Time delay in valence shell photoionization of noble gas atoms

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We use the non-relativistic random phase approximation with exchange to perform calculations of valence shell photoionization of Ne, Ar, Kr and Xe from their respective thresholds to photon energy of 200 eV. The energy derivative of the complex phase of the photoionization matrix elements is converted to the photoelectron group delay that can be measured in attosecond streaking or two-photon sideband interference experiments. Comparison with reported time delay measurements in Ne and Ar at a few selected photon energies is made. Systematic mapping of time delay across a wide range of photon energies in several atomic targets allows to highlight important aspects of fundamental atomic physics that can be probed by attosecond time delay measurements.

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I. INTRODUCTION

Time delay in atomic photoionization has become an active and rapidly expanding field of research following pioneering experiments on attosecond streaking [1] and two-photon sideband interference [2]. Both techniques use the XUV pump pulse to ionize the target atom and the IR probe to obtain the timing information on the photoemission process. In attosecond streaking, the varying time delay between the the pump and probe pulses is mapped onto the photoelectron kinetic energy. The whole valence shell is projected onto a photoelectron kinetic energy map (the so-called spectrogram) which is then modeled, in the strong field or Coulomb-Volkov approximations, with the photoionization time delay being treated as a fitting parameter. This measurement revealed a relative time delay of 21 ± 5 as between photoemission from the 2p and 2s sub-shells in Ne at 106 eV photon energy. The positive sign of the relative time delay indicates that emission of the photoelectron from the 2p sub-shell is seemingly delayed relative to that from the 2s sub-shell.

In the two-photon interferometric technique, the varying time delay between the pump and probe pulses is mapped onto the two-photon sideband (SB) oscillations. The phase of these oscillations depends on the phase difference of the two neighboring harmonics and the time delay in atomic photoionization process. The atomic time delay can be presented as the sum of time delays in the XUV photon absorption and subsequent IR photon absorption (continuum-continuum or CC transition).

\[ \tau_A = \tau_W + \tau_{CC} \]  

The \[ \tau_W \] term represents the Eisenbud-Wigner-Smith time delay (or Wigner time delay or photoelectron group delay, all these terms will be used interchangeably in the present context) which is defined as the energy derivative of the complex phase of the quantum amplitude of XUV absorption [1, 3]. More details on the Wigner time delay theory can be found in the review article [4]. The \[ \tau_{CC} \] term is modeled using the lowest order perturbation theory and asymptotic forms of the continuum wave functions thus allowing to obtain the former from an experimental measurement [5]. By reconstructing the oscillations of the SB 22 to 26 of the titanium:sapphire laser at 800 nm, Klünder et al. [2] reported the relative time delay between the photoelectron emission from the 3s and 3p sub-shells of Ar in the photon energy range of 34 to 40 eV. Whether the 3p electron was delayed relative to the 3s one or vice versa was found to depend on the photon energy. This measurement was repeated later by Guénot et al. [6] and the sign of the relative time delay was reverted with the 3s photoelectron being delayed relative to the 3p one near the top end of the photon energy scale.

This repeated measurement was prompted by observation that the photon energy of 40 eV fell very close to the Cooper minimum of the 3s shell. Photoionization process in this region is driven very strongly by the many-electron correlation between the 3s and 3p sub-shells [7]. Such a process cannot be theoretically described using an independent electron model like the Hartree-Fock (HF) theory. So the interpretation of the two-photon interferometric measurement [2] based on this theory had to be re-evaluated. A more adequate model that accounts for inter-shell correlation in noble gas atoms is the random phase approximation with exchange (RPAE or, shorter, RPA, both acronyms are used here interchangeably) [8]. However, even after including the RPA corrections, the agreement between theory and experiment did not improve [6].

Theoretical interpretation of the attosecond streaking measurement of Schultze et al [1] is also not straightforward. The group delay difference between the 2p and 2s sub-shells in Ne calculated in the HF approximation is only 6.2 as [3]. With the added RPA correction of 2.2 as, it accounts for less that a half of the experimental value of 21 ± 5 as. More accurate simulations that accounted for both the XUV and IR fields returned somewhat larger values of 10.2 ± 1.3 as [9] and ∼ 12 as [10]. These values

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are still far too small to match the experimental result.

