Time delay in atomic photoionization with circularly polarized light

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We study time delay in atomic photoionization by circularly polarized light. By considering the Li atom in an excited 2p state, we demonstrate a strong time delay asymmetry between the photoemission of the target electrons that are co- and counter-rotating with the electromagnetic field in the polarization plane. In addition, we observe the time delay sensitivity to the polar angle of the photoelectron emission in the polarization plane. This modulation depends on the shape and duration of the electromagnetic pulse.

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I. INTRODUCTION

Atomic or molecular photoionization in a circularly polarized electromagnetic (EM) field exhibits a number of features that are absent in the case of linear polarization. One such effect is dependence of the photoionization probability on the direction of the rotation of the target electron relative to the polarization plane of light. Let us assume that the EM field is right circularly polarized in the (x, y)-plane. We call the target electron co- or counter-rotating with the field if its angular momentum projection on the z-axis is m > 0 or m < 0, respectively. This difference of ionization probabilities for coor counter-rotating electrons is present both in the multiphoton and tunneling regimes. However, it manifests itself quite differently. In the multiphoton regime, the ionization probability is larger for the co-rotating electrons [1, 2]. On the contrary, in the tunneling regime, ionization for the counter-rotating electron dominates [3]. The latter effect may have important implications for the angular attosecond streaking technique, which makes it possible to trace electronic motion in atoms and molecules with the resolution of several attoseconds [4].

The earlier works [1-3] focused on the asymmetry in photoionization cross-sections for the co- and counterrotating target electrons. In the present study, we supplement these works by the time delay analysis. The notion of time delay was initially introduced by Wigner [5] for scattering phenomena and applied subsequently to photoionization [6]. A number of benchmark experimental and theoretical results have been obtained in studies of atomic photoionization using this approach [4, 7–9]. These studies have been extended to molecules [10] and double electron photoionization [11]. In the present work, we calculate the time delay following atomic photoionization by a circularly polarized EM field. For a numerical illustration, we consider ionization of the lithium atom prepared in the excited 2p state. Such a state is readily experimentally accessible (see e.g. [12]). Our particular interest will be in the time delay asymmetry between the co- and counter-rotating target electrons with $m = \pm 1$.

The paper is organized as follows. In Sec. II we outline our theoretical model. In Sec. III we present our numerical results for the photoelectron spectrum, momentum distribution and time delay. We conclude in Sec. IV by outlining the key features of the observed effects. The atomic units are used throughout the paper unless otherwise specified.

II. THEORETICAL MODEL

We solve the time dependent Schrödinger equation (TDSE) for an atom with a single active electron

$$i \frac{\partial \Psi(\boldsymbol{r})}{\partial t} = \left(\hat{H}_{\text{atom}} + \hat{H}_{\text{int}}(t)\right) \Psi(\boldsymbol{r}) .$$
 (1)

Here \hat{H}_{atom} is the Hamiltonian of the field-free atom with an effective one-electron potential [13]. $\hat{H}_{\text{int}}(t)$ describes the interaction with the EM field, either in the length or velocity gauges:

$$\hat{H}_{\rm int}(t) = \begin{cases} \boldsymbol{E}(t) \cdot \hat{\boldsymbol{r}} \\ \boldsymbol{A}(t) \cdot \hat{\boldsymbol{p}} \end{cases}, \text{ where } \boldsymbol{A}(t) = -\int_{-T_1/2}^{t} \boldsymbol{E}(\tau) \ d\tau$$
(2)

The field is right circularly polarized propagating along the z-direction (which is assumed to be the quantization axis):

$$E_x = \mathcal{E}f(t)\cos\omega t$$
, $E_y = \mathcal{E}f(t)\sin\omega t$. (3)

Here $f(t) = \cos^2(\pi t/T_1)$ is the pulse envelope. The field is present on the interval $(-T_1/2, T_1/2)$. We measure the pulse duration in units of the optical cycle T_1/T where $T = 2\pi/\omega$.

