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# Tracking autoionizing-wavepacket dynamics at 1-femtosecond temporal scale

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We present time resolved studies and Fourier-Transform-Spectroscopy of inner-shell excited states undergoing Auger decay and doubly excited autoionizing states, utilizing coherent extreme-ultraviolet (XUV) radiation continua. Series of states spanning a range of  $\sim 4\text{eV}$  are excited simultaneously. An XUV probe pulse tracks the oscillatory and decaying evolution of the formed wavepacket. The Fourier transform of the measured trace reproduces the spectrum of the series. The present work paves the way for ultra-broadband XUV spectroscopy and studies of ultra-fast dynamics in all states of matter.

Linear interferometry is equivalent to frequency domain spectroscopy, as postulated in the Wiener Khinchin theorem [1]. Time domain spectroscopic techniques such as optical Ramsey [2] or Fourier Transform Spectroscopy (FTS) are based on this principle. Optical pulses can essentially excite single or few very close lying electronic states. Broadband extreme-ultraviolet (XUV) radiation may significantly increase temporal resolution and excite coherent superpositions of state manifolds spanning a large part of an electronic spectrum. Along this perspective, the present work demonstrates the tracking of 1fs-scale electron dynamics of a spectrally ultra-broad electronic wavepacket.

The temporal evolution of a coherently excited superposition  $|\psi\rangle$  of an atomic ground state  $|g\rangle$ , a number ( $N$ ) of energetically non degenerate decaying eigenstates  $|i\rangle$  and continuum states  $|c\rangle$

$$|\psi(t)\rangle = |g\rangle + \sum_i^N n_i(t) e^{-i(\omega_i t + \varphi_i)} e^{-\Gamma_i t} |i\rangle + \int dE_c n_c(t) |c\rangle \quad (1)$$

(where  $n_i(t)$  is the time dependent probability amplitude of each excited state  $|i\rangle$  with energy  $E_i = \hbar \omega_i$ ,  $\varphi_i$  its the initial phase and  $\Gamma_i$  the decay rate of the state) features a multi-exponential decay superimposed by fast oscillations at frequencies  $\omega_i$  that beat at frequencies  $\omega_i - \omega_j = \Delta E_{ij} / \hbar$  (with  $i, j=1, \dots, N$ ). Such dynamics undergo e.g. a superposition of autoionizing states (AIS). When the spacing  $\Delta E_{ij}$  of the states is of the order of 10 meV the beating period is “long”, of the order of 100 fs, and thus fs pulses are sufficient for probing the dynamics [3]. For  $\Delta E_{ij}$  values of few eV the beating period becomes 1fs or shorter, the probing of which requires attosecond pulses [4]. Coherent broadband continuum XUV radiation, supporting sub-fs pulse durations, like the radiation produced by the recently developed interferometric

polarization gating (IPG) technique [5], at sufficiently high pulse energies [6] is an ideal tool for such studies. The tracking of the dynamics of the excited manifold may be accurately implemented utilizing two time delayed pulse replicas produced by a split mirror autocorrelator [7-10]. The first pulse excites an electronic wavepacket like the one of eq. (1), thus inducing an atomic dipole that oscillates at the frequencies  $\omega_i$ . The second pulse induces a phase shifted dipole, the phase shift being determined by the variable delay  $\Delta\tau$  between the two pulses. Alternating constructive and destructive interference between the two dipoles results in Ramsey fringing appearing in the autoionization signal measured as a function of  $\Delta\tau$ . Even when this high frequency fringing cannot be resolved, its beating at different  $\omega_i - \omega_j$  frequencies can be observable in the interferogram of the ion signal. The contrast in the beating fringes is modulated with the delay, because of the different beating frequencies present. At the same time it undergoes an exponential decay, due to the decay of the interfering amplitudes as a result of autoionization dissipation. Starting from eq. (1) and following the derivation of previous works [11], one finds that the measured ionization signal reads  $S(\Delta\tau) \propto \sum_{ij} e^{-(\Gamma_i + \Gamma_j)\Delta\tau} \cos[(\omega_i - \omega_j)\Delta\tau]$ . Asymmetry Fano-Beutler parameters [12] are further incorporated as phase shifts in the interferometric trace. The Fourier transform of the measured trace gives the frequency spectrum of the manifold.