Even though the streaking IR field is relatively weak, its interplay with the long-range Coulomb potential of the ionic core (the so-called Coulomb-laser coupling - CLC) makes an additional contribution to the streaking time delay [11]-[14]. Similar to Eq. (1), the streaking time delay can be written as

$$\tau_s = \tau_W + \tau_{CLC}.$$  \hspace{1cm} (2)

It was suggested in Ref. [11] that $\tau_{CLC}$ should also include the effect of the short-range part of the core potential and hence Eq. (2) should be modified to contain twice the Wigner time delay. This would have resolved the difference between the theoretical [1, 3] and experimental [1] time delays. However, subsequent investigation on the model two-electron system, that mimicked the energy levels of the valence shell of Ne, proved that this conjecture is invalid and that Eq. (2) holds. Therefore the controversy surrounding the experiment [1] still remains unresolved.

In the present paper, we concentrate on the Wigner component $\tau_W$, which enters the atomic time delay (1) and the streaking time delay (2) measured in the attosecond interferometric and streaking experiments, respectively. The corresponding corrections to the Wigner time delay, $\tau_{CC}$ and $\tau_{CLC}$ are more or less universal and can be readily evaluated [10, 14]. For two-electron atomic transitions, like photoionization with excitation and double photoionization, which are strongly driven by electron correlation, the streaking time delay (2) is further modified by the CLC effect on the inter-electron interaction [15]. These two-electron processes, however, are outside the scope of the present study. The target polarization by the streaking IR field can also be safely ignored as it should be minimal for tightly bound closed shell atoms.

In the present work, we perform systematic investigation of the Wigner time delay in a series of noble gas atoms from Ne to Xe across a wide range of photon energies. We demonstrate that in heavier noble gases, beyond Ne, the inter-shell correlation, in the form of direct Coulomb interaction between atomic electrons assigned to different valence sub-shells, has a strong effect on photoionization process in general, and the Wigner time delay, in particular. To account for this direct inter-electron interaction, we employ the RPA method [16]. This method can be viewed as an extension of the HF theory. The latter accounts for the Coulomb inter-electron interaction only indirectly by including some part of it in the self-consistent one-electron potential. On the contrary, the RPA method accounts for a significant part of the direct inter-electron interaction that results in creation of pairwise electron-hole excitations. When more complex excitations of two-electron-two-whole states are important (see e.g. [17]), alternative methods like R-matrix [18] can provide more accurate results.

We validate our computational technique by making an extensive comparison between the calculated and experimental valence shell photoionization cross-sections.

Based on this validation, we make specific predictions for the Wigner time delay and perform further comparison with available experimental time delay data. More generally, we demonstrate that the Wigner time delay contains important phase information that enables attosecond time delay measurements to reveal various fundamental aspects of atomic physics.

The paper is organized as follows. In Sec. II we introduce our computational models for the independent electron descriptions and that with account for the inter-shell correlations. In Sec. III we present our numerical results for outer valence $ns$ and $np$ sub-shells in Ne and Ar and $ns$, $np$, $(n - 1)d$ sub-shells in Kr and Xe. We conclude in Sec. IV by revealing the systematic trends in time delay of noble gases driven by the peculiarities of the elastic scattering phases and many-electron correlations.

**II. THEORETICAL MODEL**

1. Independent electron HF model

We adopt the photoionization formalism as outlined in the monograph [16]. We evaluate the one-photon dipole matrix element $\langle \psi_k^{(-)} | \hat{z} | \phi_i \rangle$ of the transition from a bound state $i$ to an incoming continuous wave with the given photoelectron momentum $k$. The magnitude of the momentum is restricted by the energy conservation $E \equiv k^2/2 = \omega + \varepsilon_i$, where $\omega$ is the photon energy. The atomic units are used throughout the paper with $e = \hbar = \hbar = 1$ and the atomic unit of time is approximately equal to 24.2 as.

We split the radial and angular dependence in the initial state \( \phi_i(r) = Y_{l_i m_i}(\hat{r}) R_{n_i l_i}(r) \) and use the partial wave expansion in the final state

$$\psi_k^{(-)}(r) = \frac{(2\pi)^{3/2}}{k^{1/2}} \sum_{l_m} d e^{-i\delta_l(E)} Y_{l_m}^*(\hat{k}) Y_{l_m}(\hat{r}) R_{E_l}(r), \hspace{1cm} (3)$$

where the radial orbitals are normalized to energy $\langle E_l | E_l' \rangle = \delta(E - E')$ and have the asymptotics at infinity

$$P_{E_l}(r) \bigg|_{r \rightarrow \infty} = \sqrt{\frac{2}{\pi k}} \frac{1}{r} \sin(kr - \frac{l\pi}{2} + \delta_l).$$

