Electronic structure effects in spatiotemporally resolved photoemission interferograms of copper surfaces

M. J. Ambrosio and U. Thumm
Department of Physics, Kansas State University, Manhattan, Kansas 66506, USA
(Received 28 September 2017; published 20 November 2017)

Attosecond photoelectron spectroscopy allows the observation of electronic processes on attosecond time scales (1 as = 10⁻¹⁸ s), as has been demonstrated in proof-of-principle experiments that probe the electronic dynamics in isolated atoms with unprecedented accuracy. Its recent expansion to solid targets is starting to allow the distinction of ultrafast collective electronic processes in matter with added spatial resolution, probing the electronic band structure and dielectric response in nanoplasmonically enhanced light-induced processes of relevance for photocatalysis, optoelectronics, and light harvesting. Based on a quantum-mechanical model for photoelectron emission by an attosecond pulse train from the d band of a Cu(111) surface into a delayed assisting laser pulse, we calculate two-pathway two-photon interferograms as functions of the photoelectron energy and pulse delay. Our results scrutinize the dependence of observable photoelectron interferograms on the electronic structure of and electron transport in the substrate and agree well with experimental spectra and semiclassical Monte Carlo simulations of Lucchini et al. [Phys. Rev. Lett. 115, 137401 (2015)].

DOI: 10.1103/PhysRevA.96.051403

Based on the process of higher-order-harmonics (HHs) generation in atomic gases [1,2], continued progress in ultrafast laser technology enabled the superposition of HHs to attosecond pulse trains (APTs) [3] and isolated attosecond pulses (IAPs) [4,5] that are perfectly synchronized with the driving infrared (IR) laser, allowing for time-resolved investigations of the electronic dynamics in gases and condensed-matter systems. These time-domain investigations complement traditional energy-domain spectroscopies [6] and are implemented in two types of experimental setups, streaked photoelectron emission and reconstruction of attosecond beating by interference of two-photon transitions (RABBITT). In streaking experiments, photoelectrons are emitted by a spectrally broad extreme ultraviolet (XUV) IAP into the electric field of a delayed infrared pulse [7]. Recording photoelectron energy spectra as a function of the IAP-IR delay reveals temporal information through delay-dependent photoelectron energy shifts, reducing the measurement of ultrashort time intervals (of the order of 10 as, 1 atomic unit of time = 1 Å. s = 24 as) to conveniently detectable photoelectron energies. Instead of IAPs, RABBITT experiments employ XUV APTs to emit electrons into a synchronized delayed IR pulse, imaging electric dynamics as photoelectron-yield oscillations due to two-pathway two-photon quantum interference (RABBITT). In streaking experiments, photoelectrons are emitted by a spectrally broad extreme ultraviolet (XUV) IAP into the electric field of a delayed infrared pulse [7]. Recording photoelectron energy spectra as a function of the IAP-IR delay reveals temporal information through delay-dependent photoelectron energy shifts, reducing the measurement of ultrashort time intervals (of the order of 10 as, 1 atomic unit of time = 1 Å. s = 24 as) to conveniently detectable photoelectron energies. Instead of IAPs, RABBITT experiments employ XUV APTs to emit electrons into a synchronized delayed IR pulse, imaging electric dynamics as photoelectron-yield oscillations due to two-pathway two-photon quantum interference [3] (Fig. 1).

While RABBITT spectra have been analyzed to probe isolated atoms in the gas phase for some time [3,8,9], so far their recording and analysis for solid targets has been limited to just a few recent experimental [10–13] and theoretical [14,15] investigations.

While encoding the same basic information on the static and dynamic substrate electronic structure as streaked spectra, RABBITT interferograms are more conveniently obtained in the laboratory, since APTs are easier to generate than IAPs, and require lower IR intensities (typically 10¹¹ W/cm² [10–13]), thus distorting the target less and facilitating their interpretation in terms of two-photon (absorption of one XUV and absorption or emission of one IR photon, as opposed to pathways including more than one IR photon). In order to eliminate the unknown phases {φ₁HHₙ} of the odd-order HH constituents of the ATP, recent experiments subtracted measured “RABBITT phases” (i.e., the phases of delay-dependent yield oscillations in RABBITT interferograms) from RABBITT interferograms obtained either (i) in the same experiment on metal surfaces and a reference gas target [10,11] or (ii) from energetically separable initial states of the same solid target [12]. In addition to temporal information, RABBITT spectra from solid targets reveal details on the spatial distribution of the initial electronic probability density and the dispersion and scattering of APT-excited photoelectrons inside the solid prior to their emission [13,15].

