of such states are given by

$$G_{I,J}(t) = \frac{|\rho_{I,J}^{\text{R-IDM}}(t)|}{\sqrt{P_I(t) \times P_J(t)}}$$
(4)

and

$$\phi_{I,J}(t) = \arg(\rho_{I,J}^{\text{R-IDM}}(t)), \tag{5}$$

respectively.

Spectral decomposition of the R-IDM

$$\hat{\rho}^{\text{R-IDM}}(t) = \sum_{I} r_I(t) |C_I^{N-1}(t)\rangle \langle C_I^{N-1}(t)|.$$
 (6)

allows one to express the mixed quantum state of the parent-ion system, at each time t, as an ensemble of orthogonal pure quantum states, each populated with weight  $r_I(t)$  and with the correspondent projection operator given by  $|C_I^{N-1}(t)\rangle\langle C_I^{N-1}(t)|$ . The global quantum coherence of the mixed quantum state of the ionic system can be quantified by the so-called ionic purity  $p(t) = \text{Tr}(\hat{\rho}^{\text{R-IDM}}(t) \times \hat{\rho}^{\text{R-IDM}}(t))$ , while the entanglement between the parent-ion and the photoelectron can be measured by the von Neumann entropy of entanglement  $s(t) = -\sum_{i} r_{I}(t) \times r_{I}(t)$ 

 $ln(r_I(t))$ . The more entangled the two subsystems are, the more mixed (i.e. with less quantum coherence) the state of each of them is. This can be easily seen by comparing eqn (1) and (6): maximal coherence requires the parent-ion to be described by a pure state, *i.e.* by a well-defined (N-1)-electron wavefunction, which can be expressed in any arbitrary basis as a specific fully coherent linear superposition of the basis states. Therefore, the spectral decomposition of eqn (6) must only have one  $r_i$  term different from zero (and consequently equal to 1). This requirement automatically gives us a minimum value for the entropy of entanglement, equal to 0, and a maximum value for the purity, equal to 1. Finally, it is important to note that both of these "global" quantities are independent of the specific basis set used.

### 2 Coherence in attosecond science

In the last four decades, traditional synchrotron photoionization experiments have allowed the community to address the properties of individual, incoherently populated eigenstates of the ionized system. Indeed, a narrow-bandwidth synchrotron radiation couples each molecular ionized state to an outgoing electron wave of well-defined energy. Thus, the total wavefunction of the photoionized N-electron system is highly entangled and cannot be represented as a separable product of ionic and free-electron states.

The alternative large-bandwidth photoionising laser sources based on the high-order harmonic generation (HHG) and X-ray free electron laser (FEL) technologies have been actively developed over the recent two decades, however until recently they have been suffering from drawbacks that hindered their use in exploring the onset and the effects of the ionic coherence. While the attosecond HHG sources have the required bandwidth for coherently exciting series of atomic or molecular ionic states, they lacked the intensity required for both creating and probing the excited coherent superpositions. On the other hand, the much more intense X-ray FEL sources lacked the bandwidth required to create the ionic coherence across, e.g. 10 eV energy range. Recently, the spectacular developments of both HHG sources2 and X-ray FELs3 has led to the emergence of newgeneration attosecond light pulses which can coherently excite and probe a system of interest over a broad range of photon energies. These pulses allow one to produce the quantum electronic coherences that characterize the partially-entangled state of the ionized electron and parent-ion sub-systems, as well as to explore new, spectacular physical effects which result from the non-stationary quantum superpositions of ionized atomic and molecular states, such as molecular hole migration.<sup>4</sup>

It is the set of quantities from eqn (4) and (5) that determine the ultrafast correlated dynamics that is triggered by the photoionization process and that eventually couple with the adiabatic and non-adiabatic nuclear dynamics in the ionic system. One particular example is the so-called charge migration which results from electronic correlation and assumes a coherent superposition of all the states resulting from removal of an electron from a specific hole orbital. The partially coherent superposition of such molecular ionic eigenstates can be accompanied by an ultrafast electron dynamics which has been predicted to feature a migration of the positive charge across the molecular-ion backbone, on a sub- to few-femtosecond (fs) timescale (hole migration).4 On longer timescales, this purely electronic coherent dynamics is potentially subject to decoherence<sup>5,6</sup> due to coupling to the slower nuclear motion degree of freedom, which acts as a bath and could eventually lead to the final localization of the positive charge.

At the heart of modern attosecond science is the study of the coherent many-electron dynamics resulting from photoionization by means of attosecond pump-probe spectroscopy techniques.<sup>7,8</sup> Since molecular bonds are more likely to break in the regions with higher density of the electronic hole, ultrafast charge redistribution underpinned by electronic coherence in the cation leads the way to photoionization-induced, charge-directed reactivity on attosecond timescales. In turn, this opens up the possibility of steering the molecular reaction dynamics at a very early stage of its quantum evolution by using the electronic degrees of freedom before the dephasing occurs. Exploiting the quantum electronic coherences, rather than just the eigenstates populations, provides a novel opportunity to control photochemical reactions, previously unavailable on the much longer (femto/pico-second) timescales intrinsic to nuclear dynamics. Moreover, since electronic redistribution

initiates all photochemical change,<sup>9</sup> the study of quantum electronic coherence and entanglement in photoionization is also of fundamental interest for understanding the basic processes eventually leading to photosynthesis and radiation damage. Here it is important to note that, although natural photosynthetic processes are triggered by incoherent light, ultrafast pump-probe techniques based on fully-coherent light sources represent a strong tool to investigate the evolution of quantum coherence and energy transport in photosynthetic systems and shed light on their fundamental light-harvesting mechanisms.

The goals of the attosecond community can be briefly summarized as follows:

- Prepare ultrafast quantum coherence.
- Probe ultrafast quantum coherence.
- Control ultrafast quantum coherence.

In order to realize the full potential impact of attosecond science, a close integration of theoretical and experimental efforts is crucial. On the one hand, attosecond experiments frequently ionize the system, leading to an entangled ion-photoelectron pair. The resulting entanglement limits the coherence that we can observe when we perform measurements on each of the two sub-systems. On the other hand, attosecond experiments, that investigate the type of ultrafast dynamics that can be triggered in the parent ion, rely on the existence of coherence in the latter sub-system, i.e. a welldefined phase relationship (see eqn (4) and (5)) between the various components of the quantum state that can give rise to a timedependent interference pattern when an observable sensitive to the coherences is measured in the laboratory. Because of the complexity of the photoionization dynamics involved, the capability to model laser-induced many-electron processes from first principles is absolutely key to predict new physical phenomena, and to guide experimental efforts towards their observation and characterization.

In addition, theoretical prediction of quantum coherences in photoionization is essential not only for our fundamental understanding of the physics underlying photochemical transformations at ultrashort timescales, but also for applying concepts of quantum information (QI) to the field of ultrafast dynamics. Most importantly, doing so can provide us with new routes to control photo-physical/chemical processes, i.e. it opens the way to harnessing these fundamental, inherently quantum mechanical concepts in order to develop control strategies of chemical reactivity of photoionized molecules.