We align the quantization axis $z$ with the polarization axis of light and write the dipole operator in the length gauge as $\hat{z} = \sqrt{4\pi/3} r Y_{l0}(\hat{r})$. We perform the spherical integration to arrive to the following expression:

$$\langle \psi_k^{(-)} | \hat{z} | \phi_i \rangle = \frac{(2\pi)^{3/2}}{k^{1/2}} \sum_{l_i m_i} d e^{i\delta_l(E)} Y_{l_m}^*(\hat{k}) \times \left( l \begin{array}{ccc} 1 & l_i \cr m & 0 & m_i \end{array} \right) \langle E_l || r \parallel n_i l_i \rangle \hspace{1cm} (4)$$

$$\tau_s = \tau_W + \tau_{CLC}.$$  \hspace{1cm} (2)
Here the reduced dipole matrix element, stripped of all the angular momentum projections, is defined as
\[
\langle E\ell | r || n_il_i \rangle = \hat{\Pi} \left( \begin{array}{c|c} l & 1 \\ \hline 0 & 0 \end{array} \right) \int r^2 dr R_{El}(r) r R_{n_il_i}(r),
\]
where \( \hat{l} = \sqrt{2l+1} \). The partial photoionization cross section for the transition from an occupied state \( n_il_i \) to the photoelectron continuum state \( El \) is calculated as
\[
\sigma_{n_il_i \rightarrow El}(\omega) = \frac{4}{3} \pi \alpha a_0^2 |\langle E\ell || r || n_il_i \rangle|^2.
\]
Here \( \alpha \) is the fine structure constant and \( a_0 \) is the Bohr radius.

The basis of occupied atomic states \( |n_il_i\rangle \) is defined by the self-consistent HF method and calculated using the computer code [19]. The continuum electron orbitals \( \langle El || \rangle \) are defined within the frozen-core HF approximation and evaluated using the computer code [20]. These states are found in the combined field of the nucleus and the HF potential of the frozen electron core. So the photoelectron scattering phase \( \delta(E) \) derived by this method contains both the long-range Coulomb and the short-range Hartree-Fock components.

We note that the reduced matrix element (5) is real and thus the complex phase of the dipole matrix element (4) is defined by the scattering phases \( \delta_{l\pm1}(E) \). According to the Fano’s propensity rule [21], the dipole transition with the increased momentum \( l = l_1 + 1 \) is usually dominant. In such a situation, the photoemission group delay is approximately given by \( \tau_W = d\hat{l}/dE \).

2. Inter-shell correlation

To include inter-shell correlation effects, we employ the RPA model [16]. In this approximation, the reduced dipole matrix element (5) is replaced by its correlated counterpart \( \langle El || D || n_il_i \rangle \) which accounts for correlation between different valence sub-shells. This correlated matrix element is found as a solution of the system of the integral equations:

\[
\langle E\ell || D || n_il_i \rangle = \langle E\ell || r || n_il_i \rangle + \frac{1}{3} \lim_{\epsilon \to 0^+} \sum_{n'lj} \int dE' \langle n'lj || E\ell || n_il_i \rangle \times \frac{\langle E'lj || D || n'lj \rangle \langle n'lj || E \ell || n_il_i \rangle}{\omega - E' + \epsilon_{n'lj} + i\epsilon},
\]

Here the combined sum/integral sign incorporates both the summation over the discrete excited states \( n'il' \) with the energy \( \epsilon_{n'il'} \) and the integration over the continuum \( dE' \) from the threshold to infinity. The Coulomb matrix contains both the direct and the exchange parts \( V = 2U - W \). That explains the term exchange in the name RPA(E). The direct Coulomb matrix is expressed as

\[
\langle n'l_j El || U || E'l'_njl_i \rangle = i\hat{l}_j^* l_i \left( \begin{array}{c|c} l & 1 \\ \hline 0 & 0 \end{array} \right) \left( \begin{array}{c|c} l' & 1 \\ \hline 0 & 0 \end{array} \right) \times R_{ljl_i,l'_j}(E, E', n_i, n_j),
\]

where \( R(\epsilon) \) is a Slater integral [16]. In the exchange matrix, the electron \( El \) and the hole \( n_jl_j \) states are swapped.

