Eliminating the dipole phase in attosecond pulse characterization using Rydberg wave packets

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We propose a technique to fully characterize the temporal structure of extreme ultraviolet pulses by ionizing a bound coherent electronic wave packet. The influence of the dipole phase, which is the main obstacle for state-of-the-art pulse characterization schemes, can be eliminated by angle integration of the photoelectron spectrum. We show that in particular, atomic Rydberg wave packets are ideal and that wave packets involving multiple electronic states provide redundant information that can be used to cross-check the consistency of the phase reconstruction.

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I. INTRODUCTION

Ultrafast optics allows for the generation of laser pulses with durations on the femtosecond time scale. The pulses are shorter than the response time of any photodetector but limited by the femtosecond period of optical light. Methods to reconstruct laser pulses have been developed using nonlinear optical effects and cross correlation \cite{1}. In 2001, the femtosecond barrier was broken by the generation of attosecond pulses in the (extreme) ultraviolet (uv) range in the form of isolated pulses \cite{2} and pulse trains \cite{3}. This signaled the start of a new era of ultrafast experiments, known as attosecond physics \cite{4}.

Today, the use of attosecond pulses allows for direct measurement of optical laser pulses on the femtosecond time scale \cite{5}, but the generation and characterization of attosecond pulses remains an active field of research \cite{6}. Established techniques from ultrafast nonlinear optics cannot be extended into the uv regime due to two main reasons: (i) low photon fluxes associated with the uv pulses do not favor nonlinear processes, and (ii) high photon energies generally lead to target-dependent complex-valued susceptibilities that may obscure the field-reconstruction procedure. Therefore, all current attosecond characterization techniques rely instead on laser-dressed photoionization by near-infrared (nir) fields \cite{7}. Indeed, this is the principle of the frequency-resolved optical gating–complete reconstruction of attosecond bursts (FROG-CRAB) method \cite{8}, where the FROG algorithm from nonlinear optics \cite{9,10} is adapted as a phase gate given by the laser dressed continuum. If the ir field that dresses the continuum is weak, then the signal can be analyzed in terms of the phase retrieval by omega oscillation filtering (PROOF) method \cite{11}.

In this article, we propose a scheme to measure the temporal structure of extreme ultraviolet pulses by ionization of a bound electron wave packet. The key points are as follows: (i) the pump and probe steps are sequential, (ii) the intermediate states are bound, and (iii) the photoelectron is measured over all detection angles. As we will demonstrate, these three points result in an elimination of the dependence on dipole phases.

The elimination is exact provided that the parent cation is spherically symmetric and that correlation effects are negligible. Using Rydberg states as intermediate states is advantageous as they have single-particle character and possess little correlation. Photoionization cross sections are small leading to weaker signal strengths. Alkali-metal atoms would be ideal because they fulfill perfectly the spherical symmetry condition. In addition, the preparation of Rydberg wave packets can be performed using optical laser pulses. Noble gases are also a possibility, even though the ionic state is generally not spherically symmetric, and excitation energies are in the uv regime, making it less favorable.

We will also show that multilevel wave packets are useful for analysis of pulses with complex spectral structure, and that they allow for intrinsic self-consistency checks for the retrieved phases.

Rydberg states have been exploited before in attosecond experiments, but mainly with the focus on characterizing the bound electronic wave packet and not to characterize uv pulses \cite{27,28}. Another source of inspiration for our

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method is the recent work by You, Rohringer, and Dahlström, where it was proposed that nonsequential two-photon hole transitions can be used as an alternative method for attosecond pulse characterization [29]. Nonetheless, we will demonstrate many advantages of sequential laser excitation using multiple electronic bound states as intermediate states.

In Sec. II, we discuss the theory behind our proposed method. In Sec. III A, we show examples of how to reconstruct the spectral phase of an isolated attosecond pulse and an attosecond pulse train by using a Rydberg wave packet in potassium. The influence of electronic correlation effects is discussed in Sec. III B. We show that the dipole phase dependence due to inner-shell correlations and Fano resonances is much smaller in the angle-integrated photoelectron spectrum than in the directional photoelectron spectrum. Atomic units are used: $\hbar = |e| = 4\pi\varepsilon_0 = 1$ unless otherwise stated.

II. THEORY

A. The basic idea

In this article, we propose a scheme for uv pulse characterization that relies on photoionization of an electron wave packet for direct interference of different uv components. The basic idea is illustrated in Fig. 1.

Step 1 is preparation (pump) of a coherent electron wave packet,

$$|\psi(t)\rangle = \sum_j c_j e^{-i\epsilon_j t} |j\rangle,$$

where $\epsilon_j$ and $c_j$ are the energy and complex amplitude of state $j$, respectively. The details of the preparation of the bound wave packet are not important. However, the pump process must be completed before the attosecond pulse starts the ionization process (probe), so that pump and probe steps are sequential. For characterization of the uv pulse, it is important that the wave-packet states have the same parity (in more detail, the wave-packet states must be able to reach the same final state via one-photon absorption). Populated states with opposite parity, such as a partially depleted initial state, will not interfere with photoelectrons from the wave packet when detection in all angles is assumed. Preparation of wave-packet states with the same parity as the ground state allows for interference of uv components that are several eV apart and may be useful in special cases. However, we will focus on wave packets with opposite parity to the initial state.