To date, a full quantum information perspective on attoscience is still missing and the role of electron nuclear coupling in this respect is thus far mostly unexplored. There are many open questions yet to be answered including: how would an interaction with, or a measurement performed on, the photoelectron affect or correlate, respectively, with the chemical process in the ion that this electron left behind?

### 3 Theoretical methodology and applications

All this has called for the development, in the last decade, of new theoretical and computational methodologies capable of answering fundamental questions on coherence and entanglement in ultrafast photoionization dynamics. In general, in order to calculate the ionization-induced quantum electronic coherences, it is essential for these methodologies to explicitly describe the photoionization process and preferably include in the theoretical description the following required "ingredients":

• A complete description of the full N-electron quantum system, starting from the initial state of the neutral molecule, and capturing the entanglement between the (N-1)-electron ion and the photoelectron after the photoionization process. This must be done by tracking the time evolution of the N-electron wavefunction, whose ionized part can be written in general as

$$|\Psi^{N}(t)\rangle = \sum_{j} \sum_{a} C_{j,a}(t) |I_{j}e_{a}^{-}\rangle,$$
 (7)

where  $|I_i e_a^-\rangle$  is a product state of stationary eigenstates of the ionic  $|I_i\rangle$  and photoelectron  $|e_a^-\rangle$  sub-systems, with indices j, k,  $l, \dots$  and  $a, b, c, \dots$  for their quantum numbers, respectively.

- A time-dependent description of the photoionization process in order to capture the ultrafast formation and loss of quantum coherence.
- An accurate prediction of the ionization quantum amplitudes corresponding to each of the (most physically-relevant) many-electron states of the parent ion (ionization channels), as a function of the parameters (e.g. frequency and intensity) of the ionizing laser pulse.
- Accounting for the effect of the interchannel couplings between the emitted electron and the parent ion, i.e. for correlation effects in the continuum. This type of residual interaction between the parent-ion and the photoelectron is different with respect to the so-called intrachannel coupling, where the photoelectron experiences the potential created by the charge distribution of the specific fixed cationic state, insofar as the state of the parent-ion is also allowed to change as a result of it. It is important to note that it is the interchannel coupling term that causes the loss of coherence in the parention during the departure of the emitted electron.

From a theoretical point of view, the major challenge in modelling the formation of ionic coherence in the process of attosecond and strong-field ionization of a molecular system consists of solving the multi-centre (with respect to the singlecentre atomic case), multi-channel, many-electron problem in the presence of a laser field. This is key to revealing the physical mechanisms contributing to the formation of the cationic wavefunction through the external field and electron correlation, and needs to be done across a wide energy window, which is defined by the interaction with the ionising pulse, and which leads one to face several computational difficulties. Here we refer to the electron correlation, as standard in quantum chemistry, as to the effect of electron-electron repulsion that is not accounted for within the Hartree-Fock (self-consistent field) approximation.10 First, standard methods of electronic structure calculation are poorly adapted to the description of continuum electrons. Second, attosecond (HHG and pumpprobe) experiments typically involve tunneling/multi-photon

ionization as well as (sequential) double ionization, which are even more difficult to take into account. Finally, molecular systems are characterized by a larger size, a higher number of degrees of freedom and, in general, a lower symmetry with respect to atoms, which poses additional demands on the computation. This is why, until recently, the density matrix characterising the ionic state emerging from attosecond ionization could not be predicted theoretically or characterized experimentally in the general molecular case - but only in some atomic cases, 11,12 for which a very recent experimental work has also reported the experimental reconstruction of the density matrix of the photoelectron sub-system.13

#### 3.1 Low-frequency strong-field regime

In recent years, simulation of coherence formation in infrared (IR)-driven strong-field ionization (SFI) has been performed, by solving the time-dependent Schrodinger equation (TDSE), both for atomic systems by the group of Santra et al.,14 using the time-dependent (TD) configuration interaction singles (CIS) method for electronic structure, and for molecular systems by Ruberti and Averbukh<sup>15</sup> using the first-order version of TD B-spline algebraic diagrammatic construction (ADC) method, i.e. TD B-spline ADC(1).

In the low-frequency, SFI regime, it is possible to identify at least three different physical mechanisms that, interplaying with each other, contribute to the formation and destruction of quantum electronic coherence in the parent ion system during the photoionization process. These are:

• The presence of distinct, single-channel, laser-driven nonperturbative transitions (multi-photon or tunnel) that can excite an electron to the same continuum state and, at the same time, leave the parent-ion system in different (j, k)internal states. The Hamiltonian terms responsible for this process are:

$$\langle \Psi_0^N | \hat{\mu} | I_{j/k} \mathbf{e}_a^- \rangle \neq 0$$
 and  $\langle I_{j/k} \mathbf{e}_a^- | \hat{\mu} | I_{j/k} \mathbf{e}_b^- \rangle \neq 0$ , (8)

where  $\hat{\mu}$  is the electric dipole operator. In a simple orbital picture, these mechanisms correspond to a non-zero amplitude for an electron to be excited, by the electric field of the laser pulse, from different, initially occupied (hole) orbitals into the same final orbital in the continuum of its energy spectrum. The terms in eqn (8) are Hamiltonian terms, that are independent of the intensity of the laser field. However, what does strongly depend on the strength of the laser field is what type of dynamics and transitions these Hamiltonian matrix elements lead to: non-perturbative multi-photon or tunnelling ionization in the strong-field regime, perturbative single to a few-photon transitions in the weak field regime.

 Rabi-like dynamics in the parent-ion: resonant or nearlyresonant laser-driven transitions between different states of the parent-ion that are coupled to each other by an electric dipole matrix element:

$$\delta_{a,b}\langle I_i|\hat{\mu}|I_k\rangle \neq 0. \tag{9}$$

In this particular mechanism, the photoelectron acts as a spectator and does not change its state during the rearrangement of the electronic cloud within the ionic system (note the  $\delta_{a,b}$  in the Hamiltonian term of eqn (9)). Therefore, a quantum amplitude referred to a fixed photoelectron state is transferred between the ionic states involved, changing the coherence between them. Note that this mechanism is not present for weak-field single-photon ionization in the XUV/X-ray regime, since the energy of the photon is far off-resonance with the bound-bound ionic transitions and the field intensity is low.

 Residual Coulomb interaction between the parent-ion and the photoelectron as the latter is still in the proximity of the former. In particular, the interaction of the liberated electron with the ionic hole state, from which it originates, is referred to as intrachannel interaction and leads, for large electron-ion distances, to the 1/r behaviour typical of the Coulomb potential. If the liberated electron is influenced by other molecular orbitals (ionic states) the interaction is referred to as interchannel coupling:

$$\langle I_i \mathbf{e}_a^- | \hat{V} | I_k \mathbf{e}_b^- \rangle \neq 0, \tag{10}$$

where  $\hat{V}$  represents the direct and exchange terms of the twoparticle Coulomb interaction. It is the latter scenario that can produce transitions in the parent-ion and directly impact on the coherence between the ionic states involved.