The RPA equations are represented graphically in Fig. 1. Here the straight line with an arrow to the left or right represents electron (continuum) or hole (bound) states, respectively. The wavy line exhibits the Coulomb interaction. The dashed line is used to display a photon of the frequency \( \omega \). The shaded circle is the correlated dipole matrix element whereas the bare matrix element is exhibited by a three-pronged vertex. The Coulomb interaction matrices \( \langle n'l_j El || V || E'l'_njl_i \rangle \) and \( \langle E'l_j El || V || n'il_jnjl_i \rangle \) describe the so-called time-forward and time-reverse correlation processes which are exhibited by the second and third diagrams (from left to right).

In the time-forward process, the photon absorption is followed by the inter-electron interaction in the form of creation of the virtual electron-hole pair in the neighbouring sub-shell. In the time-reverse process, the virtual electron-hole pair is created before the photon absorption takes place. Because the time-forward process is real in a sense that it conserves the energy of the system, while the time-backward process is virtual, the time-forward process makes stronger contribution to the photoionization process. However, for the completeness and gauge invariance of the theory, both processes should be taken into account.

![FIG. 1: Graphical representation of the RPA equations (7). Left: non-correlated dipole matrix element. Center: time-forward process. Right: time-reverse process.](image-url)
get access to the phase information, one has to evaluate the angular asymmetry parameter $\beta$ which contains the phase difference between the two photoionization channels $l = l_i \pm 1$ when $l_i \neq 0$ [16]. The photoelectron group delay, which is the energy derivative of the phase of the complex photoionization amplitude, gives an alternative access to the phase information. It is evaluated as

$$\tau = \frac{d}{dE} \arg f(E) \equiv \text{Im} \left[ f'(E)/f(E) \right].$$

Here the photoionization amplitude $f(E)$ is given the partial wave expansion

$$f(E) \propto \sum_{l=\pm 1} e^{i\delta_l} Y_{lm}(\hat{k}) (-1)^m \left( \frac{l}{m} 1 l_i m_i \right) \times \langle E|D|n_i k_i \rangle \tag{10}$$

The amplitude $f(E)$ is evaluated in the forward direction $k||\hat{z}$, which is usually the case in the attosecond time delay measurements. In this case, $Y_{lm}(\hat{k}) = \hat{l}(4\pi)^{-1/2} \delta_{m0}$ and hence $m_i = 0$ also. It has to be noted that the phase of the amplitude (10) contains the contribution of the HF phases $\delta_l$ in both photoionization channels $l = l_i \pm 1$ as well as the RPA correction due to the imaginary part of the RPA dipole matrix element $\langle E|D|n_i k_i \rangle$. Thus the associated group delay is labeled HF+RPA when the numerical results are presented in the following section.

### III. NUMERICAL RESULTS

#### A. Neon 2s and 2p sub-shells

On the top panel of Fig. 2 we present the partial photoionization cross-sections of valence shell photoionization of Ne. The HF cross-sections are shown by the dashed (blue) lines and the RPA cross-sections are exhibited by the solid (red) line. The recommended experimental data by Bizau and Wuilleumier [23] are displayed with error bars. In the RPA calculation, we substitute the HF bound state energies with the experimental ionization thresholds $\varepsilon_{2p_{3/2}} = 21.56$ eV and $\varepsilon_{2s} = 48.47$ eV [24] which are indicated on the upper boundary of the panel. We see that account for the RPA correlation between the 2s and 2p sub-shells improves the calculated cross-sections and makes them closer to the experimental data.

We note that even though agreement between theory and experiment is improved in the RPA model, there is a visible difference between the calculated and measured cross-sections, especially for the 2s sub-shell. This difference may arise from the fact that not all the many-electron correlations are accounted for by the RPA model which includes pairwise electron-hole virtual excitations. Other processes like admixture of the two-hole-one-electron states to the pure one-hole state in the singly charged ion are not included in the RPA model. This admixture is responsible for the shift in atomic ionization potentials relative to the corresponding HF binding energies as well as appearance of the satellite lines in the photoionization spectra [25]. These effects cannot be accounted for ab initio in the RPA model. Phenomenologically, they are partly compensated by using the experimental ionization potentials instead of the HF energies $\varepsilon_i$ in the RPA equations (7).