Besides their complex electronic structure, solid targets require the inclusion of macroscopic effects in the reflection of the assisting IR pulse for the computation and analysis of RABBITT spectra [10–12,14,15]. This is of particular importance for the considered copper surfaces that reflect most of the incident IR intensity and entail distinct Fresnel reflection rates for the two incidence angles realized by Lucchini et al. [11] (Fig. 1). In this Rapid Communication we present a quantum-mechanical model for the laser-assisted XUV photoemission from metal surfaces and apply this model to the RABBITT experiment presented in Ref. [11]. Accordingly, we assume the XUV pulse train vector potential Aₜ(XUV,ξ,τ) to be composed of odd HH orders n = 15 to 23, each having a spectral width of 1 eV [10]. The XUV APTs propagates unimpeded through the few monolayers below the copper surface, where photoelectrons are detected [15].

Our numerical results will be shown to allow for (i) the scrutiny of contributions from interfering photoemission pathways involving more than one IR photon and the investigation of the dependence of observable interferograms on the (ii) substrate’s IR dielectric response, (iii) ground-state electronic structure, and (iv) photoelectron dispersion. Unless specified otherwise, we use atomic units throughout.

Integrating around the emission direction Ω_τ = (θ_τ, φ_τ, 0) in the reflection plane over a solid-angle interval ΔΩ_τ = (dθ_τ, dφ_τ) = (10°, 5°) for a given delay τ between the IR pulse and the ATP centers, and for a photoelectron final energy ε_τ = k_τ²/2, incoherent summation over all occupied initial...
valence-band (VB) states below the Fermi level leads to the energy-differential photoemission yield

\[ P(E_f, \tau) = \sqrt{2E_f} \int_{\Delta \Omega_f} d\Omega_f \sum_{|k_i| < k_F} |T_{k_i, k_F}(\tau)|^2, \]  

(1)

where \( k_F \) is the Fermi momentum. Representing the transition-matrix element in the non-dipole velocity gauge,

\[ T_{k_i, k_F}(\tau) \propto \int_{-\infty}^{\infty} dt \langle \Psi_k^f(\mathbf{r}, t, \tau) | A_{\text{XUV}}(z,t) \cdot \nabla | \Psi_k^i(\mathbf{r}, t) \rangle, \]  

(2)

we do not impose restrictions on the size of the illuminated target volume and electron-emission direction [15–17].

We approximate the initial VB state as translationally invariant in the surface (x-y) plane, \( \Psi_k^i(\mathbf{r}, t) = e^{i\mathbf{k}_i \cdot \mathbf{r}} \psi_n, k_i, z(\mathbf{r}) e^{-ik_r t}, \) where \( k_{i, z} = (k_{i, x}, k_{i, y}, z) \) and \( \mathbf{r}_{i, z} = (x, y), \) and expand along the surface normal (z) direction in localized Hulthén generalized-Sturmian functions (GSFs) [20, 21] centered at each atomic plane \( j, \)

\[ \phi_{n, z}(z) = \sum_j e^{ik_{i,j} z} \varphi_n(E_{TB}, \alpha_H, |z - z_j|). \]  

(3)

The localized GSFs

\[ \varphi_n(E_{TB}, \alpha_H, u) \propto e^{-\kappa u} 2F_1[-n, 2\alpha_H \kappa + n, 2\alpha_H \kappa, e^{-u/\alpha_H}], \]  

(4)