A schematic representation of these mechanisms can be seen in Fig. 1. An important feature of SFI is the strong (exponential) dependence of the ionization amplitude for a specific ionic state on the value of its ionization potential. As a result, during this ionization process, only the first few lowest-excited states of the parent-ion are effectively populated. In small to medium-sized molecules, these states are characterized by a relatively small amount of electron correlation and, as a consequence, they are reasonably well described within the Koopman's picture of ionization, 10 according to which singlyionized states are obtained by removing an electron from one of the occupied orbitals in the Hartree-Fock (HF) ground state of the neutral molecule.

Therefore, in the long-wavelength, strong-field regime, the complexity given by the non-perturbative nature of the lasermolecule interaction is tempered by the fact that the modelling of the electronic structure in the outer-valence energy region of the spectrum can often be greatly simplified by adopting a single-hole description. Orbital relaxation and electron correlation in the strong-field regime can also be included without much difficulty, using a non-orthogonal, time-dependent resolution in ion states (TD-RIS) approach. 16,17

#### 3.2 High-frequency regime

The situation is radically different in the XUV and soft X-ray regimes, where the photon energy is such that more deeplybound electronic states, i.e. higher excited states of the parent ion, can be accessed by absorption of a single or a few photons. In spite of the perturbative nature of the photoionization process at these shorter wavelengths (at least for laser intensity of the order of 10<sup>15</sup> W cm<sup>-2</sup> or less), calculation of the quantum

(i) E<sub>Neutral</sub> (ii) (iii)

Fig. 1 A schematic representation of the main mechanisms of ionic coherence formation in photoionization: (i) direct single-channel ionization of the neutral ground state into different final ionic channels that share the same photoelectron state. The form of the transition operator  $\hat{T}$ depends on the (perturbative or non-perturbative) nature of the ionization process, i.e. on the laser intensity; (ii) laser-driven transitions between (mainly resonant or nearly-resonant) ionic states; (iii) residual interchannel interaction between the photoelectron and the parent ion.

electronic coherences in the ionic sub-system requires a more accurate treatment of electron correlation effects, which dominate in the inner-valence energy region of the ionic spectrum. Moreover, things are made more complicated also by the higher density of states in the region close to the double ionization potential (DIP) threshold and by the fact that each of these states of the parent-ion has to be coupled to a continuum of states for the photoelectron sub-system. All these aspects make the accurate prediction of the reduced ionic density matrix upon attosecond XUV ionization of a polyatomic molecule very challenging.

A typical electronic energy spectrum of a molecular ion in the fixed-nuclei approximation, which extends beyond the DIP threshold, is shown in Fig. 2. Each line in the spectrum represents an ionic eigenstate and is located at the corresponding ionization energy. The values of the spectral intensities reflect the contribution of the one-hole (1h) configurations to the ionic states. In the absence of electron correlation, i.e., within the range of validity of the Koopman's theorem, one indeed obtains a series of lines with a height of 1 for each occupied HF molecular orbital. This is often accurate in the outer-valence energy region, i.e. close to the first ionization potential of the system. As the energy of the ionic states increases the effect of electron correlation also becomes more important: this can either give rise to hole-mixing, where eigenstate are linear combinations of different 1h states, to the formation of shakeup satellites and ultimately, in the inner-valence energy region to the complete breakdown of the molecular orbital (MO) picture (see Fig. 2). In terms of electronic configurations, description of electron correlation in the modelling of the cation implies the inclusions of all the ionic two-hole-one-particle (2h-1p)

configurations. Within advanced descriptions, such as the one given by the second (ADC(2)x) and third (ADC(3)) order methods of the ADC hierarchy, 18 electron correlation in the description of the ionic system manifests itself in individual ionic states as a reduction of the spectral intensity line height (with the "missing" part reflecting the contribution of 2h1p configurations) and, related with the latter, as the appearance of additional (satellite) lines in the spectrum. A 2h1p configuration indeed describes the removal of an electron from a particular orbital accompanied by the excitation of another electron to an initially unoccupied orbital.

As a result, two more "ingredients" can be added to the previous list:

- Description of electron correlation, i.e. shakeup states, correlation satellites.
- Description of the electronic relaxation that happens due to the change in the mean-field potential that each electron experiences upon creation of a hole in a deeply-bound orbital.

Among the most promising new theoretical methods that have recently been developed to achieve this goal, we would highlight the time-dependent first-order perturbation theory, static-exchange density-functional-theory (DFT) method developed by P. Decleva and collaborators, tits multi-reference extension developed by Ponzi et al. 19 and applied to attosecond molecular ionization dynamics by Martin and collaborators, 20 the XCHEM method by Martin and collaborators, 21 the realtime non-equilibrium Green's function methods (NEGF) as implemented in the CHEERS code by Perfetto et al.,22 the molecular R-matrix with-time approach by Benda et al., 23 the TD B-spline ADC method by Ruberti and Averbukh, 24,25 and finally the TD B-spline restricted correlation space (RCS)-ADC by Ruberti. 26,27 For a more comprehensive list of such methods we refer the reader to ref. 28.

In particular, the TD B-spline ADC method extends the ab initio ADC approach, originally developed to describe bound state dynamics, 18 to the realm of ultrafast ionization dynamics, combining the accurate description of electron correlation of quantum chemistry with the full account of the continuum dynamics of the photoelectron, while the RCS technique<sup>26</sup> crucially turns first-principles modeling of coherence and entanglement in photoionization of polyatomic molecular systems into a tractable problem. RCS-ADC is based on the separation of the orbital space into correlation and ionization spaces<sup>26</sup> and naturally bridges the gap between multiconfigurational ab initio techniques and closed-coupling schemes based on a limited number of essential, physically relevant ionic states, combining the key advantages of both.

Approaches of this kind allow one to accurately predict the mixed state of the ionized system prepared by attosecond ionization of a molecular system.<sup>27</sup> Within TD B-spline RCS-ADC, complete theoretical characterization of the atto-ionized many-electron state and photo-induced attosecond charge dynamics is achieved by calculating the reduced ionic density matrix (R-IDM) of the bipartite ion-photoelectron system, with full inclusion of the correlated ionic shakeup states, and its interaction with yet another ionising laser pulse probing its dynamics.29

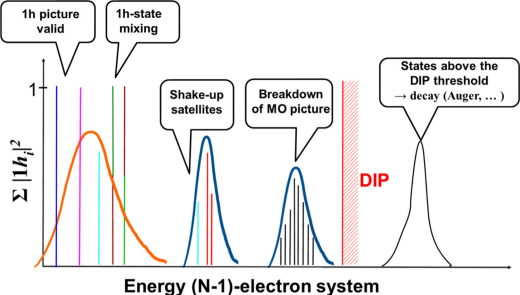


Fig. 2 Schematic representation of the typical structure of the electronic energy spectrum in a photoionized poly-atomic molecule. Each vertical line corresponds to an energy eigenstate of the cationic system; the height of each line corresponds to the so-called spectral intensity of the state, i.e. the sum of its squared coefficients on all the possible  $1h_i$  configurations,  $|1h_i|^2$ . Contributions from different 1h configurations are shown with different colours. Typical energy spectra of ultrashort pulses capable of coherently excite states in the outer-valence (orange curve) and inner-valence (blue curves) energy region are also shown.