![FIG. 2: (Color online) Top: the partial photoionization cross-sections of the 2s and 2p sub-shells of Ne. The HF and RPA calculations are shown by the dashed (blue) and solid (red) lines, respectively. The recommended experimental data by Bizau and Wuilleumier [23] are displayed with error bars. Middle: elastic scattering phases in the field of the Ne$^+$ ion for the 2s $\rightarrow$ Ep and the dominant 2p $\rightarrow$ Ed channels (dotted blue line) and the RPA phases (solid red line). The thin dotted line visualizes the Coulomb phase with $Z = 1$. Bottom: the phase derivatives are converted to the units of the group delay. The vertical bar at the photon energy of 106 eV visualizes the relative time delay between the 2p and 2s sub-shells of 21 as as measured by Schultze et al. [1] (Color online) Top: the partial photoionization cross-sections of the 2s and 2p sub-shells of Ne. The HF and RPA calculations are shown by the dashed (blue) and solid (red) lines, respectively. The recommended experimental data by Bizau and Wuilleumier [23] are displayed with error bars. Middle: elastic scattering phases in the field of the Ne$^+$ ion for the 2s $\rightarrow$ Ep and the dominant 2p $\rightarrow$ Ed channels (dotted blue line) and the RPA phases (solid red line). The thin dotted line visualizes the Coulomb phase with $Z = 1$. Bottom: the phase derivatives are converted to the units of the group delay. The vertical bar at the photon energy of 106 eV visualizes the relative time delay between the 2p and 2s sub-shells of 21 as as measured by Schultze et al. [1] ](image-url)
The Coulomb phase for the photoionization matrix element is given by 
\[ \delta_{\nu}(k \to 0) - \sigma_{\nu}(k \to 0) = \mu_{\nu}(\infty)\pi \]
for a neutral target. The RPA phase at zero energy is estimated by the number of bound target states \( N_{\nu} \) according to the Levinson-Seaton theorem. In the absence of the Coulomb potential, the \( 2s \to E\sigma \) phase tends to zero as there are no occupied sub-shells in the \( \text{Ne}^+ \) ion with \( n = 2 \). With the Coulomb potential taken into account, \( \delta_{2s,+k}\pi(k \to 0) - \sigma_{2s,+k}\pi(k \to 0) = 0.88\pi \), where \( \mu_{2s,1} = 0.88 \) is the quantum defect calculated from the \( np \) orbital energies in the \( \text{Ne}^+ \) ion \( \epsilon_{np} \sim -(n - \mu_{2s,1})^2 \) for \( n > 2 \). As \( \mu_{2s,1} \) tends to zero, the RPA phase stays flat at the value determined by the corresponding quantum defect. We may associate this behavior with the photoionization matrix element. The resulting time delay difference \( \Delta \tau_{\text{RPA-HF}} = 8.4 \) as with the difference between the corresponding CLC corrections \( \Delta \tau_{\text{CLC}} = 3.5 \) as [14]. The resulting time delay difference \( \Delta \tau_{\text{S}} = 11.9 \) as which is very similar to that reported in [10] by only half of the experimental value of \( 21 \pm 5 \) as.

1. **Argon 3s and 3p sub-shells**

An analogous set of data for Ar 3s and 3p sub-shells is shown in Fig. 3. On the top panel we make a comparison of the HF (dashed blue line) and the RPA (solid red line) partial photoionization cross-sections with the experimental data by Möbus et al. [27] for 3s sub-shell and by Samson and Stolte [28] for the sum of 3s and 3p sub-shells. The experimental ionization thresholds \( \epsilon_{3p-3s} = 15.76 \) eV and \( \epsilon_{3s} = 29.24 \) eV [24] are indicated on the upper boundary of the panel. These partial photoionization cross-sections are qualitatively different from those of Ne shown in Fig. 2. Firstly, the 3p cross-section in Ar displays the Cooper minimum whereas the nodeless 2p orbital does not [29]. Second, the inter-shell correlation changes completely the 3s cross-section, which also displays a deep Cooper-like minimum at a slightly smaller photon energy. The RPA calculation reproduces these features in fair agreement with the experiment.
gous case of Ne except that the $3s \rightarrow E_p$ phase would tend to $2\pi$ in the absence of the Coulomb singularity as there are two occupied np-shells in the Ar$^+$ ion. With the Coulomb potential taken into account, $\delta_{3s \rightarrow kp}(k \rightarrow 0) - \delta_{3s \rightarrow kp}(k \rightarrow 0) = 1.73\pi$ where the corresponding value of the quantum defect in Ar$^+$ is $\mu_{t=1} = 1.73$. The RPA phases in Ar are very different from Ne. When the Coulomb potential is taken into account, $\delta$ there are two occupied ps, and $\pi$ in the $3p \rightarrow Ed$ amplitude. This jump is easy to understand. If the amplitude was real and had a node, it would simply change its sign which would amount to adding a phase factor of $\pi$ in the complex number representation. Incidentally, this jump was investigated in an earlier model calculation \cite{30} which established validity of the attosecond streaking technique for the phase measurements.