Step 2 is field-free propagation of the wave packet for a time duration $\tau$. In step 3, the uv pulse arrives at the target,

$$E_X(\omega, \tau) = |E_X(\omega)| e^{i\phi_X(\omega)} ,$$

and ionizes the wave packet, as illustrated in Fig. 1. $|E_X(\omega)|$ and $\phi_X(\omega)$ are the amplitude and phase of the unshifted uv pulse, namely for $\tau = 0$. In the time domain, the electric field is given by

$$\tilde{E}_X(t, \tau) = \frac{1}{2\pi} \int d\omega E_X(\omega) e^{-i\omega(\tau-\tau)+i\phi_X(\omega)} = \tilde{E}_X(t-\tau, 0),$$

which shows that a pulse with a given $\tau$ corresponds to an unshifted pulse delayed by $\tau$. As step 3 is a one-photon ionization process, the photoelectron amplitude is given by first-order perturbation theory,

$$c_f(\tau) = \lim_{t' \to \infty} \frac{1}{i} \sum_j c_j d_{fj} \int dt' \tilde{E}_X(t', \tau) e^{i\omega_{fj} t'}$$

$$= -i \sum_j c_j d_{fj} |E_X(\omega_{fj})| e^{i\omega_{fj} \tau + i\phi_X(\omega_{fj})} ,$$

where $d_{fj}$ is the dipole transition moment from the bound state $|j\rangle$ to the continuum state $|f\rangle$. The lower bound of the time integral can be set to $-\infty$ because the ionization step of the wave packet is sequential and the $c_j$ amplitudes can be treated as constants.

The jitter between the pump and the probe pulses may lead to phase uncertainties. However, Rydberg states in alkali-metal atoms can be pumped with ir pulses that are phase-locked to the uv field through the ir-driven high-order harmonic generation process. In this case, jitter is removed as efficiently as for the FROG-CRAB and PROOF methods.

Different uv energies, $\omega_{fj} = \epsilon_f - \epsilon_j$, are required to reach the same final state from different bound states of the wave packet, as shown in Fig. 1(c). The interference of the different ionization pathways makes it possible to learn about the phase relation between the different spectral components of the uv

FIG. 1. Sketch of the method to characterize a uv pulse using a bound-electron wave packet. (a) Preparation of the wave packet with a laser pulse. (b) Field-free propagation of the electronic wave packet $\psi$ for the duration $\tau$. (c) Ionization of the wave packet by the uv pulse. Due to the coherent superposition of different electronic states, different spectral components of the uv pulse are absorbed to reach the same final continuum state with energy $\epsilon$. The interferences between the different spectral components contain the spectral phase information of the uv pulse that can be extracted by varying the delay $\tau$ and repeating the experiment.
pulse. The final photoelectron spectrum reads

\[
P(k_f, \tau) = |c_f(\tau)|^2 = \sum_j \left( A_j + \sum_{j' > j} B_{jj'} \cos \Theta_{jj'} \right),
\]

(5)

where \(k_f\) is the momentum of the photoelectron with energy \(\epsilon_f = k_f^2/2\). The variable \(A_j\) is the isolated contribution from the wave-packet state \(|j\rangle\) to \(|f\rangle\), while \(B_{jj'}\) and \(\Theta_{jj'}\) are the amplitude and phase of quantum interference between different parts of the wave packet with energies \(\epsilon_{j'} > \epsilon_j\). The explicit forms of the coefficients in Eq. (4) are

\[
A_j = |c_j|^2 |d_{fj}|^2 |E_X(\omega_{jj})|^2,
\]

(6a)

\[
B_{jj'} = 2|c_j| |c_{j'}| |d_{fj}| |d_{fj'}| |E_X(\omega_{jj})E_X(\omega_{jj'})|,
\]

(6b)

\[
\Theta_{jj'} = \arg[c_j c_{j'} d_{fj}^* d_{fj'} E_X(\omega_{jj}, \tau) E_X^*(\omega_{jj'}, \tau)].
\]

(6c)

The phase of the interference pattern can be rewritten to clearly show the quantum beating of the probability, at each energy difference \(\omega_{jj'} = \omega_f - \omega_{jj'} > 0\), as

\[
\Theta_{jj'} = \omega_{jj'} \tau + \phi_X^{(jj')}(\epsilon_f) + \phi_D^{(jj')}(\epsilon_f) + \phi_I^{(jj')},
\]

(7)

where the phase differences read

\[
\phi_X^{(jj')}(\epsilon_f) = \Phi_X(\omega_{jj'}) - \Phi_X(\omega_f) \approx \tau_X^{(GD)}(\omega_f') \omega_{jj'},
\]

(8a)

\[
\phi_D^{(jj')}(\epsilon_f) = \arg[d_{jj'}] - \arg[d_{fj}],
\]

(8b)

\[
\phi_I^{(jj')}(\epsilon_f) = \arg[c_j |c_{j'}|],
\]

(8c)

Equation (8a) defines the group delay \(\tau_X^{(GD)}(\omega_f)\) of the uv pulse computed as a finite-difference derivative at the mean photon energy, \(\omega_f' = (\omega_f + \omega_{jj'})/2\). The dipole transition elements \(d_{jj'}\) are complex valued functions that contribute energy-dependent dipole phases to the phase difference \(\phi_X^{(jj')}(\epsilon_f)\) in Eq. (8b). In contrast, \(\phi_I^{(jj')}\) in Eq. (8c) is inherent to the preparation process of the bound state wave packet and independent of the final photoelectron energy.

The main goal in characterizing the uv pulse is the determination of \(\tau_X^{(GD)}(\omega_f)\). If this group delay is known, then the spectral phase \(\Phi_X(\omega_f)\) can be fully reconstructed (up to an absolute value) by numerical integration. However, the oscillations in the photoelectron spectrum depend also on \(\phi_I^{(jj')}(\epsilon_f)\). The intrinsic phase shift \(\phi_I^{(jj')}(\epsilon_f)\) does not present an obstacle as it is energy-independent and corresponds to a constant shift of the pump-probe delay by \(\phi_I^{(jj')}/\omega_{jj'}\).

However, the energy dependence of the dipole phases \(\phi_D^{(jj')}(\epsilon_f)\) may prevent an accurate determination of the group delay of the uv pulse.