The solution of Eq. (1) is sought in the form of a partial wave expansion

$$\Psi(\mathbf{r},t) = \sum_{l=0}^{L_{\max}} \sum_{\mu=-l}^{l} f_{l\mu}(r,t) Y_{l\mu}(\theta,\phi), \qquad (4)$$

The radial part of the TDSE is discretized on the grid with the stepsize $\delta r = 0.05$ a.u. in a box of the size

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 R_{max} . Convergence of Eq. (4) with L_{max} and R_{max} depends on the field strength \mathcal{E} . This convergence is tested thoroughly in each numerical example presented below. To propagate the wave function (4) in time, we use the matrix iteration method developed in [14] and further tested in strong field ionization calculations [15, 16].

We project the solution of the TDSE at the end of the laser pulse at $t = T_1/2$ on the set of the ingoing scattering states:

$$\psi_{\mathbf{k}}^{(-)}(\mathbf{r}) = \sum_{l\mu} i^{l} e^{-i\delta_{l}} Y_{l\mu}^{*}(\hat{\mathbf{k}}) Y_{l\mu}(\hat{\mathbf{r}}) R_{kl}(r) , \qquad (5)$$

which gives us a set of coefficients

$$a_{\boldsymbol{k}} = \langle \psi_{\boldsymbol{k}}^{(-)} | \Psi(t) \rangle e^{iE_{\boldsymbol{k}}t} .$$
 (6)

The squared moduli of these coefficients determine the photoelectron spectrum. The energy derivative of the phase of the coefficients (6) gives the photoelectron group delay which is also known as the Wigner time delay. An equivalent form, which is more convenient in practical calculations, is the following:

$$\tau_0 = \operatorname{Im}\left(\frac{\hat{\boldsymbol{q}}}{q}\frac{\partial a_{\boldsymbol{k}}}{\partial \boldsymbol{k}}\right) \ . \tag{7}$$

Here the derivative is computed at the point $\mathbf{k} = \mathbf{q}$, corresponding to the asymptotic momentum of the photoelectron in the field-free zone [16–18]. We notice, that when a single partial wave l in the sum (5) is dominant, the photoelectron time delay is simply the energy derivative of the corresponding elastic scattering phase shift $\tau_0 = d\delta_l/dE = k^{-1}d\delta_l/dk \times 24$ as. Here one atomic unit of time is equated to 24 as.

The time delay τ_0 has a transparent physical meaning [5, 6]. It appears as a coefficient in the asymptotic expression for the trajectory of the crest of the wavepacket

$$\boldsymbol{r}(t) \sim \boldsymbol{q}(t - \tau_0) + \boldsymbol{r}'(t) \ . \tag{8}$$

Here the term $\mathbf{r}'(t)$ gives a well-known Coulomb corrections to the trajectory which grows logarithmically with t for large t.

III. NUMERICAL RESULTS

For numerical illustrations, we perform photoionization calculations of the lithium atom in the 2p excited state. We use circular polarized field pulses with the carrier frequency $\omega = 0.5$ a.u. (13.6 eV) and the pulse durations T_1 varying from three to ten optical cycles (1 to 3 fs). The field strength ranges from $\mathcal{E} = 0.01$ to 0.1 a.u. which corresponds to the field intensity of 3.5×10^{12} to 3.5×10^{14} W/cm².



FIG. 1: (Color online) The photoelectron spectrum of Li ionization from the $2p_{\pm 1}$ state. The field parameters are $\omega = 0.5$ au, $\mathcal{E} = 0.1$ au and the pulse duration is 10 optical cycles. The calculations with the following pairs of values $L_{\max}/R_{\max}(a.u.)$ are plotted. Solid (red) line: 12/500, (blue) short dash: 10/500, (magenta) dots: 10/900, all in the velocity gauge; dashed (green) line: 12/500, the length gauge.