The power of this approach is apparent in the Schmidt decomposition of the R-IDM. In fact, the latter unveils the many-electron dynamics triggered by the pump and allows for the identification of the key pure-state channels involved in the quantum coherent many-electron dynamics.<sup>29</sup> Most importantly, the methodology can be applied to the simulation of complete numerical pump-probe experiments of ultrafast dynamics in molecular systems. 15,29,30 This is because the hierarchies of both ADC and RCS-ADC methods extend across different ionization stages.

Another wavefunction method is XCHEM, 21 which uses a Close-Coupling approach based on a Restricted-Active-Space Self-Consistent-Field (RAS-SCF) description of electron correlation and on the use of a mixed basis set comprising both Gaussian type orbitals and B-splines. This method, which, as well as the ADC-based methods, includes a description of electron correlation effects in the continuum, has so far been applied to the timeindependent description of photoionization in small molecules.<sup>31</sup> The static-exchange DFT method<sup>7</sup> provides an effective singleparticle description of the photoionization dynamics, as well as a more basic description of electron correlation effects through the use of exchange-correlation functionals in the calculation of the ground-state Kohn-Sham (KS) orbitals. The multi-reference static-exchange scattering method<sup>19</sup> improves the description of electron correlation with respect to ref. 7 by considering the multi-configurational character of the parent-ion wavefunction. However, both methods neglect interchannel couplings in the electronic continuum and their first-order perturbation theory implementation is limited to description of weak-field, singlephoton ionization processes. Finally, NEGF theory allows one to describe the many-body problem from first principles by

using effectively-single-particle approaches.<sup>22</sup> Dynamical correlation effects are added to the Hartree-Fock dynamics through a many-body self-energy. The real-time implementation of the method is potentially suitable for applications to describe nonperturbative ionization dynamics. The description of the photoelectron system is limited by the use of KS continuum states a in planewave or grid basis set.

Within single-photon absorption, coherence between different single holes in the outer-valence part of the spectrum, where the energy separation between the different ionic states can be of the order of a few eV, typically requires ultra-short, attosecond laser pulses with a time duration of the order of hundreds of attoseconds.

In the inner-valence energy region, longer pulses in the few-femtoseconds range can already be capable of exciting coherently either a single hole and its shakeup satellite states, or a series of correlated states in a breakdown of the molecular orbital (MO) picture scenario (see Fig. 2). This is because of the higher density of states in the inner-valence energy region. A particular type of coherent excitation is the one pictured by the so-called sudden ionization approximation, whose validity strictly requires the photon energy to be much higher than the ionization thresholds of the system. In the sudden ionization picture, a high frequency pulse "suddenly" removes an electron from one occupied orbital in the inner-valence region. As a consequence of electron correlation, this implies the coherent population of each of the ionic eigenstates with a non-zero amplitude on the 1h configuration corresponding to the created hole.

Finally, it is worth noting that the creation of quantum coherence between states lying above the DIP can give rise to

coherent dynamics super-imposed on the non-radiative (Auger)

decay of such states, a phenomenon that we can call "Auger

In perturbative XUV ionization, direct (in this case mainly single-photon) transitions into the same continuum state and inter-channel couplings (see panel (i) and (iii) of Fig. 1, respectively) are the dominating mechanisms that contribute to the build-up of the ionic coherence. The latter always leads to an overall decrease of ionic purity and a corresponding increase of the entropy of entanglement. Increasing the intensity can also potentially introduce new mechanisms of coherence formation in the XUV regime: these include not only the dipole transitions between the populated ionic states and multi-photon ionization, but also Raman-type transitions mediated by the core-ionized states and/or the continuum.

Finally, in order to achieve a complete theoretical description of the quantum coherent dynamics triggered during the photoionization of a molecular system, the last key "ingredient" is:

 Description of the coupling between the electronic and nuclear dynamics. This is in principle a very challenging task, because the possible existence of several electronic coherences among excited electronic states requires one to model nonadiabatic nuclear dynamics across the several, correspondent, potential energy surfaces.

Before proceeding, in the next section, to discuss the correlations between electronic and nuclear degrees of freedom, we would like to remark that a deeper understanding of the mechanisms that characterize the dynamics of the quantum attosecond coherences is also key to the interpretation and the design of new experimental protocols aiming at probing such coherences, such as single-photon laser enabled Auger decay (spLEAD), 32,33 X-ray attosecond transient absorption and X-ray time-resolved Auger and photoelectron spectroscopies. 30,34 Crucial features, which are highly desirable for such new schemes, are, on the one hand, the chemical-site and ionic-state selectivity and, on the other hand, extremely high temporal resolutions up to the attosecond regime.

### 4 Correlations between electronic and nuclear degrees of freedom in ionization

Electronic-state coherences, discussed in the preceding section, are the most important, but not the only coherence effects relevant to attosecond and strong-field photoionization. All isolated systems possess the centre-of-mass degrees of freedom, as well as electronic- and nuclear-spin degrees of freedom. Molecules additionally possess rotational and vibrational degrees of freedom. Potentially, all of these degrees of freedom can become correlated to the electronic degrees of freedom, leading to non-trivial dynamical and spectroscopic consequences.

Here, we will concentrate on the vibrational motion, which occurs on the time scale closest to that of the electron dynamics. We will however note in passing that correlations between the centre-of-mass motion and ultrafast electronic dynamics, which are routinely neglected in the attosecond domain, can offer a sensitive, non-destructive probe of the electron dynamics.35,36

#### 4.1 Vibrational dynamics

In order to qualitatively understand the origin of the nuclearmotion effects in broad-band, coherent photoelectron spectra, it is helpful to adopt the time-dependent perspective. A delightful exposition of these ideas can be found in a classic review by Heller. 37 The simplest-possible, near-trivial example is one-photon ionization, which we find instructive to explore in some detail. We choose to closely follow the textbook by Tannor, 38 however casting the expressions in the S-matrix form commonly used in analytical strong-field applications. 39,40 Unless mentioned otherwise, we use Hartree atomic units  $m_e = |e| = \hbar = 1$  below.