Before we continue our discussion, we discuss the importance of higher-order effects that go beyond first-order perturbation [cf. Eq. (4)]. Pazourek et al. [28] have pointed out that depopulation of the initial states results in an additional universal energy-dependent phase contribution in Eq. (7) that varies by multiples of \(\pi\) across the pulse spectrum. This is particularly an issue when the probability of ionization is high. By using Rydberg states, we reduce this phase dependence to a minimum as uv pulses are weak and cross sections of Rydberg states are two or more orders of magnitude smaller than for bound states. In Sec. III A, we give an explicit number for this effect and show that it is indeed negligible.

Similarly, a differential ac Stark shift between the Rydberg states due to the ionizing uv pulse is a third-order effect and can lead to phase corrections. Furthermore, the ac Stark shifts of neighboring Rydberg states at uv frequencies become more similar and approach quickly the ponderomotive potential, \(U_p = E_p^2/(4\omega_0^2)\), as the principal quantum number \(n\) increases. For an 10^{12} W/cm^2 pulse with \(\omega_0 = 50\) eV, the ponderomotive potential is less than 0.1 meV and the differential ac Stark shift between Rydberg states is even smaller, allowing us to safely neglect ac Stark effects.

**B. Uncorrelated spherically symmetric systems**

For spherically symmetric systems, and more specifically for spherically symmetric cations without correlation effects, the radial dipole moment to any energy eigenstate (continuum or bound) can be chosen to be real [30]. As a result, the angle-integrated photoelectron spectrum does not have an energy-dependent \(\phi_D^{(jj')}(\epsilon_f)\) phase anymore. To see this, we start with a partial wave expansion of the final momentum state,

\[
\psi_f(\mathbf{r}) = \frac{1}{k_{ff}} \sum_{L=0}^{\infty} \sum_{M=-L}^{L} i^L e^{-i\eta L} Y_{LM}^*(\hat{k}) Y_{LM}(\hat{r}) R_{L}(r),
\]

(9)

where \(k = k_f \hat{k}\) is the momentum of the photoelectron, and its energy is given by \(\epsilon_f = k_f^2/2\) [31]. Note that \(\psi_f(\mathbf{r}) = momentum-normalized. \langle \psi_f^* | \psi_f \rangle = \delta(\mathbf{k} - \mathbf{k}')\), while the real radial wave functions are chosen to be energy-normalized, \(\langle R_{L}(r) | R_{L'}(r) \rangle = \delta(\epsilon - \epsilon')\). Inserting Eq. (9) for the final state, \(f\), in Eq. (4) leads to the following expression for the photoelectron spectrum:

\[
P(k, \tau) = |c_k(\tau)|^2 = \frac{1}{k} \sum_{L, M, L', M'} c_j c_{j'} d_{fj}^* d_{fj'} E_X(\omega_{jj}, \tau) E_X^*(\omega_{jj'}, \tau) \times e^{-i\eta L + i\eta L'} \sum_{L, M} i^{L - L'} Y_{LM}^*(\hat{k}) Y_{LM}(\hat{r})
\]

\[
\times \sum_{L, M} |c_j|^2 |c_{j'}|^2 |d_{fj}| |d_{fj'}| E_X(\omega_{jj}, \tau) E_X^*(\omega_{jj'}, \tau),
\]

(10)

where we used the partial-wave basis \(|f\rangle = |\epsilon_f LM\rangle\) and \(d_{jj'} = \langle f | \hat{d} | j \rangle = \langle \epsilon_f L | r | n_f L_f \rangle C_{LM}^{L_f M_f} \sqrt{2L + 1} \mu\) being the polarization of the ionizing uv pulse. In Eq. (10), the sums on \(j\) and \(j'\) run over all excited states of the prepared wave packet. Since the radial part of the continuum states and the bound state are chosen to be real functions, the reduced matrix element, \(\langle \epsilon_f L | r | n_f L_f \rangle\), and the dipole, \(d_{jj'}\), are real and do not contribute to a phase difference in the photoelectron spectrum.

However, Eq. (10) depends on the scattering phase differences between all final partial-wave states, resulting in nontrivial angle and energy dependencies. By performing an angle-integrated measurement, the dependencies on the dipole phase can be eliminated,

\[
P(\epsilon, \tau) = \int d\Omega_k |c_k(\tau)|^2
\]

\[
= \sum_{L, j, j'} |c_j|^2 |d_{fj}| \sum_{L, j} \sum_{L', j'} |c_{j'}|^2 |d_{fj'}| E_X(\omega_{jj}, \tau) E_X^*(\omega_{jj'}, \tau),
\]

(11)
where the orthogonality of spherical harmonics reduces for each \((j, j')\) pair to a sum over all final angular momenta \(L_f\). Rewriting Eq. (11) by grouping contributions from the same \((j = j')\) and different electronic states \((j \neq j')\) together, we obtain

\[
P(\epsilon, \tau) = \sum_j \left( \tilde{A}_j + \sum_{j' > j} \tilde{B}_{jj'} \cos \tilde{\Theta}_{jj'} \right),
\]

where the variables read

\[
\tilde{A}_j = \sum_{L_j} |c_j|^2 |d_{j}|^2 |E_X(\omega_{fj})|^2 / k,
\]

\[
\tilde{B}_{jj'} = \sum_{L_{jj'}} d_{jj'} |c_{jj'}| |E_X(\omega_{fjj'})| / k,
\]

\[
\tilde{\Theta}_{jj'} = \omega_{fjj'} \tau + \phi^{(jj')}_{X}(\epsilon_f) + \phi^{(jj')}_{X}(\epsilon_f),
\]

where we write the index \(f\) explicitly on \(L_f\) for clarity. The phase term, \(\Theta_{jj'}\), is independent of atomic scattering (or dipole) phases and of \(L_f\). As the dipoles can be positive or negative, there should be no absolute magnitude around \(\eta_{L_f}\) dipole phases and of \(L_f\) corresponding phase differences \(\phi^{(jj')}_{X}(\epsilon_f)\), but not on the final scattering phases, \(\eta_{L_f}\) —exactly what we wanted.