This jump of $\pi$ has a dramatic effect on the time delay which is shown on the bottom panel of Fig. 3. It drives the time delay in the 3s sub-shell to very large numbers in several hundreds of attoseconds. The situation is less dramatic for the 3p sub-shell. Here the normally weak $3p \rightarrow Es$ transition takes over near the Cooper minimum of the strong $3p \rightarrow Ed$ transition and the resulting time delay does not go below $-100$ as. We note that there is a strong variation of phase near the autoionization resonances in the 3p photoionization which is seen on the top panel of Fig. 3. We do not show this variation in the phase and time delay plots for clarity of presentation. Anyway, these resonances are far too narrow to be detected in time delay measurements at present energy resolution.

One can compare significant time delay near the Cooper minimum with the delay time in Breit-Wigner resonant scattering $\tau_d = 2/\Gamma$ with $\Gamma$ being the resonant width at half maximum of the cross-section \cite{31}. In the case of the Cooper minimum in Ar, which is roughly 0.5 a.u. of energy wide, the time delay is expected to be 4 atomic units of time which equates to about 100 as. Of course, this is a very rough estimate and the actual time delay is not constant but varies across the Cooper minimum. The steepness of this variation can only be estimated from an accurate numerical calculation.

On the upper boundary of the variation, we indicate the photon energies corresponding to the SB 22 to 26 of the titanium:sapphire laser at 80 nm used in the two-photon interferometric experiments \cite{2,6} We see that at this photon energy range, the RPA correction changes completely the sign of the relative 3p/3s time delay. In the HF approximation, the 3p photoemission is delayed more that the 3s ones. The inter-shell correlation changes this ordering completely. With the RPA correction, it is the 3s that is delayed more than the 3p. This is an important, strong and qualitative result which is related to the Cooper minima in the corresponding partial photoionization cross-sections. This result is confirmed by an alternative time-independent calculation by Dahlströmm et al. \cite{10} with a similar account for many-electron correlations as in RPA. As compared to the original calculation presented in \cite{10}, the group delay data shown on the bottom panel of Fig. 3 are corrected for the experimental ionization potentials \cite{32}. Without this correction, the HF ionization potential of the 3s sub-shell $\varepsilon_{3s} = 34.7$ eV makes the SB 22 inaccessible.

![](image)

**FIG. 3:** (Color online) Top: the partial photoionization cross-sections of the 3s and 3p sub-shells of Ar. The HF and RPA calculations are shown by the dashed (blue) and solid (red) lines, respectively. The experimental data for 3s \cite{27} and for 3s+3p \cite{28} are displayed with error bars. Middle: elastic scattering phases in the field of the Ar$^+$ ion for the 3s $\rightarrow E_p$ and the dominant 3p $\rightarrow Ed$ channels (dotted blue line) and the RPA phases (solid red line). Bottom: the phase derivatives are converted to the units of the group delay. The green asterisks display the calculation \cite{10} corrected for experimental ionization thresholds.

A strong modification of the relative time delay between the 3p and 3s sub-shells in Ar is more clearly seen in Table I where we present the time delay difference $\tau_{3s} - \tau_{3p}$ in the HF and RPA approximations and compare it with the experimental data of Guénot et al. \cite{6}. Even a fairly large uncertainty of $\pm 50$ as cannot reconcile the experimental data with neither of the calculations. In the same table, we present results of a multi-configurational Hartree-Fock (MCHF) close-
coupling calculation [33]. In this calculation, the Cooper minimum was displaced to significantly larger photon energies which were not probed experimentally. Hence the time delay difference at the SB 26 was not affected by this minimum as strongly as in the present RPA calculation.