A. Photoelectron spectrum

We start our calculations with the photoelectron spectrum integrated over all the escape directions. We test convergence of our results with respect to the values of L_{max} in Eq. (4) and the box size R_{max} as well as the gauge invariance between the length and velocity forms of the atom- EM field interaction (2). We illustrate this test in Figure 1 using the most challenging condition of the strong field ionization with $\mathcal{E} = 0.1$ a.u. The pulse duration is 10 optical cycles. In this figure we show the photoelectron spectrum following ionization of the $2p_{\pm 1}$ initial state. The velocity gauge results with the pair of values $L_{\rm max}/R_{\rm max}({\rm a.u.}) = 12/500, 10/500$ and 10/900 as well as the length gauge result with $L_{\rm max}/R_{\rm max}({\rm au}) = 12/500$ are indistinguishable on the scale of the figure. A small peak at higher electron energy of $\simeq 23$ eV is a manifestation of a two-photon absorption. Such two-photon processes, as can be seen from Figure 1, are not very important as long as we are interested in the electron spectra only. As we shall see below, the situation is rather different if we are interested in computing the time delay.

B. Photoelectron momentum distribution

In this and the following section, we confine our study to the equatorial (x, y) plane in which most of the photoelectrons escape. In Figure 2 we present the photoelectron momentum distributions in this plane for various field strengths and pulse durations. The upper row of Figure 2 shows the photoelectron momentum distribution obtained for the weak field of $\mathcal{E} = 0.01$ a.u. and the pulse length of 3 optical cycles. This weak field regime can be analyzed within the lowest order perturbation theory (LOPT). Under this condition, an electron in the $2p_{+1}$



FIG. 2: (Color online) The photoelectron momentum distributions in the equatorial (x, y) plane for Li ionization from the $2p_{+1}$ state (left column) and $2p_{-1}$ state (right column). The field parameters are $\omega = 13.6$ eV, the pulse duration is 3 optical cycles (upper three rows) and 10 optical cycles (bottom row), $\mathcal{E} = 0.01$ a.u. (first row), $\mathcal{E} = 0.05$ a.u. (second row), $\mathcal{E} = 0.1$ a.u. (third and fourth rows).

initial state, upon absorption of a photon from the circularly polarized EM field Eq. (3), ends up in the *d* partial wave with the angular momentum projection $\mu = 2$. An electron ionized from the $2p_{-1}$ state ends up in the superposition of the continuum *s* and *d* states with $\mu = 0$. In both cases, the photoelectron angular distribution in the equatorial plane will be independent on the polar angle ϕ , which is clearly seen in Figure 2.

Results for the higher field strengths are shown in the second row ($\mathcal{E} = 0.05$ a.u.) and third and fourth rows ($\mathcal{E} = 0.1$ a.u.). Unlike in the weak field regime, the photoelectron angular distributions for a short pulse of 3 optical cycles show some angular structure which is due

to the higher order processes.

C. Photoelectron time delay

In Figure 3 we present the time delay results as a function of the polar angle for the photoelectrons escaping in the equatorial plane. Also shown are the energy derivatives of the scattering phase shifts in the s and d partial waves, computed at the energies corresponding to the peak of the photoelectron wavepacket in the momentum space. The three sets of panels display the time delay results for the field strengths of $\mathcal{E} = 0.01$ a.u. (top), $\mathcal{E} = 0.05$ a.u. (middle), and $\mathcal{E} = 0.1$ a.u. (bottom). The pulse duration is $T_1/T = 3$ and 10 for the left and right sets of panels, respectively.

Let us first discuss the weak field regime of $\mathcal{E} = 0.01$ a.u. for which the perturbative treatment is applicable. Unlike the photoelectron angular distributions displayed in Figure 2, the time delay results for the short pulse duration show a noticeable angular anisotropy. The angular dependence of the time delay has a character of a π -periodic modulation. This effect depends on the pulse duration, and disappears completely for a longer pulse of $T_1/T = 10$. Another observation is that for all the pulse lengths and pulse strengths, the time delay for the corotating electron $2p_{+1}$ is larger than the time delay for the electron ionized from the $2p_{-1}$ state.