Because we are now interested in correlations between electronic and nuclear dynamics, it is necessary to venture into the realm of non-adiabatic vibronic dynamics (see e.g. ref. 41-44). For our purposes, it is sufficient to work within the Born-Huang Ansatz<sup>45</sup> and to ignore the challenges<sup>42,46</sup> arising due to state intersections. A more careful treatment is necessary in the general case, which however would be out of scope of this perspective. Some of the possibilities would include using diabatic representation,43 the "complete adiabatic" Ansatz,47,48 and the related exact-factorization approach. 49

The lowest-order total (electronic and nuclear) wavefunction response to a time-dependent perturbation  $\hat{V}(t)$  is given by:<sup>38,40,50</sup>

$$\left| \Psi^{(1)}(t); \hat{V} \right\rangle = -i \int_{t_0}^{t} \mathrm{d}t' \, \hat{U}_T(t, t') \hat{V}(t') \hat{U}_0(t', t_0) |\Psi(t_0)\rangle. \tag{11}$$

The initial time  $t_0$  is chosen such that  $\hat{V}$  vanishes at all earlier times. The propagators  $\hat{U}_0$  and  $\hat{U}_T$  are symbolically represented by exponentiation of the corresponding Hamiltonians  $\hat{H}_0$  and  $\hat{H}_T = \hat{H}_0 + \hat{V}$  (for brevity, we choose to omit the time index where it is unambiguous from the context):

$$\hat{U}_a(t_2, t_1) = e^{-i \int_{t_1}^{t_2} dt'' \hat{H}_a}, \tag{12}$$

where a = 0, T.

The general eqn (11) can be reduced to a computationally tractable form by introducing a sequence of approximation, as discussed in detail elsewhere. 38,50,51 Briefly, by choosing the energy origin at the energy of the initial state  $\Psi$ , which is assumed to be an eigenfunction of  $\hat{H}_0$ , and using the Born– Oppenheimer factorization for the initial state, we obtain:

$$\hat{U}_0(t',t_0)|\Psi(t_0)\rangle = |\psi_i\rangle|\chi_i\rangle,$$

where the time-independent initial electronic wavefunction  $|\psi_i\rangle$ depends at most parametrically on the nuclear coordinates, and  $|\chi_i\rangle$  is the initial, time-independent nuclear wavefunction. For one-photon ionization or excitation, operator  $\hat{V}$  can normally be treated within the dipole approximation:

$$\hat{V}(t') = \frac{1}{2}\hat{\mu}e^{-i\omega t'}f(t'),$$

where  $\omega$  is the carrier frequency, f(t) is a slowly-varying

envelope, the counter-rotating, stimulated-emission term  $(\propto e^{+i\omega t'})$  has been dropped, and operator  $\hat{\mu}$  absorbs the field polarization properties and the choice of the interaction gauge. Finally, approximating  $\hat{U}_T$  by  $\hat{U}_0$ , neglecting non-adiabatic contributions  $^{42,43}$  to  $\hat{H}_0$ , projecting  $\Psi^{(1)}(t)$  onto a time-independent target electronic state  $\psi_f$ , and rearranging the terms, we recover the expected  $^{38,50}$  result for the vibronic wave-packet propagating on the electronic potential energy surface f:

$$\left|\Psi_{f}^{(1)}(t);\hat{V}\right\rangle = -\frac{i}{2}\left|\psi_{f}\right\rangle e^{-i\varepsilon_{f}t} \int_{t_{0}}^{t} dt' f(t') e^{i(\varepsilon_{f}-\omega)t'} \hat{u}_{f}(t,t') \mu_{fi}\chi_{i},\tag{13}$$

where  $\varepsilon_{fi}$  is (in principle arbitrary) "characteristic" electronic transition energy, and  $\mu_{fi}$  is the electronic transition dipole moment between surfaces i and f:

$$\mu_{fi}(R) = \langle \psi_f | \hat{\mu} | \psi_i \rangle. \tag{14}$$

Finally, the nuclear propagator  $\hat{u}_f$  on the electronic surface f is given by:

$$\hat{u}_f(t,t') = e^{-i\int_{t'}^t dt'' \hat{h}_f}$$
(15)

$$\hat{h}_f(R) = \int \! \mathrm{d}r \psi_f^* \hat{H}_0 \psi_i - \varepsilon_{fi} \tag{16}$$

Eqn (13) represents bound and dissociative nuclear wave-packet on an equal footing.<sup>37</sup> It is applicable to both bound-to-bound and ionizing electronic transitions. In the latter case, electronic dynamics in the continuum, which is the subject of the preceding sections, appears as an additional integration over the target states f. Furthermore, the general form of the eqn (13) is preserved for multiphoton and strong-field transitions,<sup>38</sup> although the definitions of the specific operators may need to be modified. It therefore forms the perfect departure point for the discussion below.

#### 4.2 Broadband, 1-photon ionization

We advance by examining the familiar example of 1-photon ionization. Except for the (presently immaterial) normalization choice of the target states, the same treatment applies to 1-photon excitation. Because we are ultimately interested in coherent electronic dynamics, we will consider a coherent, *two-colour* perturbation:

$$\hat{V}(t') = \sum_{j=1,2} \hat{V}_j(t)$$

$$\equiv \sum_{j=1,2} \frac{1}{2} \hat{\mu} e^{-i\omega_j t'} f_j(t').$$
(17)

(eqn (17), and the discussion below can be trivially generalized to a general coherent, broadband perturbation if desired). Perturbation (17) will produce a vibronic wavepacket:

$$|\Psi^{(1)}(t)\rangle = \sum_{f} \sum_{j=1,2} |\Psi_f^{(1)}(t); \hat{V}_j\rangle,$$
 (18)

where each  $|\Psi_f^{(1)}(t); \hat{V}_j\rangle$ , wavepacket component on electronic

surface f produced by perturbation component  $\hat{V}_j$ , is given by eqn (13).

Because we assume that only the electronic state of the system is observed, we need to form the reduced density matrix  $\hat{\rho}$  of the system, with the vibrational degrees of freedom traced out, exactly as we did for the photoelectron in eqn (3):

$$\hat{\rho}(t) = \sum_{ba} |\psi_a\rangle \rho_{ba}(t)\langle \psi_b|,\tag{19}$$

$$\rho_{ba}(t) = \sum_{j,k=1,2} \rho_{ba}^{jk}(t), \tag{20}$$

$$\rho_{ba}^{jk}(t) = \langle \Psi_b^{(1)}(t); \, \hat{V}_i | \psi_b \rangle \langle \psi_a | \Psi_a^{(1)}(t); \, \hat{V}_k \rangle, \tag{21}$$

where the partial density-matrix elements  $\rho_{ba}^{jk}$  describe the vibrational coherence between vibronic wavepacket on electronic surface b induced by perturbation component j, and wavepacket on surface a induced by perturbation k.