2. Krypton 4p, 4s and 3d sub-shells

Our results for the 4p, 4s and 3d photoionization of Kr are displayed in Fig. 4. On the top panel we make a comparison of the HF (dashed blue line) and the RPA (solid red line) partial photoionization cross-sections with the experimental data of Ehresmann et al. [34] for 4s and of Samson and Stolte [28] for 4p + 3d (error bars). The data from Aksela et al. [35] for 3d are displayed with asterisks. The experimental ionization thresholds $\epsilon_{4p,3/2} = 14.00$ eV, $\epsilon_{4s} = 27.51$ eV [24] and $\epsilon_{3d,5/2} = 93.83$ eV [36] are indicated on the upper boundary of the panel. The 4p and 4s cross-sections in Kr behave similarly to the 3p and 3s cross-sections in Ar (see the top panel of Fig. 3). The 4p $\rightarrow$ Ed cross-section goes through its Cooper minimum which is offset somewhat by the weaker 4p $\rightarrow$ Es channel. So the total 4p cross-section displays a shoulder rather than a true minimum. The 4s cross-section is driven strongly by its inter-shell correlation with 4p to a very deep minimum which is missed completely in the HF approximation. The 3d cross-section from its threshold displays a strong maximum associated with its shape resonance. This resonance is known to be due to electron correlation within a single shell [37] and indeed the 3d photoionization cross-section is well described by the HF approximation.

The HF phases in Kr (middle panel of Fig. 4) behave similarly to the analogous cases of Ne and Ar except that the 4s $\rightarrow$ Es phase would tend to $\pi$ and the 4p $\rightarrow$ Ed phase would tend to $\pi$ in the absence of the Coulomb potential. With this potential, the HF phases are determined by the corresponding quantum defect values $\mu_{l=1} = 2.67$ and $\mu_{l=2} = 1.04$. The RPA phases in Kr are also similar to Ar. Every time the cross-section goes through the Cooper minimum, the corresponding phase makes a jump of $\pi$: upwards in the 4s $\rightarrow$ Es amplitude and downwards in the 4p $\rightarrow$ Ed amplitude. The RPA phase in the 3d $\rightarrow$ Ef transition is rather stationary.

This behavior of the phases translates into the corresponding time delays plotted on the bottom panel of Fig. 4. The RPA time delay in 4p sub-shell is not dramatically different from the HF calculation. Even though the dominant 4p $\rightarrow$ Ed transition displays a Cooper minimum, it is offset by the weak 4p $\rightarrow$ Es transition and is not as prominent in the total 4p cross-section as in the 3p cross-section of Ar. There are some variation of the time delay near the autoionizing resonances close to the 4s threshold which are seen in the RPA calculation but not in HF one. The time delay in the 3d sub-shell is almost entirely due to intra-shell effects and the HF and RPA results

| $\omega$ (eV) | 34.1 | 37.2 | 40.3 |
|---------------|------|------|------|
| $\tau_W^{3s} - \tau_W^{3p}$ (as) | 3    | -36  | -38  |
| HF            | 76   | 53   | 215  |
| RPA           | 45   | 10   | -5   |
| Expt          | 70   | -30  | 50   |

TABLE I: Relative time delay between the photoemission from the 3s and 3p sub-shells $\Delta \tau = \tau_W^{3s} - \tau_W^{3p}$ in Ar at three fixed photon energies corresponding to the SB 22 to 26 in the experiment of Guenot et al. [6]. The experimental uncertainty is $\pm$50 as. The MCHF calculation [33] has typical error bars of $\pm$40 as due to pseudo-resonance structure.
are very close. The situation is very different in the 4s sub-shell where the time delay is strongly affected by the inter-shell correlation with the 4p sub-shell and reaches 300 as in its peak. Similarly to Ar, there is a complete reversal of the relative time delay between the 4p and 4s sub-shells in the RPA calculation in comparison with the HF one.

3. Xenon 5p, 5s and 4d sub-shells

The analogous set of data for the 5p, 5s and 4d sub-shells of Xe is presented in Fig. 5. On the top panel we compare the partial photoionization cross-sections in the HF (dashed blue line) and RPA (solid red line) approximations with the experimental data [38, 39] which are shown with the blue asterisks for 5s and error bars for 5p and 4d. The experimental ionization thresholds \( \epsilon_{5p_{3/2}} = 12.13 \text{ eV} \), \( \epsilon_{2s} = 23.40 \text{ eV} \) [24] and \( \epsilon_{4d_{5/2}} = 67.50 \text{ eV} \) [40] are indicated on the upper boundary of the panel.