In the present work, the above method is applied to a manifold of inner-shell and doubly excited AIS of xenon [13-18] using coherent broadband continua spanning from 15 to 25eV the FWHM. More precisely, the absorbed radiation excites a large part of the spectrum of the odd parity Rydberg autoionizing series that includes doubly excited  $5s^25p^4[{}^3P, {}^1D, {}^1S]mlm'l'$  [15] states and 5s inner-shell excited  $5s5p^6({}^2S_{1/2})np[{}^1P_1]$  states (with  $6 < n < 12$ ) undergoing Auger decay. Figure 1a depicts

the excitation scheme. The spectrum of this region is known from photoabsorption studies [14, 17]. The energy differences between these states range from 4 meV to 2.4 eV. As the maximum delay  $\Delta\tau_{\max}$  in this experiment was 220 fs, the frequency resolution was limited to  $4.5 \text{ ps}^{-1}$  so that peaks with separation smaller than  $\sim 50 \text{ meV}$  could not be resolved.

The experiment has been performed utilizing a 10 Hz repetition rate Ti:Sapphire laser system delivering pulses of 80 mJ/pulse energy and  $\tau_L=40 \text{ fs}$  duration at a carrier wavelength of 800 nm. The XUV broadband coherent continuum radiation is generated by an Interferometric Polarization Gating (IPG) device [5, 6]. The experimental set-up which is shown in Fig. 2a, is described in detail elsewhere [8]. The generation occurs in a Xe gas jet and the XUV radiation is reflected by a Si plate, placed at the Brewster's angle of  $75^\circ$  of the fundamental laser frequency. After reflection the XUV beam passes through an aperture of 3 mm diameter and a 150 nm thick Sn filter in order to select the central part of the radiation with central wavelength of  $\sim 60 \text{ nm}$ . The spectral intensity distribution of the XUV radiation was determined by measuring the energy resolved single-photon ionization photoelectron spectra of Ar gas. The electrons produced by the interaction of the Ar gas with the XUV beam were detected by a  $\mu$ -metal shielded time-of-flight (TOF) spectrometer. The spectrum of the radiation used in the experiment is shown in Fig. 2b. This radiation can support isolated pulses with a duration as short as  $\sim 450 \text{ asec}$ . The duration of the pulse has not been measured, as the available intensity was not sufficient for a 2<sup>nd</sup> order intensity volume autocorrelation (IVAC) [8]. The XUV beam was subsequently focused by a wave-front divider (split spherical gold mirror) of 5 cm focal length into a Xe pulsed gas jet. One half of the mirror is mounted on a feedback-controlled piezo-crystal translation stage with minimum displacement in z-

direction of 1.5 nm. The Xenon ions were recorded by the TOF ion mass spectrometer as a function of the delay  $\Delta\tau$  between the two XUV replicas produced by the split mirror. It should be stressed, that the ionization process in this work is not a two-XUV-photon process [19] in contrast to recent 2<sup>nd</sup> order IVAC measurements [8]. It is a single photon ionization and thus a first order process, for which the split mirror wavefront divider is not an appropriate device [8], since in principle it operates only as a non-linear autocorrelator. The reason that a first order process produces an observable beating trace here, is that the measured signal does not originate from the entire interaction volume. The experimental set up (gas jet dimensions and focusing conditions) introduces a confinement of the interaction volume along the propagation axis. Moreover the data of the trace are line outs of the mass spectra and not integrals of the ion mass peak, which introduces a partial confinement in a second dimension (along the TOF axis) as the TOF was not operated as to have a temporal focus. Due to this volume confinement the signal beating is not entirely vanishing through the spatial integration. This has been verified by 1<sup>st</sup> order autocorrelation measurements (no excited states involved) and confirmed through modelling using the model reported in [10]. Measurements have been performed in two types of scans, a coarse scan with a delay step  $\Delta\tau = 1.33$  fs addressing low frequency components (autoionizing state pairs with small energy differences) and a fine scan with a delay step  $\Delta\tau = 0.333$  fs addressing high frequency components (autoionizing state pairs with large energy differences) of the spectrum. In both scans 100 data points were accumulated for each delay step.