Substituting  $\Psi^{(1)}$  from eqn (13), rearranging the terms, changing the time variables, neglecting the difference between  $f_j(t'+\tau)$  and  $f_j(t')$  for the slowly-varying envelope  $f_j$ , and considering the limit of  $t \to \infty$ ,  $t_0 \to -\infty$ , we obtain:<sup>37,50</sup>

$$\rho_{ba}^{jk}(t) = \frac{\pi}{2} e^{i\varepsilon_{ba}t} \delta(\Delta\omega_{jk} - \varepsilon_{ba}) \int_{-\infty}^{\infty} dt' f_j^*(t') f_k(t') \tilde{C}_{ba}$$
 (22)

$$\tilde{C}_{ba} = \int_{-\infty}^{\infty} d\tau e^{i(\omega_j - \varepsilon_{bi})\tau} C_{ba}$$
 (23)

$$C_{ba} = \langle \gamma_i | \mu_{bi}^* \hat{u}_b(\tau, t - t') \hat{u}_a(t - t', 0) \mu_{ai} | \gamma_i \rangle \tag{24}$$

where  $\delta$  is Dirac's delta function,  $\varepsilon_{ba} = \varepsilon_{bi} - \varepsilon_{ai}$ ,  $\Delta \omega_{jk} = \omega_j - \omega_k$ , and quantity  $C_{ba}$  is the cross-correlation function between nuclear wavepackets evolving on electronic surfaces b and a, and  $\tilde{C}_{ba}$  is its Fourier transform. In deriving eqn (22) we used the invariance of propagator  $\hat{u}$  with respect to time origin and the relation  $\hat{u}^*(t',t) = \hat{u}(t,t')$ . For brevity, we suppress the arguments of  $C_{ba}(t,t',\tau)$  and  $\tilde{C}_{ba}(t,t',\omega_j-\varepsilon_{bi})$  in the expressions above. (Note that  $\omega_j - \varepsilon_{bi} = \omega_k - \varepsilon_{ai}$  due to the presence of the  $\delta$ -function term in eqn (22).)

The individual terms in eqn (22) have a transparent physical interpretation. The  $\delta(\Delta\omega_{jk}-\varepsilon_{ba})$  term reflects energy conservation. The  $\int\! dt'$  integral is, up to a constant, the fluence of the ionizing laser pulse. Finally, the  $\tilde{C}_{ba}$  term reflects the vibrational density of states available at the target energy.

The effects of  $\tilde{C}_{ba}$  on the reduced density matrix are qualitatively different for the diagonal (population) and off-diagonal (coherence) terms. It is therefore useful to examine the two separately. For the diagonal term, the nuclear propagators  $\hat{u}_a$  and  $\hat{u}_b$  in eqn (24) coincide. The observation-time argument (t-t') therefore drops out, and the resulting autocorrelation function depends only on the time difference  $\tau$  between the two ionization times. The time-dependent phase factors in eqn (22) also cancel, leading to:

$$\rho_{aa}^{jk} = \frac{\pi}{2} \delta(\Delta \omega_{jk}) \tilde{C}_{aa} \int_{-\infty}^{\infty} dt' f_j^*(t') f_k(t')$$
 (22a)

$$C_{aa}(\tau) = \langle \chi_i | \mu_{ai}^* \hat{u}_a(\tau, 0) \mu_{ai} | \chi_i \rangle$$
 (24a)

where we formally kept the  $\delta(\Delta\omega_{ik})$  factor to preserve energy normalization. From eqn (22a) and (24a), it is clear that the final electronic state populations  $\rho_{qq}$  do not depend on the observation time t. The effect of the nuclear motion is to modulate the electronic transition probabilities by the vibrational density of states, at all times beyond the end of the laser pulse.

The situation is quite different for the off-diagonal elements of the density matrix, describing the electronic coherence. This is clearly seen by rewriting eqn (24) in an equivalent form:<sup>6,52</sup>

$$C_{ba} (\Delta t, \tau) = \langle \hat{u}_b(\Delta t, \tau) \mu_{bi} \chi_i | \hat{u}_a (\Delta t, 0) \mu_{ai} \chi_i \rangle$$
 (24b)

where  $\Delta t = t - t'$  is the time elapsed between wavepacket preparation and its observation. Thus, the reduced electronic coherence is modulated by the overlap of the vibrational wavepackets on the two electronic surfaces at the time of observation. This coherence will continue to evolve after the end of the laser pulse.

In very small molecules, where the vibrational motion remains low-dimensional, periodic revivals of the cross-correlation function  $C_{ba}$ , and therefore the electronic wavepacket motion, can be expected to persist for many vibrational periods.<sup>53,54</sup> A similar situation can be expected when the electronic wavepacket evolves on electronic surfaces which are nested either due to the symmetry reasons, or accidentally. In all other systems, we should expect the cross-correlation function to decay rapidly, leading to a suppression of the electronic wavepacket dynamics.

We should emphasize that the coupled vibronic dynamics discussed above is fully coherent; no "true" decoherence due to interaction with the environment is included in our model. The electronic and nuclear wavefunctions remain entangled throughout. The reason the electronic dynamics described by the off-diagonal reduced density-matrix elements "disappears", is the loss of overlap between the nuclear wavepackets. This is turn prevents factorization of the electronic and nuclear components of the wavefunction, and suppresses interference between the electronic eigenstates.

We also note that the time-domain, cross-correlation treatment of the reduced electronic coherence, exemplified by the eqn (22), (23) and (24b), is fundamentally different from the approach considered in ref. 5 and 55. The latter treatment assumes, as its very departure point, that electronic and nuclear wavefunctions are factorizable, neglecting the entanglement between the two. As the result, the treatment of ref. 5 and 55 only accounts for the structural inhomogeneity of the electronic transition, giving the upper bound for the electronic coherence lifetime due to nuclear motion. On the other hand, the approaches taken by ref. 6 and 52 fully account for the electronic-nuclear entanglement, are equivalent to our treatment above, and reach conclusions similar to ours.

### 4.3 An example: coherent hole dynamics in glycine cation

Following the seminal work by Kuleff, Breidbach, and Cederbaum, 4,56 electron-hole dynamics in glycine cation provides the paradigmal example of attosecond electron dynamics in molecular systems. It is therefore fitting to examine the effect nuclear motion has on these dynamics.

We limit ourselves to the case of the 3<sup>2</sup>A', 4<sup>2</sup>A' state pair, which has large contributions due to electron removal from the Hartree-Fock molecular orbital 14a' of the neutral species.<sup>56</sup> The corresponding Dyson orbitals (see Fig. 3) have norms of 0.852 and 0.877 respectively, confirming that these are 1h-type states. In the fixed-nuclei approximation, sudden removal of an electron from the 14a' orbital induces hole dynamics with the period of  $\approx 3.6$  fs, lasting for at least tens of femtoseconds.<sup>56</sup> In our calculations the two states are separated by  $\approx 1.19$  eV, corresponding to the natural oscillation period of  $\approx 3.47$  fs, in a good agreement with ref. 56.

We calculate the nuclear correlation functions on quadratic potential energy surfaces, determined at the occupationrestricted multiple active space configuration-interaction singles [ORMAS-CI(S)]<sup>57</sup>/cc-pVDZ<sup>58</sup> level, with the ROHF ground-state wavefunction used as the reference. We perform the expansion in the vicinity of the RHF/cc-pVDZ optimized geometry of the ground-state, neutral glycine. All quantum-chemistry calculations used GAMESS-US code. 59,60

The nuclear dynamics is performed in the basis of the products of the harmonic vibrational states of the neutral species, using an internally-developed code. 61,62 The implementation relies on a binary-tree representation of Franck-Condon factors<sup>63</sup> for efficient, sparse storage of vibrational wavepackets. The electronic potential-energy surfaces are implemented within the quadratic vibronic-Hamiltonian approach, 64 fitted to ab initio energy points. Non-adiabatic and electric-dipole coupling terms, although available in the code, were neglected presently. Wavepackets were propagated using the 4th-order Runge-Kutta integrator with the time step of 0.02 atomic units, independently on the 3<sup>2</sup>A' and 4<sup>2</sup>A' surfaces. Atomic masses of the most-abundant isotopes were used in the simulations. (In passing, we note that much more general and powerful wavefunction-dynamics packages, such as MCTDH,65 are readily available. Our choice here is driven more by familiarity and personal convenience than any technical advantage.) While this level of theory is far from reaching spectroscopic accuracy, it suffices to illustrate the main features of the dynamics, at least semi-quantitatively.