The origin of these effects can be elucidated most readily within the LOPT. The corresponding expression for the ionization amplitude a_k reads:

$$a_{\mathbf{k}} = -i \int_{-T_1/2}^{T_1/2} \langle \psi_{\mathbf{k}}^{(-)} | \hat{H}_{\text{int}}(\tau) | \psi_i \rangle e^{i(E_k - E_0)\tau} d\tau , \qquad (9)$$

where ψ_i is the initial electron state. By plugging expressions (3) into the electromagnetic interaction operator $\hat{H}_{int}(t)$ in the length gauge, and writing the scalar product $\boldsymbol{E} \cdot \boldsymbol{r} \propto E_x x + E_y y = -E_1 r_{-1} - E_{-1} r_1$ in the spherical coordinates [19], we obtain:

$$\hat{H}_{\rm int}(t) = \frac{\mathcal{E}}{\sqrt{2}} f(t) \left(e^{i\omega t} r_{-1} - e^{-i\omega t} r_1 \right)$$
(10)

By using the partial wave expansion (5) for the scattering state $\psi_{\mathbf{k}}^{(-)}$ and setting the initial electron state to $2p_{\pm 1}$ we transform the perturbative expression (9) into

$$a_{\boldsymbol{k}} = \sum_{l=0,2} \sum_{\mu=-l}^{l} i^{-l} e^{i\delta_l} Y_{l\mu}(\hat{\boldsymbol{k}}) \langle kl ||r||2p\rangle$$

$$\times \left[A \begin{pmatrix} l & 1 & 1 \\ -\mu & -1 & m \end{pmatrix} - B \begin{pmatrix} l & 1 & 1 \\ -\mu & 1 & m \end{pmatrix} \right]$$
(11)

where

$$A = \frac{\mathcal{E}}{\sqrt{2}} \int_{-T_1/2}^{T_1/2} f(t) e^{i(E_k + \omega - E_0)t} dt$$
$$B = \frac{\mathcal{E}}{\sqrt{2}} \int_{-T_1/2}^{T_1/2} f(t) e^{i(E_k - \omega - E_0)t} dt$$



FIG. 3: (Color online) The photoelectron time delays as functions of the polar angle ϕ in the equatorial plane for the field strengths $\mathcal{E} = 0.01$ a.u. (top), $\mathcal{E} = 0.05$ a.u. (middle), and $\mathcal{E} = 0.1$ a.u. (bottom). Pulse duration: $T_1/T = 3$ (left), $T_1/T = 10$ (right). (Red) solid line: ionization from the $2p_{+1}$ state, (green) dash: ionization from $2p_{-1}$ state. (Blue) short dash: $\tau_0^d = d\delta_2/dE$; (magenta) dots: $\tau_0^s = d\delta_0/dE$

and the reduced matrix element $\langle kl || r || n_i l_i \rangle$ is given by:

$$\langle kl \| r \| n_i l_i \rangle = \hat{l} \hat{l}_i \begin{pmatrix} l & 1 & l_i \\ 0 & 0 & 0 \end{pmatrix} \int r^2 dr \, R_{kl}(r) \, r \, R_{n_i l_i}(r) \,, \tag{12}$$

with $\hat{l} = \sqrt{2l+1}$. The terms with the coefficients A and B in the square brackets on the right hand side of Eq. (12) describe emission and absorption of a photon, respectively.

It is easy to see that A = 0 on the energy shell where $E_q = q^2/2 = \omega + \epsilon_i$. This property does not depend on the particular form of the envelope function f(t) or the pulse length T_1 in Eq. (3) and it holds as long as the envelope function is even on the interval of the pulse duration. On the energy shell Eq. (12) can be rewritten, therefore, as:

$$a_{\boldsymbol{q}} = \begin{cases} \alpha_{0,0,-1} Y_{00}(\hat{\boldsymbol{q}}) + \alpha_{2,0,-1} Y_{20}(\hat{\boldsymbol{q}}) &, \ m = -1 \\ \alpha_{2,2,1} Y_{22}(\hat{\boldsymbol{q}}) &, \ m = +1 \end{cases}$$
(13)

where

$$\alpha_{l,\mu,m} = -Bi^{-l}e^{i\delta_l}\langle ql||r||2p\rangle \begin{pmatrix} l & 1 & 1\\ -\mu & 1 & m \end{pmatrix}$$
(14)

Dependence of the amplitudes on the direction of the photoelectron escape is contained in the spherical harmonics appearing in Eq. (13). If we consider the modulus squared of the amplitude in the Eq. (13) in the equatorial plane, we will obtain the probability independent of the polar angle, a LOPT prediction which the TDSE calculation for weak fields illustrated in Figure 2 conforms to.