Figure 3 shows traces of a coarse (black line in Fig. 3a) and a fine scan (black line in Fig. 3c). The clearly observable interference fringes are the signature of the excited state superposition and reflect the temporal evolution of the formed wavepacket.

Figures 3b and 3d show the frequency spectrum (grey dot-line) resulting from the Fourier Transform (FT) of the raw data for the coarse and fine scans, respectively and the red line yellow filled area shows the spectrum resulted by an Interpolated Discrete Time Fourier Transform (IDTFT) procedure [20] on the raw data. The peak frequencies are the excitation frequency differences of the autoionizing states. In Fig 3c the first part of the spectrum, where peak 2 appears with large height, is not shown in order for the rest of the peaks to become better visible. Due to the stated limited energy resolution defined by the maximum delay available, not all states can be resolved. Each peak of Figure 3b and 3d originates from the decay of more than one state. The states contributing to each peak are summarized in Table 1. The configurations and numbers in parenthesis give the pair of states contributing to each of the recorded frequency components. Configurations are given for the inner-shell excited states, while numbers represent doubly excited states  $mlml'$  as described in the footnote of Table 1. Lifetimes of the states are given at the foot of the table. In order to enhance the visibility of the low frequency beating ( $< 0.05 \text{ fs}^{-1}$ ), the moving average of the raw data in the coarse and fine scan are taken over 7 and 20 points, respectively and are shown in the red line yellow filled areas in fig. 3a,c.

To calculate the fringe contrast we have used the ionization probability of Xe as obtained from the theoretical model of ref. 11. In the calculation, the widths  $\Gamma$  of the states, the Fano-Beutler parameters  $q$ , and the relative ionization cross-sections  $\sigma$  are taken from ref. 14 and 15. The values of  $\sigma$  for the inner-shell excited autoionizing Rydberg states and the values of  $q$  for the states with energy in the region  $22.5 \text{ eV} < E < 22.7 \text{ eV}$  have been extracted from ref. 13. The widths  $\Gamma$  of Rydberg states  $np$  with  $9 < n \leq 12$  have been obtained through an extrapolation of literature  $\Gamma$  data using  $\Gamma(n) \propto 1/(n - \delta)^3$  as fit function. The parameter  $\delta$  is the quantum defect, which in the

present extrapolation is found to be  $\delta = 3.06$ . It has been found that the states with energies higher than 23.2 eV have minor contribution in the general behavior of the calculated trace. The blue-line yellow-filled area in Fig. 4 shows the calculated fringe contrast together with the contrast extracted from the experimental data (grey line). The moving average of the raw data is shown in red line. In this figure, the calculated data have been multiplied by a constant in order to bring the calculated and measured data at approximately the same scale. This does not affect the positions of the maxima and minima. The calculated and experimental data appear to be in reasonable agreement. The dash green line in Fig. 4 is an exponential ( $y \propto \exp(-\Delta\tau/\tau_D)$ ) fitted to the maxima of the fringe contrast. The resulting  $\tau_D = 140 \pm 90$  fs is a “weighted average” life time of the manifolds and is in good agreement with measured lifetime values.

In conclusion, we present time resolved studies and Fourier-Transform-Spectroscopy utilizing extreme-ultraviolet radiation continua. The method allows the tracking of ultrafast electronic wavepacket dynamics exhibiting oscillatory and exponential decay evolution. The method is applied here to a superposition of a coherently excited autoionizing manifold, the decay of which facilitates detection, but can be applied to non self decaying states by absorption of a second photon leading to photoionization [21]. Total spectral widths that can be treated in one run may be extended to few tens of eV or equivalently to a sub-100 attosecond temporal resolution. At the other end energy resolution can be increased by simply increasing the total length of the delay line. The method, demonstrated here for the electronic motion of an atomic system, can be applied in electronic or non-electronic ultrafast dynamics of more complex systems in all states of matter.