The 10-atomic glycine molecule possesses 24 vibrational degrees of freedom, which makes a full-dimensional, quantum

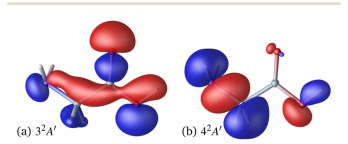


Fig. 3 Dyson orbitals for ionization of the ground-state glycine molecule, leading the 3A' (panel a) and 4A' (panel b) states of glycine cation. The red (blue) isosurfaces are at  $\pm 0.05 a_0^{-3/2}$ . The orbitals are shown at the optimized geometry of the neutral species

Table 1 Key parameters of the neutral-species normal modes (RHF/ccpVDZ, numbered sequentially, frequencies  $\omega$  in cm<sup>-1</sup>, harmonic vibrational periods T in fs) included in the simulations of glycine cation crosscorrelation functions. Modes "activated" by the ionization leading to the 3<sup>2</sup>A' and 4<sup>2</sup>A' cation states are indicated by checkmarks

No.	Symm.	ω	T	$3^2 A^\prime$	$4^2 A^\prime$	Dominant character
7	a"	87	383		<u> </u>	CC torsion
8	$a^{\prime\prime}$	267	125			CN torsion
14	a'	910	36.7		1	CC stretch
16	a'	1048	31.8			NH2 rock
17	a'	1228	27.2	1		CN stretch
19	a'	1301	25.6	1	1	COH bend/CH2 rock
20	a'	1328	25.1	1	1	COH bend/CH2 rock
22	a'	1555	21.4		1	CO(H) stretch
23	a'	1567	21.3		1	CH2 scissor
24	a'	1806	18.5		1	NH2 scissor
25	a'	2022	16.5		1	CO stretch
26	a'	3209	10.3			CH2 symm. stretch
28	a'	3724	8.96		1	NH2 symm. stretch
30	a'	4086	8.16			OH stretch

simulation of vibrational dynamics quite challenging. We instead choose to include the 14 degrees of freedom, along which the quadratic forcefields of the parent neutral and the two cations differ significantly. Ionization event is expected to trigger nuclear motion along these modes. The list of the "activated" modes is given in Table 1. Along the remaining normal modes, the neutral and cationic energy surfaces remain almost-perfectly nested. Dynamics along these modes can be neglected at short times. None of the "activated" modes are dominant, with each giving an incremental effect on the autoand cross-correlation functions. Such behaviour appears to be universal for sufficiently-large molecules.<sup>52</sup>

The calculated magnitude of the cross-correlation function  $C_{3^2A',4^2A'}(\Delta t,0)$  (eqn (24b), where we have taken  $\mu_{bi} = \mu_{ai} = 1$ ) is shown in Fig. 4. We are restricting ourselves to the case of zero inter-surface wavepacket delay  $\tau$ , as appropriate for the sudden preparation of the vibronic wavepacket. In the simulation, we allowed maximum of  $n_{\text{max}} = 2$  vibrational quanta per normal

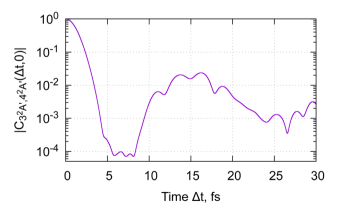


Fig. 4 Magnitude of the nuclear cross-correlation function for sudden, vertical preparation of the coherent superposition of the 3A' and 4A' electronic states in glycine cation.  $C_{3^2\mathrm{A'},4^2\mathrm{A'}}=1$  corresponds to a fully coherent electronic wavepacket;  $C_{\rm 3^2A^\prime,4^2A^\prime}=0$  indicates complete loss of the reduced electronic coherence.

mode, for the total of  $(1 + n_{\text{max}})^{14} \approx 4.8 \times 10^6$  vibrational basis functions. To check convergence of the results, we have performed test calculations with  $n_{\text{max}} = 3 (2.7 \times 10^8 \text{ basis functions})$ , which we continued to  $\Delta t = 6$  fs. We further evaluated the fulldimensional cross-correlation functions up to  $\Delta t = 3$  fs while allowing excitation up to  $n_{\text{max}} = 3$  in each of the 24 normal modes  $(2.8 \times 10^{14})$  basis functions). The results of the test calculations qualitatively agree with data in Fig. 4, confirming that our model is adequate for discerning the main features of the dynamics.

As we can see from Fig. 4, the nuclear cross-correlation function decays below 5% of the initial value after 3 fs. In the case of glycine cation, both nuclear surfaces remain bound. As long as the external sources of decoherence can be neglected, the dynamics therefore remains periodic, and the cross-correlation function (and the electronic-hole dynamics) will eventually revive. Due to the large number of the normal modes accessed by the dynamics, and their disparate characteristic times (see Table 1), the full-revival time is expected to be quite long - certainly in excess of 380 fs, the period of the C-C torsion mode. Due to the number of modes involved in the dynamics, partial revivals are expected to be weak. Indeed, in our simulation, the largest revival is at the 2% level, reached 14 fs after the start of the dynamics.

We therefore conclude that the coherent electronic wavepacket oscillation upon sudden preparation of the  $3^2A'$ ,  $4^2A'$ state pair in glycine cation is expected to disappear within 3 fs. We can expect to observe at most a single, decaying oscillation of these electronic dynamics, even under the ideal experimental conditions. This conclusion is at stark variance with the result obtained for the frozen nuclei, where pronounced charge oscillations are predicted to persist for tens of femtoseconds, with hardly any decay.<sup>56</sup> Thus, even though the nuclear displacements and velocities reached on a few-femtosecond time scales remain minuscule in absolute terms, their effect on the electronic dynamics is profound, and can't be neglected.

We note that coherent electronic dynamics in the glycine cation was previously examined using Ehrenfest dynamics,66 obtaining electronic dephasing times similar to ours. On the other hand, the study,55 also using Ehrenfest dynamics, found nuclear motion effects to be negligible for short-time electronic dynamics, and only become visible beyond 15 fs. The reasons for the discrepancy between the conclusions reached by ref. 55 and 66 are not immediately apparent.

We can reasonably suspect that the rapid-dephasing scenario generally applies to valence electronic wavepackets in medium-sized molecules.52 As long as the components of the electronic wavepacket modify the bonding situation in different parts of the molecule, the ensuing nuclear dynamics will rapidly drive apart the nuclear wavepackets evolving on different electronic eigensurfaces. The sheer volume of the available vibrational phase space will then ensure that the vibrational revival does not occur before the system has a chance do decohere due to interactions with the environment.