When \(N\) states are involved in the electron wave packet, \(\psi(t)\), we obtain \((\tfrac{N}{2})\) different \(\omega_{fjj'}\) oscillations with their corresponding phase differences \(\phi^{(jj')}_{X}(\epsilon_f)\). It turns out there exist groups of three phase differences that obey the relation

\[
\phi^{(j,j,j)}_{X}(\epsilon_f) = \phi^{(j)}_{X}(\omega_{fj}) - \phi^{(j)}_{X}(\omega_{fj}) - \phi^{(j)}_{X}(\omega_{fj}) = \phi^{(j)}_{X}(\omega_{fj}) - \phi^{(j)}_{X}(\omega_{fj}) - \phi^{(j)}_{X}(\omega_{fj}),
\]

\[
\phi^{(j,j,j)}_{X}(\epsilon_f) = \phi^{(j)}_{X}(\omega_{fj}) + \phi^{(j)}_{X}(\omega_{fj}) + \phi^{(j)}_{X}(\omega_{fj}) = \phi^{(j)}_{X}(\omega_{fj}) + \phi^{(j)}_{X}(\omega_{fj}) + \phi^{(j)}_{X}(\omega_{fj}).
\]

Equation (14) holds true also for the retrieved phase differences \(\phi^{(jj')}_{X}(\epsilon_f) + \phi^{(jj')}_{X}(\epsilon_f)\), up to a constant, because \(\phi^{(j,j,j)}_{X}(\epsilon_f)\) does not need to fulfill any special relation. This connection can be used to cross-reference the extracted phases and to check the consistency of the retrieved phases. To our knowledge, no other ultrafast pulse reconstruction method (for femtosecond or attosecond pulses) possesses such a feature.

Our scheme to characterize uv pulses has several advantages that can be summarized as follows:

(i) Angle-integrated detection increases the total signal and ensures that ionization pathways to different final angular momenta of the photoelectron do not affect the phase reconstruction.

(ii) Rydberg states make it possible to probe spectral phase differences, bridging small energy distances as the energy difference between Rydberg states goes like \(n^{-3}\).

(iii) A combination of small and large energy differences, \(\omega_{fjj}\), makes it possible to probe phase differences between close and distant spectral phase components. This is especially useful if certain spectral regions have negligible intensity and need to be bridged, as is the case for pulse trains.

(iv) There is no constraint on the polarization of the uv pulse. It works for linearly, circularly, and elliptically polarized pulses.

(v) Wave packets with \(N\) electron states (with the same angular momentum) lead to \((\tfrac{N}{2})\) beating frequencies in the photoelectron spectrum, providing redundant information [Eq. (14)] to cross-check the consistency of the reconstruction.

In more detail, electron correlation results in complex-valued dipole moments that may affect the pulse reconstruction, even if the photoelectron is detected over all angles. As we will show in Sec. III B for the case of Rydberg wave packets, these effects are surprisingly small even in correlated energy regions close to autoionizing resonances.

## III. RESULTS

In this section, we first use Rydberg wave packets to characterize a single attosecond pulse (Secs. III A 1 and III A 2) and an attosecond pulse train (Sec. III A 3). We then use photoionization of Rydberg wave packets to study the electronic-structure properties of angle-resolved photoelectrons (Sec. III B 1). Finally, we study the influence of electron correlation in the case of potassium in a resonance-free region (Sec. III B 1) and in neon near the 2s−13s Fano resonance (Sec. III B 4).

We choose potassium as our main system of interest. Atomic alkali metals possess several advantages and have been used for ultrafast experiments [32,33]: (i) After removal of the outermost \(s\) electron, alkali-metal cations are closed-shell systems with relatively small amount of correlation. (ii) The mean-field potential of alkali-metal cations is spherically symmetric, fulfilling exactly the conditions for real reduced dipole matrix elements. (iii) The excitation energies of the Rydberg states in all alkali metals lie within 1.4 and 5.4 eV [34]. This is a convenient energy range as it is easily accessible with common laser systems and standard nonlinear optics that make it possible to transfer up to 100% of the population into Rydberg states.

A practical question that arises when using Rydberg states is the signal strength because Rydberg states have much lower photoionization cross sections \((\propto n^{-3})\) than valence shells [35]. To characterize attosecond pulses, low-lying Rydberg states with \(n < 10\) are sufficient. Their cross section is roughly 100 times weaker than the outermost valence shell. Given common gas parameters \((10^{17} \text{ at./cm}^3, 1 \text{ mm gas jet diameter} [36]\) and attosecond pulse parameters \((10^{11} \text{ photons/s}, 1 \text{ kHz–1 MHz repetition rate} [4]\), we estimate a total number of \(10^3–10^6\) electrons/s to become ionized for photon energies up to 100 eV.

 Dipole-dipole interactions between Rydberg states can lead to corrections to the energy level and the Rydberg wave function. The energy corrections are on the 1 meV scale for the Rydberg states considered here \((n < 10)\) and gas densities up to \(10^{17} \text{ at./cm}^3\). Rydberg states up to \(n < 10\) have periods in the tens of femtosecond regime, which are 100 times faster than the picosecond-scale corrections corresponding to 1 meV changes. Consequently, the interaction between Rydberg states or neighboring atoms is negligible for this type of measurement.

All results obtained by explicit time propagation [37] utilize the XCIID program [38] with the following numerical parameters [39] and the wave-function splitting method to analyze the photoelectron spectrum as described in Ref. [40]. This method
has been successfully used to analyze photoelectron spectra in the multiphoton regime [41,42]. In this work, we are staying in the one-photon regime. Correlation effects for potassium are studies by means of the random-phase approximation with exchange (RPAE) by adapting existing numerical codes [43] with the initial state being a given np virtual orbital obtained by the Hartree-Fock (HF) equation of the K⁺ core [44,45]. The time-dependent configuration-interaction singles method (TDCCIS) is also used to study correlation effects in neon, as it has been shown to describe well many-body physics in the attosecond [46] and strong-field regime [47].