Situation is different for the time delay calculations. Indeed, according to Eq. (7), to compute the time delay we have to calculate energy derivative of the amplitude. This includes the energy derivative of the coefficients Aand B in Eq. (12). It is easy to see that on the energy shell dB/dE = 0, while $dA/dE \neq 0$. Similarly to the statements $A = 0, B \neq 0$, these are quite general properties, which do not depend on the particular pulse shape as long as the envelope function is an even function of time. Moreover, dA/dE depends rather sensitively on the pulse duration as shown in Figure 4.



FIG. 4: (Color online) Coefficients -dA/dE (red solid line) and B(E) (green dashed line) on the energy shell as functions of the pulse duration T_1/T . The envelope function f(t) is defined in Eq. (3).

For the energy derivative of the ionization amplitude from the $2p_{-1}$ state we then obtain:

$$\frac{da_{\boldsymbol{q}}}{dE} = \frac{d\alpha_{0,0,-1}}{dE}Y_{00}(\boldsymbol{q}) + \frac{d\alpha_{2,0,-1}}{dE}Y_{20}(\boldsymbol{q}) + \eta Y_{2-2}(\boldsymbol{q}),$$
(15)

where the coefficient η is proportional to the derivative dA/dE computed on the energy shell. In the equatorial plane, the presence of the term with $Y_{2-2}(q)$ introduces a modulation proportional to $\exp(-2i\phi)$. This π -periodic modulation will also be present in the time delay (7), which is a logarithmic derivative of the amplitude. The depth of the modulation is determined by the value of the derivative dA/dE on the energy shell which, in turn, is determined by the particular pulse shape and pulse duration. As we observe in Figure 4, the derivative dA/dE decreases fast with the pulse duration. Consequently, the depth of the modulation decreases with increasing pulse length.

No modulation effect is present in the $2p_{+1}$ case under the perturbative regime. Emission of a photon in this case leads to a continuum state with $l = \mu = 2$, which



FIG. 5: (Color online) Top: scattering phase shifts $\delta_{l=0}$ (red dotted line) and $\delta_{l=2}$ (blue solid line) ionization channels. Bottom: the energy derivatives $d\delta_{l=0}/dE$ (red dotted line) and $d\delta_{l=2}/dE$ (blue solid line) are converted to the units of time delay.

does not introduce angular dependence for the time delay. These LOPT effects are clearly seen in our TDSE calculation in the weak field regime which are illustrated on the top row of panels in Figure 3. The π -modulation of the time delay for the $2p_{-1}$ is clearly visible for the 3-cycle laser pulse, and disappears completely for the 10cycle pulse.

As for the relative values of the time delays for ionization from $2p_{\pm 1}$ states, this question can be easily addressed using the LOPT expressions. Formulas are simpler for the case of a long pulse, when the term with the coefficient η is negligible in Eq. (15). In this case the time delay for the ionization from the $2p_{+1}$ state is simply $\tau_0(2p_{+1}) = d\delta_2/dE$. The time delay ratio between the $2p_{-1}$ and $2p_{+1}$ states can be expressed as

$$\tau_0(2p_{-1})/\tau_0(2p_{+1}) \approx \frac{(1+\gamma\cos\Delta)}{(1+\gamma\cos\Delta)^2 + \sin^2\Delta} < 1$$
, (16)

where $\gamma = \alpha_{0,0,-1}Y_{00}(\boldsymbol{q}) \left[\alpha_{2,0,-1}Y_{20}(\boldsymbol{q})\right]^{-1}$ and $\Delta = \delta_0 - \delta_2$. Here we used the fact that for the presently considered photoelectron energies $d\delta_2/dE \gg d\delta_0/dE$.