Below the 4d ionization threshold, the 5s and 5p cross-sections in Xe behave similarly to the 4s and 4p sub-shells in Kr (top panel of Fig. 4). However, above this threshold, the 4d sub-shell goes through a very steep shape resonance, sometimes even called a “giant resonance”. This resonance is then turns into a Cooper minimum. By strong inter-shell interaction, this behavior is replicated in the 5p and 5s partial photoionization cross-sections which are well reproduced by the RPA calculation. Accordingly, the corresponding RPA phases displays steep \( \pi \) jumps (middle panel) which are reflected in the corresponding time delays (bottom panel). In the case of the 5s sub-shell, the RPA phase jump near the Cooper minimum merges with the Coulomb singularity and produces a very large, nearly 300 as time delay at the photon energies below 30 eV. The 5p sub-shell shows a large and negative time delay due to its Cooper minimum at around 50 eV. Both the 5s and 5p sub-shells display a large and negative time delay near the local cross-section minima around 150 eV induced by the correlation with the 4d sub-shell. The time delay in the 4d sub-shell is driven from the strongly positive due to the Coulomb singularity at low photon energies to a large negative jump near the Cooper minimum at about 180 eV. At larger energies, the cross-sections are rather structureless and there is no significant time delay variations.

A phase jump of \( \pi \), smoothed by the interaction between the two channels, has already been observed both theoretically and experimentally by analyzing the anisotropy parameter in photoionization of Xe 5p sub-shell [41]. This parameter contains the phase shift between the two photoionization channels with \( l = l_i \pm 1 \). In the case of 5p photoionization, these are 5p → Ed and 5p → Es transitions. Their partial photoionization cross-sections and the relative phase shift are presented on the top and bottom panels of Fig. 6. On both panels, we show the present RPA and HF calculations displayed with the solid red and blue dotted lines, respectively. On the bottom panel, we exhibit the RPA (open circles) and HF (filled circles) phase shifts reported by [41].

On the top panel of Fig. 6 we observe a significant shift of the Cooper minimum in the 5p → Ed channel towards the lower photon energies and appearance of the secondary minimum due to the correlation with the 4d sub-shell. In the meantime, the inter-shell correlation does not change the 5p → Es partial photoionization cross-section in such a dramatic way. Accordingly, on the bottom panel of Fig. 6, we see a strong variation of the RPA phase shift with the two successive jumps near the Cooper minima of the 5p → Ed cross-section. In the meantime, the HF calculation returns quite a smooth and monotonous phase shift. Agreement between the two sets of calculations, the present and the one reported by [41], is rather good. A small shift between the present calculation and the reference one is most likely due to scanning.
FIG. 6: (Color online) Top: Partial photoionization cross-sections of Xe in the 5p → Ed and 5p → Es channels in the RPA (solid red line) and HF (dotted blue line) approximations. Bottom: Phase shift between the partial 5p → Ed and 5p → Es waves. The present RPA and HF calculations (solid red and blue dotted lines, respectively) are compared with the RPA and HF calculations reported in Zimmermann et al. \[41\] (open and filled circles, respectively).

and digitizing the analog data of Fig. 3 in Ref. \[41\].

IV. CONCLUSION

In the present work, we perform a systematic study of the photoemission time delay from the valence shells of noble gas atoms in sequence from Ne to Xe. We cover the photon energy range from the ionization threshold to 200 eV. We test the accuracy of our calculation by making comparison with available partial photoionization cross-sections. We derive the complex phase of the photoionization amplitude in the non-relativistic HF and RPA calculations and convert it to the photoelectron group delay by taking the energy derivative.

The time delay results display a very diverse landscape due to an interplay of three major factors. The first two are the logarithmic Coulomb singularity and the Levinson theorem which drive the photoelectron scattering phase in the field of the singly charged ion. The third factor is the phase jump of π near the Cooper minimum which is smoothed by the inter-shell interaction. The two former factors are revealed in the HF calculations whereas the third one is most vividly reflected in the RPA calculations. Experimentally, photoionization measurements near the Cooper minima may be challenging but it is the area where the time delay effects are expected to be largest.

These time delay results are compared with experimental data derived from the attosecond streaking measurement \[1\] and the two-photon interferometric technique \[6\]. This comparison is inconclusive as the difference between the theoretical and experimental results clearly exceeds the reported error bars. We are fairly confident about the accuracy of the present calculation which is tested by comparison of the partial photoionization cross-sections with a large set of independent experimental data and the angular asymmetry parameters as in the case of Xe \[41\]. It is hard to give a numerical estimate on the accuracy of the group delay results. In lighter atoms we expect it to be within 10%. For Xe, it may be more significant as suggested by larger difference between the calculated and experimental cross-sections. Even for this heaviest of the atoms studied in the present work, the relativistic effects are not expected to change considerably the complex phase \[41\] and hence the associated group delay. It is therefore an open question why the time delay results cannot be verified experimentally even after the corresponding CLC or CC corrections are made. Such a verification would be a very welcoming development both for the attosecond time delay measuring techniques and the complete theory of atomic photoionization.

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