A. Pulse characterization

1. Single pulse + two-level wave packet

As Rydberg wave packets can be easily prepared with common laser systems, we omit the description of their preparation. In our first example, we choose the wave packet in potassium (K) to be a coherent superposition between the 3p⁶4p and 3p⁶5p Rydberg states,

\[ |\psi(t)\rangle = \frac{1}{\sqrt{3}} \sum_{n=4,5} e^{-i \epsilon_n t} |np\rangle. \]  (15)

We omit reference to the inner-shell configuration as we are only interested in the Rydberg electron, and we assume that the ionic configuration will not change (see Sec. III B 4 for a more general discussion and the influence of interchannel coupling). Potassium is treated within the HF level, where the HF procedure is done for K⁺. The valence electron interacts with a mean-field potential of K⁺, and interchannel correlation is neglected.

In the probe step, the 4p-5p Rydberg wave packet is ionized by the attosecond pulse. We consider an isolated attosecond pulse with a constant spectral phase: \( \phi_X(\omega) = 0 \), with a quadratic spectral phase: \( \phi_X(\omega) = 100(\omega - \omega_0)^2 \), and with a cubic spectral phase: \( \phi_X(\omega) = 100(\omega - \omega_0)^3 \). Each attosecond pulse has a central photon energy of \( \omega_0 = 74 \text{ eV} \) (=2.72 a.u.) and a full width at half-maximum (FWHM) spectral intensity width of 7.4 eV (=0.27 a.u.) corresponding to a pulse duration of 247 as for a Fourier-limited pulse. The pulse with linear chirp (quadratic phase) is symmetrically stretched in time, increasing the pulse duration, while the pulse with quadratic chirp (third-order phase) experiences an asymmetric pulse deformation.

In Figs. 2(a)–2(c), the photoelectron spectra, \( P(\epsilon, r) \), are shown for the attosecond pulse with (a) no chirp, (b) linear chirp, and (c) quadratic chirp. In accordance with Eqs. (8a) and (13c), the chirps are directly visible as a constant, linear, and quadratic shift of the beating patterns in Figs. 2(a)–2(c), respectively.

In Figs. 2(d)–2(f), the photoelectron spectra, \( P(\epsilon, v) \), are shown where the delay axis is Fourier-transformed.

In Figs. 2(g)–2(i), the retrieved phases of the 1.36 eV beating signal for the different attosecond pulses are shown.

According to Eq. (13c), these phase differences correspond to \( \phi_{jj}(\epsilon_f) + \phi_{jj}'(\epsilon_f) \), where \( j = 4p \) and \( j' = 5p \). The intrinsic phases of the Rydberg wave packet \( \phi_{jj}'(\epsilon_f) \) are subtracted in Figs. 2(g)–2(i) such that the retrieved phase at the spectral peak of the beating signal is 0. The phase \( \phi_{jj}'(\epsilon_f) \) does not affect the reconstruction of the spectral chirp of the pulse (see Sec. II). The analytic solutions of the phase differences, \( \phi_{jj}'(\epsilon_f) \), are plotted as black-dashed lines, and they agree perfectly with the retrieved phases.

2. Single pulse + multilevel wave packet

The Rydberg wave packet is not limited to be a superposition of two Rydberg states for characterization of uv pulses. Involving \( N > 2 \) states (with the same angular momentum \( l \)) in the Rydberg wave packet is even more beneficial as \( \langle \langle \rangle \rangle \) beating frequencies occur in the photoelectron spectra—all containing spectral information about the pulse.

We consider the three-level Rydberg wave packet,

\[ |\psi(t)\rangle = \frac{1}{\sqrt{3}} \sum_{n=4,7,8} e^{-i \epsilon_n t} |np\rangle, \]  (16)

which provides two high-frequency beatings, \( \omega_{64} = \epsilon_{5p} - \epsilon_{4p} = 2.26 \text{ eV} \) and \( \omega_{74} = \epsilon_{7p} - \epsilon_{4p} = 2.12 \text{ eV} \), and one low-frequency beating, \( \omega_{87} = \epsilon_{8p} - \epsilon_{7p} = 0.14 \text{ eV} \). The reference time, where all phases of the wave packet are the same, is arbitrarily chosen to be \( t_0 = -600 \text{ a.u.} \). In Fig. 3(a), the pulse spectrum shifted by the ionization potentials is shown to indicate where the corresponding photoelectron distributions are expected.

Figure 3(b) shows the photoelectron spectrum, \( P(\epsilon, r) \), for ionization of the three-level Rydberg wave packet with the quadratic chirped pulse from Sec. III A 1. The photoelectron spectra, \( P(\epsilon, v) \), for the three frequencies \( v = \omega_{47}, \omega_{48}, \omega_{78} \) are shown in Fig. 3(c).
The analytic results are shown as black dashed lines. (d) Retrieved phases of the three frequency components. The static and the 0.14, 2.12, and 2.26 eV oscillating signals of the photoelectron spectrum. (c) The static and the 0.14, 2.12, and 2.26 eV oscillating components of the photoelectron spectrum. (d) Retrieved phases of the three frequency components. The analytic results are shown as black dashed lines. The analytic results match perfectly for the three beating patterns [see Fig. 3(d)]. All three curves are parabolas with curvatures $3(\nu_j \pm \pi/4)$ depending on the quadratic chirp, $\beta = 100$, and on the energy difference between the states, $\omega_j \pm \pi/4$. Also the relations between $\phi_X(j)$ [see Eq. (14)] hold for all combinations. This demonstrates that our method provides an internal mechanism to confirm the consistency of the retrieved phases.