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## Figure Captions

**Figure 1.** Single photon ionization scheme of Xe induced by a coherent broadband XUV radiation continuum.

**Figure 2.** a) Experimental apparatus. IPG: Interferometric Polarization Gating device; L: lens; GJ1: Pulsed Xe gas jet; Si: Silicon plate; F: Sn filter; GJ2: Pulsed gas jet; TOF: Time of flight spectrometer; SM: Split mirror. b) Spectrum of the coherent continuum XUV radiation. The black solid and dash lines in b) show the spectral position of the autoionizing states within the XUV radiation spectrum.

**Figure 3.** Ultra-fast autoionization dynamics measurement. a) The trace shown by the grey line is the autoionization signal as a function of the delay between the two XUV pulses, in a coarse scan. A higher-order autocorrelation trace of the fundamental laser field has been used for the calibration of the zero delay value ( $\Delta\tau = 0 \pm 3$  fs) of the trace. b) Frequency spectrum obtained from the Fourier transform of the raw data of the trace shown in (a). The numbering of the peaks refers to the autoionization states in Table 1. c) Traces as in (a) in a fine scan. d) Frequency spectrum obtained from the Fourier transform of the raw data of the trace shown in (c).

**Figure 4.** Comparison of the calculated (blue-line) and measured (grey-line) fringe contrast of the autoionization signal. The grey solid line depicts the fringe contrast of the raw data of the recorded trace in Fig. 3a. The moving average of the raw data is shown as a red line. The green dash line is an exponential decay fit on the raw data.

**Table 1.** AIS contributing to the spectral distribution of figure 3b and d. The numbers in the first column correspond to the numbering in Fig. 3b and d. The second column gives the central frequencies  $\nu$  of the peak in Fig. 3b and d. The third column shows the combination of the pairs of AIS contributing to each peak.