model the active electron in the screened Coulomb field of its parent lattice plane, \( V(u) = \frac{e^{-u/\alpha_H}}{\sqrt{2E_{TB}}} \), with an adjustable energy \( E_{TB} \), corresponding to the rate of exponential decay of the electron’s probability density with the distance \( u = z - z_j \), They are written in terms of hypergeometric functions \( 2F_1 \) as solutions of the 1D Hulthén generalized-eigenvalue problem \( -\frac{d^2}{dz^2} + \beta_n V(u) - E_{TB} \varphi_n(E_{TB}, \alpha_H, u) = 0 \) with \( \kappa = \sqrt{2E_{TB}} \) and for \( u \geq 0 \) [20]. We select GSFs with positive parity by defining \( \varphi_n(E_{TB}, \alpha_H, u) \equiv \varphi_n(E_{TB}, \alpha_H, -u) \) for \( z < z_j, \) \( \alpha_H \) determines the nodal structure of \( \varphi_n(E_{TB}, \alpha_H, u) \) near its parent lattice plane. For our numerical applications discussed below, we select Cu(111) 3d orbitals with a single node (setting \( n = 1 \)) and adjust \( \alpha_H = 0.2 \alpha_t \) with the Cu(111) interlayer spacing \( \alpha_t = 3.94 \) [22]. The generalized eigenvalue \( \beta_n \) corresponds to an effective core charge and is determined by the adjustable parameters \( E_{TB} \) and \( \alpha_H \). A similar tight-binding approach [16] based on charge-scaled hydrogenic ground-state wave functions (HGWFs) was found suitable to model Pt(111) core levels (see Refs. [18, 19] with regards to the localization properties of the Cu d band). We note, however, that the present expansion in terms of GSFs with two adjustable parameters allows a more versatile representation of the target electronic structure than the use of HGWFs.

We model the final state based on Volkov functions for the interaction of the free electron with homogeneous continuum-wave electric fields [16],

\[ \psi_k^f(k_x, k_y, z, t) = \frac{f_{k_x, k_y}(z, \tau)}{(2\pi)^{3/2}} e^{i\mathbf{k}_f \cdot \mathbf{r}} \psi_k^f(z, t_0) e^{-i\varepsilon_f t}, \]  

(5)

with \( t_f = t - \tau \) and \( k_f = (k_{f, x}, k_{f, y}, z) \), including three heuristic modifications: (i) Inside the substrate \((z < 0)\), we introduce the energy-dependent mean free path \( \lambda(E_f) \) with the damping factor

\[ f_{k_x, k_y}(z < 0) = e^{z/\lambda(E_f)} \cos(\theta_f), \]  

(6)

in order to account for the loss of emission probability from deeper layers, setting \( f_{k_x, k_y}(z > 0) = 1 \) and using the numerical values for \( \lambda(E_f) \) of Ref. [23]. (ii) With

\[ \psi_k^f(z) = \begin{cases} e^{k_{f, z} z}, z \geq 0 \\ \psi_{\text{well}}^f(z), z < 0, \end{cases} \]  

(7)

we include the energy loss and partial reflection of the photoelectron at the surface \((z = 0)\) in terms of the outgoing plane wave \( e^{i\mathbf{k}_{f, z} \cdot \mathbf{r}} \) with asymptotic momentum component \( k_{f, z} \) that matches with continuous derivative the superposition of a substrate-bound plane wave with momentum \( k^n_{f, z} = \sqrt{(k_{f, z})^2 + 2U_0} \) and its reflection into the bulk at a potential step of height \( U_0 \) at the surface,

\[ \psi_{\text{well}}^f(z) = \cos(k^n_{f, z} z) - \frac{k_{f, z}}{k^n_{f, z}} \sin(k^n_{f, z} z). \]  

(8)

Including the effect of \( U_0 > 0 \) on the final photoelectron state strongly affects photoelectron spectra, as will be shown below. This inclusion of \( U_0 \) in \( \psi_{\text{well}}^f \) extends the representation of the final state in Ref. [16] in terms of exponentially screened Volkov wave functions and is supported by recent experimental evidence for a distinct non-free-electron character of photoelectrons emitted from Ni(111) surfaces [12]. (iii) Generalizing the Volkov phase as

\[ \phi_k^f(z, t) = \int d\tau' \mathbf{k}_{f, z} \cdot \mathbf{A}_{IR}(z, \tau') \psi_k^f(z, \tau') \]  

(9)

with \( \mathbf{A}_{IR} = (A_{IR, x}, A_{IR, y}) \), we allow for inhomogeneous IR vector potentials \( A_{IR}(z, t) = \int d\tau' \mathbf{E}_{IR}(z, \tau') dt' \). The spatial dependence of \( \psi_{\text{well}}^f(z, t) \) is due to both Fresnel reflection of the incident IR pulse and the inclusion of \( U_0 > 0 \). In the limit of homogeneous IR fields \( \mathbf{A}_{IR}(t) \), Eq. (9) yields the phase of Volkov wave functions [16].