3. Pulse train + multilevel wave packet

We will now demonstrate that the 4$p$-7$p$-8$p$ three-level Rydberg wave packet from Sec. III A 2 can be applied to probe the phase of pulse trains on the attosecond and femtosecond time scales simultaneously. We choose a pulse train that consists of the 48th, 49th, and 50th harmonic of 800 nm with some rather complicated phase relations: The 48th harmonic has no spectral chirp but a phase constant of $\pi/4$. The 49th harmonic has a linear chirp, $\phi_X(\omega) = -0.5/\delta\omega^2(\omega - \nu_{84})^2$. Finally, the 50th harmonic has a quadratic chirp, $\phi_X(\omega) = 0.5/\delta\omega^3(\omega - \nu_{84})^3 + \pi/3$. Each harmonic has a spectral intensity FWHM width of $\delta\omega = 400$ meV.

The three-level Rydberg wave packet allows for interference of spectral components within an individual harmonic and between adjacent harmonics. This is visualized in Fig. 4(a), where the uv spectrum is shifted by the three different binding energies. The energy difference $\omega_{4p,8p}$ is smaller than the spectral width of the harmonics, and it therefore allows for probing the phase relation within each harmonic individually [by overlap of red and blue in Fig. 4(a)]. The energy differences $\omega_{4p} \approx \omega_{50}$ are comparable to the spectral separation between harmonics and allow for probing of the phase relation between adjacent harmonics [by overlap of the brown curve with the blue or red curves in Fig. 4(a)].

Figure 4(b) shows the photoelectron spectrum $P(\epsilon, \tau)$ from the three-level Rydberg wave packet ionized by the attosecond pulse train. The three main delay-independent peaks correspond to the ionization of the 4$p$ electron by the three harmonics. The contributions of 7$p$ and 8$p$ cannot be spectrally distinguished, but their interference results in three distinct low-frequency beatings in Fig. 4(b).

The $\omega_{4p,7p}$ and $\omega_{4p,8p}$ modulations are located at kinetic energies 73.7 and 75.3 eV in Fig. 4(b), corresponding to interference of harmonic 48 with 49 and harmonic 49 with 50, respectively. The extracted phases from all three modulations, shown in Fig. 4(d), agree perfectly with the analytic predictions, and Eq. (14) holds for kinetic energies where all three beating patterns have a nonvanishing signal. In this way, we can reconstruct the phase within each harmonic (up to a constant) using the 7$p$-8$p$ beating signal. This is shown by the violet curve in Fig. 4(d), which indicates that harmonic 48 has no chirp, harmonic 49 has a linear chirp, while harmonic 50 has a quadratic chirp. To determine the phase between two neighboring harmonics, the phase at one specific kinetic energy of the 4$p$-7$p$ or 4$p$-8$p$ oscillations needs to be found. At 73.7 and 75.3 eV, the phase differences of the 4$p$-7$p$ modulation are $-0.34\pi$ and 0.49$\pi$, respectively [see the red dots in Fig. 4(d)]. Using our knowledge about the spectral phase within each harmonic, which we just determined, we find that the constant phase difference between harmonics 48 and 49 is $-\pi/4$, and that between harmonics 49 and 50 is $\pi/3$. Hence, we have fully reconstructed the spectral phases of the pulse train up to a constant global phase. This shows that our method is useful for characterization of pulse trains (on both attosecond and
femtosecond time scales) and that it can be used to benchmark the accuracy of the RABBITT method [3].

B. Electronic structure

1. Directional dipole phases

As we discussed in Sec. II and showed in Sec. III A, the dipole phase drops out when studying the angle-integrated photoelectron spectrum. In this section, we analyze the dipole phase dependence of the angle-integrated photoelectron spectrum for an electron ionized along the field polarization $\hat{z}$.

In Fig. 5(a), we show the dipole phases $\psi_D^{4p/5p} = \arg\{\epsilon, \hat{z}, \epsilon|4p/5p\}$ for ionizing an electron in the $\hat{z}$ direction with kinetic energy $\epsilon$. The dipole phases vary nonmonotonically by $\pi/2$ over an energy region from 20 to 90 eV. This behavior can be attributed to a Cooper minimum in the photoionization cross section of K from the $np$ states. The dipole phases of the Rydberg states $4p$ and $5p$ of K are almost exactly the same. What enters in our scheme is the phase differences between the two dipoles.

In Fig. 5(b), we compare the delay due to the dipole phase difference of $4p$ and $5p$, $\phi_D^{(4p,5p)}(\epsilon) = \arg\{\epsilon, \hat{z}, \epsilon|4p,5p\} = (\psi_D^{4p}(\epsilon) - \psi_D^{5p}(\epsilon))/(-\epsilon_{5p} - \epsilon_{4p})$, that enters in our proposed method (blue dashed line) with the Wigner delay, $\delta t_{\psi D}^{(4p,5p)}(\epsilon)$, that enters in the FROG-CRAB, PROOF, and RABBITT methods (red dotted-dashed line) (in addition to an ir-induced delay—not considered here). The Wigner delay is an order of magnitude larger than the delay of our directed method. This shows that even without angle integration, the influence of the dipole phase is strongly reduced thanks to the similarity between neighboring Rydberg states. The delay due to the angle-integrated phase difference, $\phi_D^{(4p,5p)}(\epsilon)$, is exactly zero (blue solid line), as expected from our discussion in Sec. II.

2. Critical pulse duration

The curvature (second spectral derivative) of the dipole phase is important for the pulse characterization because it leads to an energy-dependent Wigner delay and, thus, artificial deformations in the reconstructed pulse. The curvature is around 3 as/eV for directed photoelectrons at photon energies up to 50 eV in potassium from $4p$ or $5p$ (a value that is reasonable to assume also for noble-gas atoms). Given a Gaussian Fourier-limited pulse with FWHM-pulse duration $\delta t$, a constant dipole phase curvature, $c$, implies that the reconstructed phase becomes $\phi = \frac{\epsilon}{c}(\omega - \omega_0)^2$ (plus an unknown constant). The reconstructed FWHM-pulse duration becomes

$$\delta t_{\text{rec}} = \delta t \sqrt{1 + 16 \ln^2(2) c^2/\delta t^4} = \delta t \sqrt{1 + (\delta t_{\text{crit}}/\delta t)^4},$$

where we have defined a critical pulse duration, $\delta t_{\text{crit}} = \sqrt{4 \ln^2(2)c}$. Expressed in commonly used units, the critical pulse duration is $\delta t_{\text{crit}}(\text{as}(\text{eV})) = 42.7 \sqrt{c}(\text{as}/\text{eV})$.

