Extraction of the attosecond time delay in atomic photoionization
using the soft-photon approximation

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We use the soft-photon approximation to extract the Wigner time delay from atomic two-color photoionization experiments. Unlike the strong-field approximation, the present method does not require introduction of the Coulomb-laser coupling corrections and enables one to extract the Wigner time delay directly from attosecond time delay measurements.

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I. INTRODUCTION

The concept of time delay was developed in formal scattering theory by Wigner [1] and his contemporaries (see Ref. [2] for a comprehensive review). It is a quantity related to the phase of the complex scattering amplitude which provides insight into development of the scattering process in time. In recent years, this idea has made a dramatic comeback when it was realized that the time delay can be measured experimentally in photoionization processes. This has led to many interesting and not yet fully understood results such as observation of a considerable time delay between photoelectrons emitted from the 2s and 2p subshells in neon [3] and the experimental determination of the tunneling time in an ionization event [4].

The timing information in photoionization process is extracted experimentally by applying an ionizing XUV pulse (the pump pulse) and an infrared (IR) probe pulse. In the attosecond streaking experiments, the time delay between the pump and probe pulses is mapped onto the kinetic energy of the photoelectron in the form of a spectrogram. In such experiments, duration of the probe pulse may be several optical cycles of the IR field [3]. Alternatively, one may use the so-called reconstruction of attosecond bursts by ionization of two-photon transitions (RABITT) technique [5], which employs a monochromatic IR probe. In this technique, the pump-probe delay is mapped onto the phase of the sideband oscillations caused by interference of alternative two-photon ionization processes. A detailed description of these techniques can be found in Ref. [6].

To extract the Wigner time delay related to the XUV photoionization, one has to take into account the effect of the probe IR field on the system under investigation. In the RABITT experiments with monochromatic probes, the IR field is typically weak, which allows the perturbation theory treatment [7,8]. In the attosecond streaking approach, where the IR probe intensity is typically in the range of $10^{11}$ to $10^{12}$ W/cm$^2$, the nonperturbative treatment is called for. In the first interpretation of the attosecond streaking experiment [9], the well-known classical equation was invoked:

$$ p_f(t) = p_0 - A_{IR}(t), \tag{1} $$

relating the unperturbed asymptotic momentum of the photoelectron $p_0$ and the final momentum $p_f(t)$ for emission at time $t$ in the presence of an IR field $A_{IR}$. This implies that the interaction of the photoelectron with the ionic core is neglected. To account for the corrections due to this interaction (the so-called Coulomb-laser coupling), a further refinement of this model has been developed [10–12]. This refinement results in renormalizing the vector potential in Eq. (1) and shifting its argument by adding the the so-called Coulomb-laser coupling correction in the argument of the function $A_{IR}(t)$. In Ref. [13] the authors were able to derive a similar formula using an analytical approach based on the eikonal approximation. Yet another approach, based on the direct numerical evaluation of the quantity defining the time delay (which is essentially the additional time the particle would spend in a region of space as compared to the free particle) has been proposed in Ref. [14].

Below we present an alternative procedure of extraction of the time delay from the experimentally observable photoionization cross sections. This procedure introduces an accurate description of the IR field influence from the outset and no further corrections are needed.

II. THEORY AND COMPUTATIONAL DETAILS

The procedure is based on the so-called soft-photon approximation [15]. Under condition of the IR photon frequency being small in comparison with the photoelectron energy, this approximation has been shown to reproduce quite accurately the angle-integrated cross sections of the process of two-color ionization by the XUV and IR fields [16]. To extract timing information, one has to know the phase or, rather, the energy derivative of the phase of the amplitude of the ionization process. It is unclear whether the soft-photon approximation can cope with this problem. Below, we address this question.

We consider a typical configuration of the XUV and IR fields used in the attosecond streaking experiments. The time dependence of the electric field of the IR pulse is

$$ E_{IR}(t) = E_0 \sin \Omega t, \tag{2} $$

with the base frequency $\Omega = 0.057$ a.u. (photon energy of 1.55 eV). The IR field is present on the interval of time $(0, T_1)$, where $T_1 = 2\pi/\Omega = 2.7$ fs is an optical cycle corresponding to the IR frequency $\Omega$.