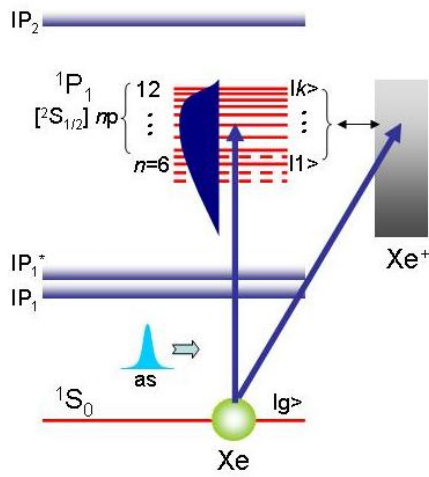


Figure 1

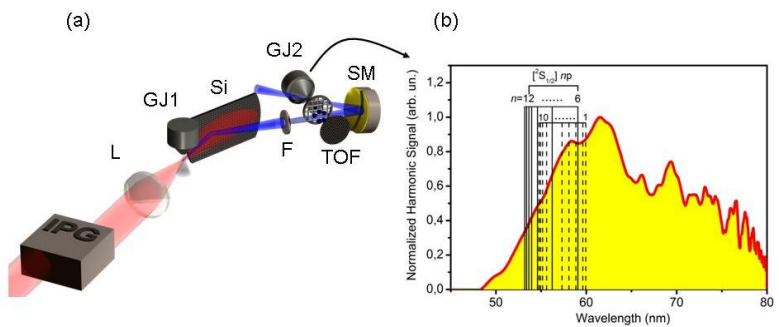
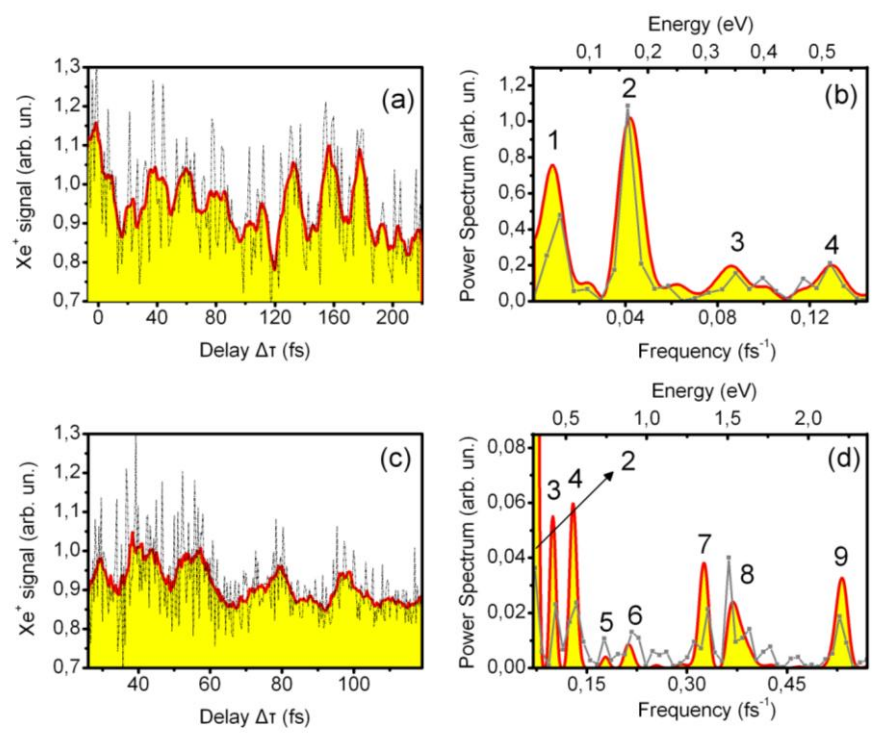
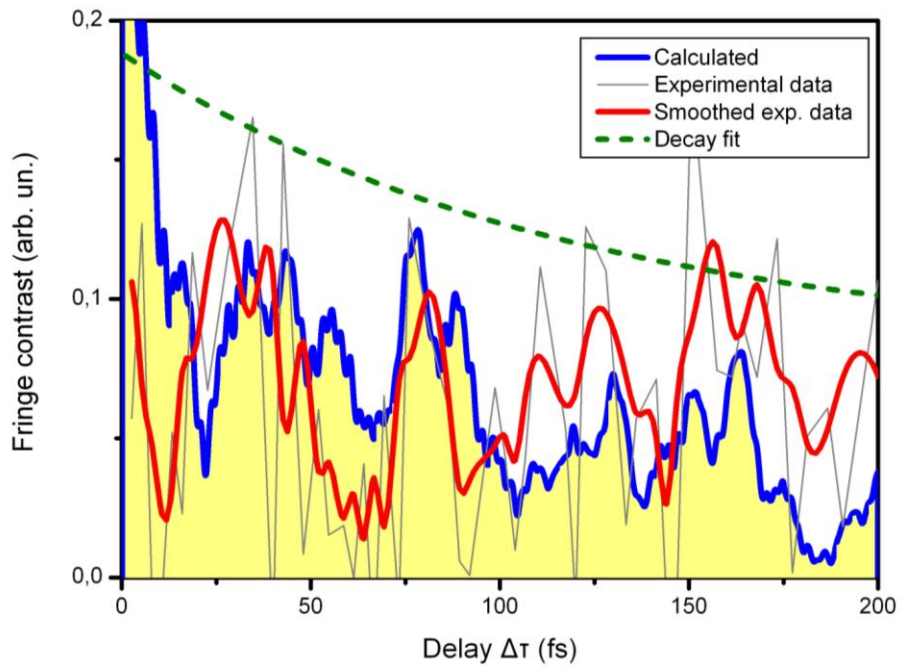