If we use $c \sim 3$ as/eV, the critical pulse duration is 74 as. This shows that, not only for the characterization of the shortest attosecond pulses or zeptosecond pulses [48,49], the dipole phase cannot be ignored. Already for a 100 as Fourier-limited pulse, the error is 14%. In our approach, the dipole phase curvature that enters (without angle integration) is 40 times smaller, reducing the critical pulse duration to 11 as. With angle integration, $c = 0$ and no critical duration exist.

As mentioned in Sec. II, depopulation introduces a universal energy-dependent phase. The small cross sections and weak uv intensity lead to small depopulations and consequently to small phase effects. Ionization of K $4p$ by an 100 as pulse (10$^{12}$ W/cm$^2$ peak intensity and 70 eV photon energy) leads to relative depopulation of $10^{-4}$ and a negligible depopulation-induced phase curvature of $c < 2 \times 10^{-3}$ as/eV.

Next, we estimate how much $c$ is affected by electron correlation effects.

3. Residual correlation effects

Once electronic correlations are considered, the dipole phase affects the photoelectron spectrum even after angle integration. Photoionization of alkali-metal atoms is a prototypical test case for correlation effects for close-to-threshold photoionization [50]. In this work, we consider larger photon energies where, in general, correlation effects are expected to be smaller. In Fig. 6, we compare the retrieved phases of the $\omega_{4p,5p}$ oscillations of the directional (dashed lines) and angle-integrated (solid lines) photoelectron spectra of potassium treated within the Hartree-Fock approximation (HF; blue lines). This calculation builds upon the perturbation diagram around 3 as/eV for directed photoelectrons at photon energies up to 50 eV in potassium from $4p$ or $5p$ (a value that is reasonable to assume also for noble-gas atoms). Given a Gaussian Fourier-limited pulse with FWHM-pulse duration $\delta t$, a constant dipole phase curvature, $c$, implies that the reconstructed phase becomes $\phi = \frac{\epsilon}{c}(\omega - \omega_0)^2$ (plus an unknown constant). The reconstructed FWHM-pulse duration becomes

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shown in Fig. 7(a), where \( j = 4p, 5p \) are the excited Rydberg states and \( f = ks, kd \) are the final photoelectron states, both computed within HF. This type of calculation contains no correlation effects by definition, and the corresponding delay of the angle-integrated calculation is zero, as expected from Sec. II. In Fig. 6, we also show the random-phase approximation with exchange (RPAE; yellow lines), which includes additionally correlation effects with the electrons from the Ar\(^+\) core, \( c = 3s, 3p \), as illustrated by the perturbation diagrams in Figs. 7(b)–7(e). More details about RPAE are found in Ref. [45].

The direction of the photoelectron is chosen to be parallel to the laser polarization direction (\( \theta = 0 \)). The energy range of 70–80 eV is chosen to avoid on-shell excitation of autoionizing resonances in the continuum (see Sec. III B 4) and to focus on the residual effect (virtual coupling) to other particle-hole excitations.

When correlation effects are included, the reduced dipole moments become complex-valued even in spherically symmetric systems, resulting in a dipole phase dependence in the angle-integrated photoelectron spectrum. In Fig. 6, the RPAE results, which include correlation, show a weak energy dependence. The correlation-induced delay in the angle-integrated spectrum (solid yellow line) is centered around 6 as. A constant delay difference is, however, not of interest and does not influence the determination of the spectral chirp. The energy dependence is, thanks to the Rydberg states, again quite weak.

The average slope is \(-0.052\text{ as/eV}\) in the 70–80 eV range, resulting in a critical pulse duration of \( \tau_{\text{crit}} = 9 \text{ as} \). The influence of correlation on the directional photoelectron (dashed yellow line) is with an average slope of 0.12 as/eV more than twice as strong as in the angle-integrated case. It shows also in the presence of correlations that the dipole phase dependency is reduced when studying the angle-integrated photoelectron spectrum.

The influence of correlation can be further reduced when going to higher Rydberg states, or when going to lighter atoms (e.g., sodium with a neonlike core), which generally contain less electronic correlation.

4. Autoionizing resonances

When the ionizing test pulse is resonant with autoionizing inner-shell excitations, Fano resonances become visible in the photoelectron spectrum, which are spectrally highly localized around the autoionizing resonance energy (in contrast to the residual effects discussed in Sec. III B 3). Since autoionization is a correlated process, the dipole phase dependence will survive in the photoelectron spectrum after angle integration in the form of amplitude and phase modulations in the beating patterns.

To demonstrate the influence of an autoionizing state, we choose neon and we target the lower 2s\(^{-1}\)ns autoionizing states, which are even-parity states and can only be reached with an even number of photons. First, we prepare a Rydberg wave packet between 2p\(^{-1}\)3s and 2p\(^{-1}\)4s with two 4.8 fs (=200 a.u.) Gaussian pulses with center frequencies 16.8 and 19.7 eV. After the wave packet is prepared, we ionize it with a 508 as (=21 a.u.) Fourier-limited Gaussian pulse with a center frequency of 25.9 eV (=0.95\(E_h\)), which ionizes the Rydberg electron but also drives the ionic transition between 2p\(^{-1}\) and 2s\(^{-1}\), leading to population of the autoionizing 2s\(^{-1}\)ns states. The calculations [51] are based on TDCIS, which accurately describes the 2s\(^{-1}\)np Fano resonances in neon [52].