The XUV pulse is present on the time interval $\Delta = 4T, \Delta + 4T$, where $T = 2\pi/\omega$ is an optical cycle of the XUV pulse. Parameter $\Delta$, therefore, characterizes the relative shift between beginning of the IR pulse and arrival of the center of the XUV pulse. We use the soft-photon approximation to extract the Wigner time delay from atomic two-color photoionization experiments. Unlike the strong-field approximation, the present method does not require introduction of the Coulomb-laser coupling corrections and enables one to extract the Wigner time delay directly from attosecond time delay measurements.

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pulse. On this interval the XUV field time dependence is

$$\mathcal{E}_{\text{XUV}}(t) = \mathcal{E}_{\text{XUV}}^0 f(t') \cos \omega t',$$  

where $t' = t - \tau$, and we use a cosine squared envelope function $f(t') = \cos^2(\omega t'/16)$. The XUV field strength is $\mathcal{E}_{\text{XUV}}^0 = 0.01$ a.u. (intensity of $3.5 \times 10^{12}$ W/cm$^2$). Both pulses are assumed to be linearly polarized along the z axis. As a target system, we consider the Ne atom described by a localized model potential [17] within the single active electron (SAE) approximation.

The amplitude of the photoionization process can be defined as

$$f(k) = \lim_{t \to -\infty} e^{i[E(k) t - E_0 t]} \langle \Psi_k^- | \hat{U}(t, \tau) \phi \rangle,$$  

where $\Psi_k^-$ is the (ingoing) scattering wave function describing the photoelectron with the kinetic energy $E(k)$, $\hat{U}(t, \tau) = \exp(-i H_0 t) \exp(-i H(t, \tau) t)$. The perturbative expression (5) allows us to express $f^{\text{XUV}}(k, \Delta)$ in terms of the “reference” amplitude $f^0(k, \Delta)$ as $f^{\text{XUV}}(k, \Delta) = e^{iE(k)-E_0 \Delta} f^{\text{XUV}}(k, 0)$. We can thus relate the two-color amplitudes $f(k, \Delta)$ and the reference amplitude $f^{\text{XUV}}(k, 0)$ for different values of $\Delta$.

Our goal is to determine the phase, or rather the phase derivative, of the reference amplitude with respect to the electron momentum, since the quantity of interest for us, the time delay $\tau_0$, can be expressed as [20]

$$\tau_0 = \frac{1}{k_z} \text{Im} \left( \frac{\partial f^{\text{XUV}}(k, 0)}{\partial k_z} \right).$$  

Here the derivative is to be taken at the point $k_z$ satisfying the energy conservation $E_0 + \omega = k_z^2/2$, $E_0$ being the energy of the initial atomic state. By using this equation and Eq. (8), it is not difficult to devise a procedure to obtain information about the phase of the reference amplitude for the process of photoionization by the XUV field from the experimentally measurable cross sections of the photoionization process in the presence of both the XUV and IR fields. Before describing implementation of such a procedure, we have to ascertain first that Eq. (8) is accurate enough.

### III. Numerical Results

We solve the time-dependent Schrödinger equation (TDSE) for the Ne atom described by means of a localized one-electron potential [17] in the presence of the XUV and IR fields given by Eqs. (2) and (3). We employ the procedure to solve numerically a three-dimensional (3D) TDSE, which is described in detail in Refs. [20,21]. By projecting the solution of the TDSE on the scattering state $\Psi_k^-$ of the Ne atom, as prescribed by Eq. (4), we obtain the photoionization amplitude $f(k)$ in the presence of both the XUV and IR fields. A separate calculation of atomic evolution in the presence of the XUV pulse alone described by Eq. (3) with $\Delta = 0$ gives us a “reference” amplitude $f^{\text{XUV}}(k, 0)$. By using the relation connecting $f^{\text{XUV}}(k, \Delta)$ and $f^{\text{XUV}}(k, 0)$ and Eq. (8), we can compute values of $f(k, \Delta)$, which is the amplitude of the two-color ionization for different values of the delay $\Delta$ between the XUV and IR pulses, and compare them with the ab initio values of $f(k, \Delta)$ provided by the TDSE calculation. Such a comparison is shown in Figs. 1–5. The data were obtained retaining the terms with $|m| \leq 3$ in Eq. (8) for $\mathcal{E}_{\text{IR}}^0 = 0.001$ a.u. and $|m| \leq 5$ for $\mathcal{E}_{\text{IR}}^0 = 0.004$ a.u. The number of the terms which are to be retained in Eq. (8) depends on the strength of the IR field and can be easily determined for each particular value of $\mathcal{E}_{\text{IR}}^0$ by checking that convergence with respect to $m$ is achieved.