Naturally, other scenarios can and will arise, where the electronic wavepacket is confined to a manifold of nested

states, so that the nuclear-wavefunction coherence, together with the electron-hole dynamics could persist for longer times. As we have seen on the example above, the existence of such long-lived electronic oscillations is by no means guaranteed, and must be investigated, individually for each specific case.

#### Other types of electronic transitions

Although our analysis of nuclear-motion effects has focussed on 1-photon ionization, very similar results apply for other attoand femto-second electronic transitions. Thus, the nuclear autocorrelation function  $C_{aa}$  ( $\tau$ ) (eqn (24c)) modulates the high-harmonics yield in molecules, 67 as beautifully demonstrated in PACER ("Probing attosecond dynamics by chirp-encoded recollisions") experiments. 68,69 It has been recently shown that the shorttime autocorrelation functions can be extracted directly from the intensity-dependent HHG spectra for medium-sized molecules, 70 without resorting to elaborate theory modeling. These femtosecond nuclear dynamics in turn give access to the local potential-energy surface parameters of the transiently-created cations, 67,71 despite their fleeting, sub laser-cycle existence.

The nuclear autocorrelation function has also been shown to make an appearance in the strong-field streaking spectra of molecules, 72 creating a potentially very rich source of information on atto- and femto-second vibronic dynamics. We can further expect the nuclear autocorrelation to play a role in all processes whether the observables can be expressed in terms of the diagonal part of the electronic reduced density matrix.

Similarly, the nuclear cross-correlation is expected to modulate the attosecond electronic wavepacket dynamics in molecules, regardless of the specific preparation mechanism. It has been long used in the theory of molecular Raman transitions, 38,73 and was recently applied to modeling the dephasing of attosecond electronic dynamics in molecules. 6,52 We expect it to appear wherever molecular observables are determined by an electronic coherence.

## 5 Coherence in high-intensity high-frequency ionization

Broad-band ionization with attosecond X-ray pulses can lead to coherent population of a series of outer-valence-, inner-valenceor core-ionized states spanning energy ranges of 10 s of eV. In the perturbative regime, such as has been considered by us here in relation to high-frequency (XUV, X-ray) ionization so far, the relative populations of and coherences between these states are defined by the pulse shape, bound-continuum dipole moments and the subtle many-electron effects of loss of coherence due to break-up of an N-electron system. Within this perturbative picture, applying higher intensity ionising pulses will simply lead to a proportional increase of the populations of the ionized states, without changing their relative populations or degrees of coherence. However, it is well-established that multi-level systems driven by non-resonant high frequency, high-intensity fields exhibit fast transitions between the levels once the interaction energy associated with the bound-bound

transitions exceeds the inter-level spacing, see ref. 74 and 75:

$$\vec{E} \cdot \vec{\mu} > \hbar \omega_{\text{ion}},$$
 (25)

where  $\vec{E}$  is the electric field amplitude,  $\vec{\mu}$  is the transition dipole and  $\hbar\omega_{\rm ion}$  is the energy spacing between the two coupled bound levels. Therefore, ionising pulse of sufficient intensity can lead to redistribution of populations among the ionized states. This redistribution is expected to be strongly intensity-dependent and cannot be accounted for by the traditional perturbation theory where the interaction energy in the units of level spacing is a small parameter. On the contrary, the inverse perturbative treatment becomes possible in this case, in which the level spacing is much smaller than the interaction energy. 74,75 We postulate that transition to this regime signifies the onset of strong-field X-ray physics.

This suggested strong-field regime can be reached at much lower intensities than would be necessary for reaching any appreciable values of the ponderomotive potential on the scale of the ionization energy, as required by for the strong-field dynamics at the low (e.g. IR) field frequencies. As an order of magnitude estimation, one can think of ionic states separated by several eV ( $\approx 0.1$  a.u.) with transition dipole moments of the order of 0.1 a.u. driven by the ionising field of the frequency of several hundreds of eV ( $\approx$  10 a.u.). In this case, the interaction energy exceeds the level spacing for the electric field strength of the order of 1 a.u., at which the ponderomotive potential for the high frequency field is still negligible. This does not mean, however, that the suggested high-frequency strong-field regime is guaranteed to be observable. The main competing perturbative process that can obscure the suggested strong-field dynamics is the sequential multiple ionization. Indeed, apart from driving the non-resonant transitions between the ionized states, as assumed above, the high-frequency field can also lead to further ionization of the system. For hard X-ray fields and relatively long (e.g. 100 fs) pulses, multiple ionization clearly dominates at intensities of about 10<sup>19</sup> W cm<sup>-2</sup> and higher, as has been shown in a series of studies by Santra and co-workers, see e.g. ref. 76. Using soft X-rays would lower this intensity threshold due to larger photoionization cross-sections, which could be partly mitigated by using the presently available sub-fs X-ray pulses.3 The precise window of intensities available for the strong-field high-frequency photoionization dynamics proposed here remains to be established.

### 6 Conclusions

In conclusion, the partial loss of quantum coherence in the course of a break-up of an atomic or molecular system interacting with a short pulse of ionising radiation, as well as the decay of the remaining quantum coherence by both purely electronic and electronic-nuclear mechanisms are central issues in attosecond physics. Due to the complexity of the problem, involving many-electron dynamics, coupling of electronic and nuclear degrees of freedom, as well as strong laser field effects, theoretical description of the onset and decay of

coherence in attosecond ionization is still very much an open problem. Spectacular successes have been made in the theoretical modelling of purely electronic dynamics within frozen nuclei approximation, even in the most complicated cases of the breakdown of the molecular orbital picture of the ionization. Similarly, the role of nuclear dynamics (including, at the simplest level, the role of finite distribution of nuclear coordinates due to zero-point energy) is presently well-understood for the cases of simple (Koopmans-like) spectrum of the ionized states. Combining the two, however, as is necessary for example for the description of attosecond ionization in the inner-valence region of polyatomic molecules, has not yet been achieved and should be subject of future studies. The theoretical advances towards understanding the quantum coherence in attosecond photoionization allow one, for the first time, to look at the photoionization problem from the point of view of the quantum information theory. Quantifying the interplay between quantum entanglement and quantum coherence in photoionization will therefore form another avenue of future research. Finally, while a good understanding of the mechanisms underlying the quantum coherence in photoionization by high-intensity low-frequency fields has been achieved, the more notion of a high-intensity field in the opposite, highfrequency regime is yet to be defined. One possible definition has been proposed here. Further theoretical and experimental work needs to be done in order to verify the validity of this proposition.

### Author contributions

All authors contributed equally to the conceptualization of this work. The contributions to the writing were as follows: M. R. wrote Sections 1-3, S. P. wrote Section 4, V. A. wrote Sections 5 and 6. All authors contributed to the reviewing and the editing of the whole manuscript.

### Conflicts of interest

There are no conflicts to declare.

# Acknowledgements

Marco Ruberti acknowledges support from the Quantum entanglement in attosecond ionization EPSRC grant EP/V009192/1.

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