Figure 2



**Figure 3**



**Figure 4**

**Table 1**

Peak Nr.	$\nu$ (fs <sup>-1</sup> ) $\times 10^{-3}$	Autoionization state pairs
1	9±6	([ <sup>2</sup> S <sub>1/2</sub> ]12p, [ <sup>2</sup> S <sub>1/2</sub> ]11p), ([ <sup>2</sup> S <sub>1/2</sub> ]8p, 7), (9, 8), (10, 9)
2	42±10	([ <sup>2</sup> S <sub>1/2</sub> ]11p, [ <sup>2</sup> S <sub>1/2</sub> ]9p), ([ <sup>2</sup> S <sub>1/2</sub> ]6p, 2), ([ <sup>2</sup> S <sub>1/2</sub> ]8p, 8-9), (10,7), (6, 8), (2, 1)
3	93±10	([ <sup>2</sup> S <sub>1/2</sub> ]11p-[ <sup>2</sup> S <sub>1/2</sub> ]10p, 8p), ([ <sup>2</sup> S <sub>1/2</sub> ]7p, 10), ([ <sup>2</sup> S <sub>1/2</sub> ]8p, 6), ([ <sup>2</sup> S <sub>1/2</sub> ]9p, 8-9), (4, 3)
4	128±10	([ <sup>2</sup> S <sub>1/2</sub> ]12p, [ <sup>2</sup> S <sub>1/2</sub> ]8p), ([ <sup>2</sup> S <sub>1/2</sub> ]7p,5), ([ <sup>2</sup> S <sub>1/2</sub> ]10p, 8-9)
5	179±10	([ <sup>2</sup> S <sub>1/2</sub> ]9p, [ <sup>2</sup> S <sub>1/2</sub> ]7p), ([ <sup>2</sup> S <sub>1/2</sub> ]6p, 5), ([ <sup>2</sup> S <sub>1/2</sub> ]10p, 6), ([ <sup>2</sup> S <sub>1/2</sub> ]11p, 7), ([ <sup>2</sup> S <sub>1/2</sub> ]11p, 9-10), ([ <sup>2</sup> S <sub>1/2</sub> ]12p, 7-8), (7, 5), (4, 1)
6	214±15	([ <sup>2</sup> S <sub>1/2</sub> ]7p, [ <sup>2</sup> S <sub>1/2</sub> ]11p-[ <sup>2</sup> S <sub>1/2</sub> ]10p), ([ <sup>2</sup> S <sub>1/2</sub> ]12p, 6), (5, 2), (4, 5-6), (5, 10)
7	325±15	([ <sup>2</sup> S <sub>1/2</sub> ]6p, [ <sup>2</sup> S <sub>1/2</sub> ]7p), ([ <sup>2</sup> S <sub>1/2</sub> ]6p, 6), ([ <sup>2</sup> S <sub>1/2</sub> ]8p, 4), ([ <sup>2</sup> S <sub>1/2</sub> ]10p, 5), (6, 3)
8	371±20	([ <sup>2</sup> S <sub>1/2</sub> ]6p, 7-10), ([ <sup>2</sup> S <sub>1/2</sub> ]7p, 1), ([ <sup>2</sup> S <sub>1/2</sub> ]9p, 4), (2, 6-7), (3, 9-10)
9	532±20	([ <sup>2</sup> S <sub>1/2</sub> ]6p, [ <sup>2</sup> S <sub>1/2</sub> ]10p-1[ <sup>2</sup> S <sub>1/2</sub> ]2p), ([ <sup>2</sup> S <sub>1/2</sub> ]9p, 1-2), ([ <sup>2</sup> S <sub>1/2</sub> ]10p, 2), ([ <sup>2</sup> S <sub>1/2</sub> ]11p, 3), ([ <sup>2</sup> S <sub>1/2</sub> ]12p, 3)

States 1-10 are doubly excited states with energies in eV 20.664, 20.805, 21.03, 21.407, 21.721, 22.333, 22.457, 22.514, 22.56, 22.617, respectively. The lifetimes (in fs) of the 1-10 doubly excited and [<sup>2</sup>S<sub>1/2</sub>]6p-[<sup>2</sup>S<sub>1/2</sub>]12p Rydberg states are 177, 80, 45, 113, 91, 100, 82, 110, 134, 94 and 21, 50, 105, 156, 276, 414, 590, respectively.