Phase modulations in the beating pattern will only be due to \( \phi_{ij}(\epsilon_f) \) because \( \Phi_{ij}(\epsilon_f) = 0 \) in Eq. (7). In Fig. 8(a), the angle-integrated photoelectron spectrum around the 2s\(^{-1}\)3s is shown, and Fig. 8(b) shows the amplitude of the static and oscillating contributions. The 2s\(^{-1}\)3s and higher Fano resonances are visible in the static and especially in the oscillating amplitudes. The static signal increases beyond 25 eV [see Fig. 8(b)] to two-photon absorption of the attosecond pulse. This process is not delay-dependent and contributes only to the static background.

The retrieved phases are shown in Fig. 8(c), where the energy positions of the 2s\(^{-1}\)ns resonances are highlighted by vertical dashed lines. Also the intrachannel TDCIS result for the angle-integrated photoelectron spectrum is shown (blue dashed line), where the interchannel interactions that are responsible for the autoionization of all 2s\(^{-1}\)nl states are not included. Ignoring interchannel effects eliminates correlation effects, and as a result no phase modulations

![FIG. 7. Perturbation diagram for (a) HF, (b) direct-forward RPAE, (c) direct-forward RPAE, (d) exchange-forward RPAE, and (e) exchange-backward RPAE. Up (down) arrows label electron (hole) states. Further details are given in the main text.](image)

![FIG. 8. (a) Angle-integrated photoelectron spectrum of the 2p\(^{-1}\)3s–2s\(^{-1}\)4s wave packet ionized by an isolated attosecond pulse. (b) The static (black dashed line) and oscillating components (solid red line) of the angle-integrated photoelectron spectrum. (c) Retrieved phase of the oscillating components of the angle-integrated (red solid line) and directional (green dashed-dotted line) photoelectron spectrum for the full TDCIS model. The intrachannel TDCIS results for the angle-integrated photoelectron spectrum are shown as a reference (blue dashed line). The corresponding group delay of the phase differences is shown on the right.](image)
around the resonance energies are seen. The phase changes due to the \(2s^{-1}3s\) Fano resonance are small for the angle-integrated photoelectron spectrum. The induced delay does not exceed \(\pm 3\) as.

For the directional photoelectron spectrum, the correlation-induced delay is, by up to \(-30\) as, an order of magnitude larger than for the angle-integrated result, as we have already found in Sec. III B 3 for residual correlation effects. In both cases (directional or angle-integrated), the delay is relatively small compared to the \(2p^{-1}3s\) resonance lifetime of 6.4 fs (\(\Gamma = 0.1\) eV within TDCIS). The derivative of the absolute dipole phase, measured in FROG-CRAB, reflects the lifetime of the resonance. In our method, we measure the difference in the influence of the \(2s^{-1}3s\) autoionizing state on the Rydberg states \(2p^{-1}3s\) and \(2p^{-1}4s\), which is two orders of magnitude smaller than the absolute dipole phase.

IV. CONCLUSION AND OUTLOOK

We have proposed a method to characterize attosecond uv pulses with the help of bound-electron wave packets. Different spectral components of the uv pulse interfere as an electron can be ionized from different energy levels. This leads to quantum beats (an oscillating photoelectron signal) as a function of the pulse delay. We showed that angle integration eliminates the influence of the dipole phase in the photoelectron spectrum, which becomes critical for pulses with duration below 100 as.

Rydberg wave packets have favorable properties for the proposed method. The energy spacing between electronic states decreases with \(1/n^3\), offering high spectral energy resolution and the possibility to bridge energy regions with no spectral weights (as in the case of pulse trains). The downside of using the Rydberg state is the small photoionization cross section leading to weak signal strengths. Multilevel wave packets open up the possibility for consistency checks of the retrieved phases—a feature that does not exist in other pulse reconstruction techniques.

We studied the role of correlation effects. We found that the dipole phase cannot be completely eliminated by angle integration. The effect is, however, much weaker than the changes in the Wigner delay and the ir-induced continuum-continuum delay. In particular, Rydberg wave packets minimize correlation effects. We have shown in potassium and neon correlation effects result indeed in negligible phase corrections.

When vibrational Rydberg-like wave packets instead of electronic ones are used and the pulse spectrum is fully known, this approach can be turned around and used to probe non-Born-Oppenheimer dynamics. Correlated electronic-vibrational motion results in an extra phase contribution, which is equivalent to the dipole phase dependence in atoms due to electronic correlations.

We believe ionizing Rydberg wave packets is a versatile approach to determine the spectral phases of complex pulses, e.g., generated by table-top attosecond laboratories or FEL facilities. We have focused here on pulses in the uv regime. This technique can be easily extended to x rays with shorter wavelengths and into the optical and near-infrared domains with longer wavelengths.

As pulse durations approach the zeptosecond regime [53], ionization from different inner shells cannot be distinguished anymore, making established attosecond pulse characterization techniques more prone to error [54]. Our method is not affected by inner-shell ionization. Together with the spectral phase accuracy and the applicability to a very broad spectral range, we believe this approach offers several key properties for characterizing the next generation of attosecond pulses.

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[39] A pseudospectral grid with a radial box size of 120a₀, 750 grid points, and a mapping parameter of ζ = 0.5 are used. There is no complex absorbing potential. The splitting function starts around R_{split} = 70a₀, a smoothness of 2a₀, and a splitting interval of dt_{split} = 2 a.u. The maximum angular momentum is 2, and Hartree-Fock (HF) orbitals up to an energy of 5E₀ are considered. The propagation method is Runge-Kutta 4 with a time step dt = 0.05 a.u.


[51] A pseudospectral grid with a radial box size of 220\(a_0\), 1000 grid points, and a mapping parameter of \(\xi = 0.5\) are used. The splitting function starts around \(R_{\text{split}} = 170a_0\), a smoothness of \(2a_0\), and a splitting interval of \(dt_{\text{split}} = 1\) a.u. The maximum angular momentum is 3, and Hartree-Fock orbitals up to an energy of \(10E_\text{h}\) are considered. The propagation method is Runge-Kutta 4 with a time step \(dt = 0.05\) a.u.


[54] J. Biegert (private communication).