The data displayed in these figures show that Eq. (8) allows us to compute values of the two-color ionization amplitude $f(k, \Delta)$ with a reasonable accuracy for ionization from 2$s$ and 2$p$ states of Ne provided we know the reference amplitude $f^{\text{XUV}}(k, 0)$ as a function of the momentum. Results, as can be surmised from the figures, are slightly more accurate for smaller IR field intensities.

We now try to solve an inverse problem of the reconstruction of the amplitude $f^{\text{XUV}}(k, 0)$ provided that absolute values

By using Eq. (8) for various delays $\Delta$ between the IR and XUV fields, we can obtain a set of relations between $f(k, \Delta)$ and $f^{\text{XUV}}(k, \Delta)$. The perturbative expression (5) allows us to express $f^{\text{XUV}}(k, \Delta)$ in terms of the “reference” amplitude $f^{\text{XUV}}(k, 0)$ as $f^{\text{XUV}}(k, \Delta) = e^{iE(k)-E_0 \Delta} f^{\text{XUV}}(k, 0)$. We can thus relate the two-color amplitudes $f(k, \Delta)$ and the reference amplitude $f^{\text{XUV}}(k, 0)$ for different values of $\Delta$.

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of the two-color amplitudes $f(k_z, \Delta)$ are known for some selected values of the delays $\Delta$ and momenta $k_z$. This can be demonstrated as follows. We choose a trial form for the amplitude $f^{XUV}(k_z, 0)$:

$$f^{XUV}(k_z, 0) = Ae^{-a(E-\epsilon)^2+i\tau(E-\epsilon)},$$  

where $E = k_f^2/2$; $a$, $\tau$, and $\epsilon$ are fitting parameters; and $A$ is a complex number which does not depend on the energy $E$. Parameter $\epsilon$ has a meaning of the energy at which the cross section of the photoionization by the XUV pulse is peaked. The first guess for the value of this parameter can be obtained from the energy conservation $E_0 + \omega = \epsilon_0$. We could fix the
value of this parameter to $\epsilon_0$. However, more accurate results are obtained if we treat it as a fitting parameter. The parameter $\tau$, as can be immediately seen from the Eq. (9), has a meaning of the time delay.

The ansatz (10) does a very good job at reproducing the amplitude $f^{\text{XUV}}(k_z,0)$, as Fig. 6 testifies. This figure shows comparison of a fit using the functional form (10) to the “exact” amplitude $f^{\text{XUV}}(k_z,0)$, which we obtain from the TDSE solution for ionization of the 2s subshell of the Ne atom by the XUV pulse.

By using Eqs. (8) and (10), we can compute the trial amplitude $f'(k_z,\Delta)$ and, consequently, the trial electron spectrum $P'(k_z,\Delta)$ of the two-color ionization of the Ne atom as a function of the momentum for various values of the delay $\Delta$ between the IR and XUV pulses. Using this distribution we can compute the trial expectation values of the electron momentum $\tilde{k}_z(\Delta)$ for various $\Delta$.

$$\tilde{k}_z(\Delta) = \int P'(k_z,\Delta) k_z \, dk_z,$$

and compare them with the values $\tilde{k}_z(\Delta)$ which we obtain from the TDSE calculation (and which can be measured in the experiment).