This property of the scattering phase shifts in the field of the Li⁺ ion are illustrated in Figure 5. At lower photoelectron energies, the scattering phases both in the *s*-wave (l = 0) and the *d*-wave (l = 2) display the Coulomb singularity. At larger photoelectron energy, δ_0 acquires a constant shift due to an occupied $1s^2$ orbital of the matching angular momentum in the ionic core (the Levinson-Seaton theorem [20]). No such orbital exists to match the *d*-wave and hence δ_2 tends to zero at large photon energies. This turnover from the growing to falling energy dependence of the *s*-phase at the photon energy of about 15 eV makes the energy derivative $d\delta_0/dE$ small while the constantly growing *d*-phase keeps $d\delta_2/dE$ large in this photon energy range.



FIG. 6: (Color online) The time delays as functions of the polar angle ϕ in the equatorial plane for the field strength $\mathcal{E} = 0.05$ a.u. and the pulse duration $T_1/T = 3$. The full TDSE calculations for ionization from the $2p_{+1}$ and $2p_{-1}$ states are plotted with the solid (red) and dashed (green) lines, respectively. The reduced TDSE calculations with the $\mu = 1,3$ terms suppressed in Eq. (4) for the $2p_{+1}$ state and $\mu = \pm 1$ terms suppressed for the $2p_{-1}$ state are drawn with the short dash (blue) and dotted (purple) lines, respectively.

To explain the features of the delays for higher field strength, we have to go beyond the LOPT. As we have already observed in Figure 1, two-photon ionization processes are clearly contributing, but their role is not very important in forming the photoelectron spectrum even for fields as high as $\mathcal{E} = 0.1$ a.u. Situation is rather different for the time delays. We see in Figure 3 that for the short pulse duration and a large field intensity the time delays are modulated but, unlike the case of the weak field, the period of modulation is 2π .

A simple numerical test allows us to clarify the origin of this modulation. We note that absorption and emission of a single photon from the circularly polarized field changes the electron angular momentum projection by ± 1 . So we can gauge importance of different ionization channels just by evaluating contributions of different μ states in the expansion (4). Results of such reduced calculations are shown in Figure 6, where we suppressed the contributions of the terms with $\mu = 1, 3$ in (4) for ionization from the $2p_{\pm 1}$ state, and terms with $\mu = \pm 1$ for ionization from the $2p_{-1}$ state. This excludes the second order processes with participation of two photons and removes the modulation of the time delay completely. For the $2p_{+1}$ initial state, the final state with $\mu = 1$ arises as a result of emission and absorption of a photon, while to get to the $\mu = 3$ final state the atom has to absorb two photons from the circularly polarized field. Similar observation applies to the initial state with $2p_{-1}$. Both these second order processes lead to the electron states which are far in energy from the energy shell. Their contribution, therefore, should decay very rapidly with the increase of the pulse duration. The plots of Figure 3 support this conclusion, 2π -periodic modulation effect is clearly visible for $\mathcal{E} = 0.05$ and 0.1 a.u. for the laser pulse duration of $T_1/T = 3$ and disappears completely for a longer pulse with $T_1/T = 10$.

IV. CONCLUSION

We performed time delay calculations for the photoionization process driven by a circularly polarized laser pulse. We considered initial electron states that are coand counter-rotating with respect to the electric field vector. We found, that similarly to the photoelectron spectra, studied extensively in the literature, the time delays are markedly different for these two orientations, depending sensitively on the field strengths and pulse durations. The time delays for the short pulses exhibit modulation effect with respect to the polar angle in the equatorial plane. The period of this modulation varies with the strength of the electric field. For stronger fields the modulation patterns of the time delays for co- and counter-

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rotating electrons look similar, both patterns can de described as 2π -modulation superimposed on a nearly flat background. This effect can be explained as manifestation of the two-photon processes which do not conserve energy and contribute noticeably only in the case of the ionization by a short laser pulse. Situation is different in the weak-field limit, where modulation effect is present only for the time delay of a counter-rotating electron, and can be described as a π periodic modulation on a constant background. Perturbation theory analysis allowed us to identify the precise absorption mechanisms responsible for this effect.

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