Delay measurement of attosecond emission in solids

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Abstract

We characterize an attosecond pulse without using a secondary laser pulse. Instead, we use the carrier-envelope-phase dependence of the extreme ultraviolet (XUV) emission to measure the dipole current from a solid material. To confirm, we apply our approach to data derived by solving the time-dependent Schrödinger equation with a periodic potential. Finally, using MgO, we measure the delay shift at the strong XUV absorption feature and explain it as the quantum interference between the continuum and bound states that participate in the generation process.

Keywords: strong-field physics, attosecond pulse measurement, delay in recollision process

(Some figures may appear in colour only in the online journal)
with its second harmonic [17]. All of these methods have the same purpose and can be qualitatively understood as grating the XUV emission in time to a single attosecond burst.

We propose to characterize the temporal structure of an attosecond pulse by taking the narrow envelope of the few-cycle driving laser pulse itself as a gating function. One can decompose the strong driving laser pulse into two parts—a high frequency carrier and its envelope. The carrier is a sinusoidal function responsible for the creation of XUV emission in every half optical cycle. The envelope limits the number of XUV emissions in time by limiting the number of cycles in the pulse.

Before beginning, it is useful to consider a grating analogy. Each attosecond pulse within a train of pulses is similar to the diffracted light from the grooves of a grating. The gating function corresponds to the width of the grating, defining the number of diffracting grooves. The CEP determines the relative position of the attosecond pulse train with respect to the gating function, equivalent to the determining the relative position of the groove lines on the grating. When there are many lines on the grating, the diffracted spectrum is insensitive to the relative position of the groove. However, it becomes extremely sensitive when the grating has only a few illuminated lines. We will use this analogy as we apply the formalism of grating physics to the attosecond pulse measurement. We will show that the attosecond pulse and the envelope of the driving laser pulse can be measured from the CEP-dependence of the dipole spectrum.

Dipole radiation created by light–matter interaction with a strong laser field is well understood by the following equation from literature when considering the driving laser field as a continuous wave [18, 19]:

$$X(t) = \int_{-\infty}^{\infty} d\tau \left( \frac{\pi}{\xi + i\tau/2} \right)^{1/2} R(t) S(t, \tau) T(t - \tau).$$

(1)

$T(t - \tau)$ is the ionization rate for the electron from the ground state to continuum by tunneling at $t - \tau$ (ionization time). $S(t, \tau)$ is the time evolution function of the ionized electron in the continuum from $t - \tau$ (ionization time) to $t$ (recombination or emission time) while it is accelerated by the laser field during $\tau$ (excursion time). $R(t)$ is the dipole recombination rate from the continuum to the ground state at $t$. The first factor in the $\tau$-integration originates from the 1D momentum integration with a regularized Gaussian function around the saddle point by taking into account the small value of $\xi$. We define the Fourier conjugate function for $R(t)T(t - \tau)$ as in the reference [15]:

$$b(v, \tau) = \int_{-\infty}^{\infty} R(t) T(t - \tau) e^{ivt} dt.$$  

(2)

Then the spectrum of the dipole radiation is obtained by performing a Fourier transform according to the convolution theorem:

$$X(\omega) = \int_{-\infty}^{\infty} dv \int_{-\infty}^{\infty} d\tau \left( \frac{\pi}{\xi + i\tau/2} \right)^{1/2} S(\omega - v, \tau) b(v, \tau).$$  

(3)

Now, we consider a non-uniform envelope $G(t - \tau - \phi(\omega))$ of the driving laser pulse, which limits the number of attosecond emissions in time. Here $\phi$ is the CEP and $\omega$ is the angular frequency of the driving laser field. The CEP-dependent dipole spectrum is written as the following by replacing $T(t - \tau)$ with $T(t - \tau)G(t - \tau - \phi(\omega))$:

$$H(\omega, \phi) = \int_{-\infty}^{\infty} dt e^{-i\omega t} \int_{-\infty}^{\infty} d\tau \left( \frac{\pi}{\xi + i\tau/2} \right)^{1/2} R(t) \times S(t, \tau) T(t - \tau) G(t - \tau - \phi(\omega))$$

$$= \int_{-\infty}^{\infty} d\Omega e^{i\Omega (\omega - \phi(\omega))} G(\Omega) \int_{-\infty}^{\infty} d\tau e^{i\Omega \tau} \left( \frac{\pi}{\xi + i\tau/2} \right)^{1/2}$$

$$\times \int_{-\infty}^{\infty} dv S(\omega - \Omega - v, \tau) b(v, \tau).$$  

(4)