We can now form a functional:

$$D(a,\tau,\epsilon) = \sum_i (\tilde{k}_z(\Delta_i) - \tilde{k}_z(\Delta_i))^2.$$  

(12)

Sum in Eq. (12) is taken over a set of delays $\Delta$ between the IR and XUV pulses, for which the data are available. Presently, we use the set $\Delta = 0.2T_1, 0.3T_1, 0.4T_1, 0.5T_1,$ and $0.7T_1$. By minimizing the functional (12) with respect to the parameters $a$, $\tau$, and $\epsilon$ in Eq. (10), we obtain the time delay $\tau$.

Convergence of this procedure is illustrated in Fig. 7 displaying the function $D^{-1}(a,\tau,\epsilon_0)$ for fixed value of $\epsilon = \epsilon_0$, where $\epsilon_0 = E_0 + \omega$ is the value given by the energy conservation. This figure shows that $D^{-1}(a,\tau,\epsilon_0)$ has a well-defined pronounced maximum in the space of the parameters $a, \tau$. This property is very useful since it implies that $D^{-1}(a,\tau,\epsilon)$ in Eq. (12) has a deep minimum, which is easy to locate.
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FIG. 5. (Color online) Ionization from the 2p state of Ne atom. Re $f(k_z, \Delta)$ computed using Eq. (8), solid (red online) line, and TDSE calculation, dashed (green online) line. Im $f(k_z, \Delta)$ given by Eq. (8), short dashed (blue online) line, and TDSE, dotted (magenta online) line. XUV photon energy $\omega = 106$ eV; $\varepsilon_0^lf = 0.004$ a.u. Delays $\Delta$ (top to bottom) are $\Delta = 0.37\tau$, $\Delta = 0.47\tau$, $\Delta = 0.57\tau$, where $T$ is an optical cycle of the XUV pulse. Vertical solid line corresponds to the momentum $k_z$ for which $E_0 + \omega = k_z^2/2$.

Results of the procedure based on the minimization of the functional (12) are illustrated in Fig. 8, where we present the data for the time delay for several base frequencies $\omega$ of the XUV pulse for ionization from the 2s and 2p subshells of the Ne atom. These results are compared with the values for the time delays which we can extract directly from the TDSE calculation using the computed amplitudes of XUV photoionization and Eq. (9).

Figure 8 indicates that the results of the fitting procedure described above agree well with the results of the ab initio TDSE calculation. For completeness, in the same figure we display the time delay results obtained from the Hartree-Fock elastic scattering phases $\tau^{HF} = d\delta^{HF}/dE$. These phases are calculated in the frozen-core Hartree-Fock approximation to electron scattering in the field of the Ne$^+$ ion [22]. The scattering phase in the dominant photoionization channel $l = l_i + 1$ is taken according to the Fano propensity rule [23], where $l_i$ is the angular momentum of the initial bound state. Although these results are not directly comparable to the present calculations, which employ a localized effective potential, they demonstrate a qualitatively similar dependence of the time delay on the photon energy. We should emphasize that it is the Hartree-Fock results only that rely explicitly on the Fano propensity rule. In contrast, the time delay defined by Eq. (9) is evaluated using the full amplitude $f_{XUV}(k_z, 0)$. In the partial wave expansion of the scattering state $\psi_0^f$, entering this amplitude, both the channels $l = l_i \pm 1$ are included.

We may note that the present method of extraction of the attosecond time delay, based on the soft-photon approximation, can be linked to the approach developed in Refs. [10–12].

FIG. 6. (Color online) Re $f_{XUV}(k_z, 0)$ computed using Eq. (10), solid (red online) line, and TDSE calculation, dashed (green online) line. Im $f_{XUV}(k_z, 0)$ given by Eq. (10), short dashed (blue online) line, and TDSE, dotted (magenta online) line. XUV photon energy $\omega = 106$ eV; ionization from the 2s state of Ne. Vertical solid line corresponds to the momentum $k_z$ for which $E_0 + \omega = k_z^2/2$.

FIG. 7. (Color online) Quantity $D^{-1}(a, \tau, \varepsilon_0)$ as function of $a$ and $\tau$. Value of $\varepsilon_0$ is fixed at the energy conservation value $\varepsilon_0 = E_0 + \omega$.