We determine the transmitted and reflected electric fields of the assisting IR pulse from Fresnel’s equations [15, 24],
the electric field at the interface, we introduce the gradual macroscopic in nature, implying a discontinuous change of $\mu$ assuming (2), we approximate the IR field screening by a stepwise decay, employing a Lorentz-Drude-model complex-valued dielectric function $\varepsilon(\omega_{IR})$ [25]. Since Fresnel's equations are macroscopic in nature, implying a discontinuous change of the electric field at the interface, we introduce the gradual transition function

$$\mu(z) = e^{(z - z_{IR})/\delta_{IR}}, \ z < z_{IR},$$

exponentially damping the total external IR field inside the substrate. We use the IR-pulse intensity, photon energy, and pulse width specified in Ref. [11] $[5 \times 10^{11} \ W/cm^2, 1.557 eV (786-nm wavelength), and 10 fs]$ and adjust the IR skin depth $\delta_{IR} = 2.25a_0$ and the decay-onset position $z_{IR} = 0.75a_0$ for best overall visual agreement with the experimental interferograms. Evaluating the transition matrix (2), we approximate the IR field screening by a stepwise decay, assuming $\mu(z) \approx \mu(z_j)$ across each atomic layer $j$. This allows us to perform the $z$ and $t$ integrations in (2) analytically.

In Fig. 2, we compare our calculated RABBITT interferograms with the experimental spectra of Ref. [11]. We adjusted the delay axis in our spectra for maximal overlap between the XUV APT and IR pulse to occur at $\tau = -1.8$ fs and $-0.84$ fs, respectively, for $75^\circ$ and $15^\circ$ incidence. For $75^\circ$ incidence, we observe sideband (SB) peaks to have a larger temporal width than HH peaks [Figs. 2(a) and 2(b)] in obvious deviation from the expected [3] $\cos(2\omega_{IR} \phi) \ v$ yield oscillation. We attribute this deviation to interference terms between quantum pathways collectively involving more than two IR photons. We designate the transition matrix element of a photoemission pathway involving the absorption of a $(2n + 1)\omega_{IR}$ XUV photon and the absorption (emission) of $m$ IR photons as $T_{2n+1}^{(m-1)} (T_{2n+1}^{-(m-1)})$. The coherent superposition of all involved $T_{2n+1}^{(m-1)}$ generates the observed photoelectron yield oscillations.

To lowest order, $\cos(2\omega_{IR} \phi)$ higher-order-harmonics (HH)-yield oscillations originate in $T_{2n+1}^{(0)} T_{2n+3}^{(-2)}$ and $T_{2n+1}^{(2)} T_{2n+3}^{(0)}$ interferences. The expected [3] $\cos(2\omega_{IR} \phi) \ v$ oscillations stem from the interference $T_{2n+1}^{(1-1)} T_{2n+1}^{(-1)}$. The narrowing of the HH and widening of SB peaks [relative to $\cos(2\omega_{IR} \phi)$, Figs. 3(a) and 3(b)] thus indicates contributions from pathway combinations adding up to four IR photons. These contributions are less noticeable at $15^\circ$ incidence due to the smaller net IR electric-field component along the electron-emission direction $k_f$ [Figs. 3(c) and 3(d)]. For $15^\circ$ incidence, the generalized Volkov phase (9) in (2) thus oscillates with a smaller amplitude, reducing the number of relevant multipoles (and hence the number of photons) in the expansion $\varepsilon(\phi_{k_f}(z,t_d)) = 1 + i\phi_{k_f}(z,t_d) - \phi_{k_f}(z,t_d) + \ldots$ in (5).

The 4$\omega_{IR}$ HH oscillations result from the three possible combinations of interfering multiphoton processes depicted in Fig. 4(a), $T_{2n+1}^{(-1)} T_{2n+1}^{(-1)}$, and Figs. 4(b) and 4(c), $T_{2n+1}^{(0)} T_{2n+1}^{(-1)}$. We observed the peak heights in Fig. 3(b) of the 4$\omega_{IR}$ peaks to change by roughly 25%, when excluding the pathways involving four IR photons shown in Figs. 4(b) and 4(c), demonstrating that all two-path interferences in Fig. 4 contribute to the observed HH yield. Analogously, the lowest order pathway interferences that lead to 4$\omega_{IR}$ oscillations in a given SB$_{2n}$ are $T_{2n+1}^{(1)} T_{2n+3}^{(1)}$.