FIG. 8. (Color online) Time delays computed using Eq. (9) and the fitting procedure. Ionization from the 2s state: empty box (red online), Eq. (9); solid box (green online), the fitting procedure. Ionization from the 2p state: empty circle (magenta online), Eq. (9); solid circle (blue online), the fitting procedure. The HF results: $2s \rightarrow E_p$ transition, solid line (red online); $2p \rightarrow Ed$ transition, dashed line (blue online).
which is a refinement of the strong-field approximation Eq. (1). As we mentioned above, this refinement consists in introducing the correction factor, multiplying the vector potential in Eq. (1), and adding the so-called Coulomb-laser coupling correction in the argument of the function $A^{\text{IR}}(t)$. It is easy to see from Eqs. (8) and (10) that the photoelectron spectrum of the two-color ionization $P(k, \Delta)$ in the soft-photon approximation can be represented as $P(k, \Delta) = h(k, \Delta + \tau)$, where $\tau$ is the time delay and $h$ is a periodic function of the second argument with the period $T_1$ equal to the optical cycle of the IR field. The first moment of $P(k, \Delta)$ can therefore be represented as $\bar{k}_c(\Delta) = q(\Delta + \tau)$, where $q$ is periodic with period $T_1$. We can write, therefore,

$$\bar{k}_c(\Delta) = \sum_{n=0}^{\infty} B_n \sin \Omega(\Delta + \tau) + C_n \cos \Omega(\Delta + \tau).$$

(13)

The vector potential of the IR field described by Eq. (2) is $A^{\text{IR}}(t) = \frac{eA^2}{c^2}(1 - \cos \Omega t)$. By retaining the terms with $n = 0, 1$ in Eq. (13) we obtain

$$\bar{k}_c(\Delta) - b \approx cA^{\text{IR}}(\Delta + \tau + \delta),$$

(14)

where the coefficients $b, c, \text{and} \delta$ can be expressed in terms of the coefficients of the Fourier expansion (13). By identifying the coefficients $b, c, \text{and} \delta$ in the Eq. (14) with $k_0$, the correction factor multiplying the vector potential in Eq. (1), and the Coulomb-laser coupling correction, we obtain the equation replacing the strong-field relation (1) in the approach developed in Refs. [10–12].

**IV. CONCLUSION**

In the present work, we examined applicability of the soft-photon approximation for evaluation of the amplitudes of two-color XUV and IR ionization. We have found that the two-color ionization amplitudes, computed using the soft-photon approximation, agree well with the \textit{ab initio} TDSE amplitudes. This fact can be used to extract phase information and, in particular, the time delay from the experimental photoelectron spectra detected in attosecond streaking measurements.

We tested the range of validity of the soft-photon approximation. We demonstrated that this approximation renders the two-color ionization amplitudes accurately for the IR field intensities in the range $E^{\text{IR}} = 0.001-0.004 \text{ a.u.}$. The latter can be furnished by various computational methods. In [26] we obtained the “bare” amplitude by solving the TDSE with the \textit{ab initio} Hamiltonian of the He atom.

In principle, the soft-photon approximation can be used for theoretical studies of two-color ionization in systems described by \textit{ab initio} Hamiltonians. In Ref. [26], we performed such a study for the process of double ionization of the He atom. The formulas for the amplitudes of the two-color ionization in this case are quite similar in structure to Eq. (8). In order to obtain the ionization amplitude in presence of the IR and XUV fields, one needs to know the “bare” amplitude of the ionization by the XUV field alone as a function of the photoelectron energy. The latter can be obtained by various computational methods. In [26] we obtained the “bare” amplitude by solving the TDSE with the \textit{ab initio} Hamiltonian of the He atom. We found that the two-color amplitudes, obtained in the soft-photon approximation, and the \textit{ab initio} ones, provided by the TDSE calculation, agree quite well. In Ref. [26] our concern was to reconstruct the phase of the amplitude from the experimentally observable data. With minor modifications, we can apply this procedure to study time delays for the double ionization in a completely \textit{ab initio} setting.

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