In order to assess to what extent RABBITT interferograms are sensitive to the Cu(111) electronic structure, we illustrate in Fig. 5 the variation of simulated spectra with changes of parameters in our model. We find that changing $\lambda(\varepsilon_f)$ with

\begin{align*}
\text{FIG. 3. (a) Measured [11] and calculated oscillations in the photoelectron yields integrated over 0.3-eV intervals around HH$_{19}$ and SB$_{20}$ energies for $75^\circ$ incidence. (b) Fourier transform of panel (a) with frequencies in units of $\omega_{IR}$. [(c), (d)] Same as panels (a) and (b) for $15^\circ$ incidence.}
\end{align*}
respect to the values tabulated in Ref. [23] by a factor of 4 significantly modifies RABBITT spectra, while a factor of 2 does not [Fig. 5(a)]. Figure 5(b) shows energy profiles for 75° incidence calculated for the maximum, minimum, and average bulk value the oscillating Chulkov potential [22]. The theoretical interferograms agree better with the two experiments when the effect of the potential step on \( \Phi_{1f} = \lambda \) of Ref. [8]. In our calculations, we set all XUV HH phases (ii) substituting the Ne atomic phases with calculated phases from Ref. [11]. Experimental yield adapted from Ref. [11].

Lucchini et al. [11] obtained Cu(111) RABBITT phases \( \phi_{2n}^{RAB} \) by (i) eliminating the unknown HH phases using interferograms measured on gaseous Ne as a reference and (ii) substituting the Ne atomic phases with calculated phases of Ref. [8]. In our calculations, we set all XUV HH phases to zero, resulting in RABBITT phases \( \phi_{2n}^{RAB} \) comparable to the experimental values. We obtain \( \phi_{2n}^{RAB} \) by fitting with parameters \( a_{1,\ldots,5} \) our calculated photoelectron yields, integrated over 0.3-eV intervals around the given SB2n energy, to the expression [11]  
\[
[a_{1} \cos (2\omega_{IR}(\tau - a_{3}) + \phi_{2n}^{RAB}) + a_{2}]e^{-\frac{(\omega_{IR} - a_{4})^{2}}{a_{5}}} + a_{5},
\]


For 75° incidence, Fig. 6 reveals that our quantum-mechanical calculations and the semiclassical Monte Carlo (MC) simulations of [11] have an overall shift of approximately \(-0.5\) rad (approximately \(-100\) as) with respect to the experimental phases. Two fundamentally different theories pointing to the same offset relative to the measured Cu(111) phases may indicate either (i) an anisotropic behavior of the dielectric constant not accounted for by our isotropic \( \epsilon(\omega_{IR}) \) model or (ii) the limitation of our translationally invariant initial-state model.

In summary, comparing our calculated RABBITT spectra and phases for Cu(111) with experimental data [11], we find that our localized-orbital initial-state expansion and inclusion of the potential step at the surface in the final state provide spectra in good agreement with the measurements. In addition, our model explains the measured SB temporal-width broadening and HH narrowing as due to four-IR-photon-interference pathways. We find the Cu(111) RABBITT energy profile to be fairly robust with regard to variations of the simulation parameters \( \lambda, U_{0}, \delta_{IR}, \) and \( E_{TB} \). The potential well depth \( U_{0} \) affecting the final state modifies the yields the most. This seemingly simple adjustment to the final state noticeably improves the agreement with the experiment and emphasizes the sensitivity of the RABBITT spectra on the electron-emission dynamics near solid surfaces. An IR-skin-depth increase (decrease) visibly increases (decreases) the SB yields relative to HH yields. Mean-free-path effects on the energy profile are moderate. The most favorable comparison with the experimental energy profile is achieved when the orbital asymptotic decay is characterized by the 3d-band binding energy. Our RABBITT phases are strongly affected by the inclusion of the Fresnel-reflected IR pulse and in reasonable agreement with the classical simulations in Ref. [11].

We thank M. Lucchini and U. Keller for providing experimental data and L. Argenti for useful comments. This work was supported by the Chemical Sciences, Geosciences, and Biosciences Division, Office of Basic Energy Sciences, Office of Science, US Department of Energy under Award DE-FG02-86ER13491 (attosecond interferometry, photoelectron final states in spatially varying external fields) and the U.S. National Science Foundation (NSF) under Award No. PHY 1464417 (general theory and numerical tools development for photoemission from surfaces).