Here $\Omega$ is the energy shift by the gating function $G(t - \tau - \phi(\omega))$, which is the Fourier conjugate variable for $t - \tau - \phi(\omega)$. Since the factor for the wave function diffusion comes from the momentum integration of the regularized Gaussian wave packet, we can combine the first two terms in the $\tau$-integration. Note that the electron kinetic energy $p^2/2$ is very large compared to $\Omega$. It allows us to change the variable of integration $p$ to $p^\prime$:

$$e^{i\Omega \tau} \left( \frac{\pi}{\xi + i\tau/2} \right)^{1/2} = e^{i\Omega \tau} \int_{-\infty}^{\infty} dp e^{-i2\pi \xi p^2} e^{-i2\pi b^2}$$

$$\approx e^{-2\xi(\omega - \Omega)} \int_{-\infty}^{\infty} dp' e^{i2\pi \xi p'^2}$$

$$= \left( \frac{\pi}{\xi + i\tau/2} \right)^{1/2} (\text{due to } \xi \approx 0)$$

(5)

which follows from $p'^2 - 2\Omega \approx p^2$.

Consequently, the CEP-dependent dipole spectrum can be described as a cross-correlation of the dipole radiation and the gating function in frequency, which is equivalent to the principle equation used for the pulse characterization [20–23]:

$$H(\omega, \phi) = \int_{-\infty}^{\infty} d\Omega e^{i\Omega (\omega - \phi(\omega))} G(\Omega) \int_{-\infty}^{\infty} d\tau \left( \frac{\pi}{\xi + i\tau/2} \right)^{1/2}$$

$$\times \int_{-\infty}^{\infty} dv S(\omega - \Omega - v, \tau) b(v, \tau)$$

$$= \int_{-\infty}^{\infty} d\Omega e^{i\Omega (\omega - \phi(\omega))} G(\Omega) X(\omega - \Omega)$$

$$\times \int_{-\infty}^{\infty} dv S(\omega - \Omega - v, \tau) b(v, \tau).$$  

(6)

To test our approach, we artificially construct $H(\omega, \phi)$ by assuming $G(\Omega)$ and $X(\omega)$, representing a gating function with a narrow window of 1.2 fs repeating by a $\pi$-CEP period and a positively chirped attosecond pulse for a bandwidth of 20
eV around a photon energy of 22 eV with a dispersion of 0.06 fs² [24]. Figure 1(a) shows the spectral amplitude and phase (depicted by dashed lines) of the attosecond pulse that we have constructed for the numerical simulation. \( G(\phi) \), a CEP-dependent gating function, is presented in figure 1(b) by taking the Fourier transform of \( G(\Omega) \). Finally, we obtain the CEP-dependent dipole spectra as shown in figure 1(c), which is used as an intensity constraint in the phase retrieval method.

We then verify that an attosecond pulse and a gating function can be measured by applying a phase retrieval algorithm to the CEP-dependent dipole spectra [25]. This is a reverse process to determine two functions, \( G(\Omega) \) and \( X(\omega, \phi) \). Figure 1(d) is the reconstructed \( H(\omega, \phi) \) by the phase retrieval method after running 20,000 iterations. The error in figure 1(e) reaches \( 9 \times 10^{-17} \) after 7000 iterations, monotonically decreasing exponentially. The reconstructed \( X(\omega) \) and \( G(\phi) \) in the figures 1(a) and (b) are nearly identical to the functions that we assumed in the beginning of the simulation.

To demonstrate our method, we now apply the method to experiment. We use the CEP-dependent dipole spectra obtained from MgO. For the experiment, we generate a CEP-stabilized two-cycle laser pulse by delivering a 5 mJ 25 fs 800 nm Ti:sapphire laser pulse to an optical parametric amplifier (OPA). The 1.8 μm idler is passively CEP stabilized with a standard deviation of 260 mrad. This pulse is compressed in an Ar-filled hollow-core fiber where its spectrum is stretched and chirped to cover a wavelength bandwidth from 1.4–2.2 μm. The pulse is then compressed by a 2 mm-thick piece of fused silica to produce a two-cycle pulse [26]. Finally, this two-cycle laser pulse, with an energy of 100 μJ, is focused into a 200 μm-thick MgO crystal, where it reflectively generates XUV radiation from the crystal surface [27].

The XUV spectrum (figure 2(a)) is measured as a function of the CEP by using an x-ray imaging spectrometer, consisting of two toroidal mirrors and an XUV grating. Due to the very short duration of the driving laser pulse, in the near cut-off region of the dipole radiation we observe CEP-modulated XUV spectra around 22 eV with a bandwidth of 3 eV. The modulation is similar to that in the figure 1(c).

We apply our method to the measured XUV spectra of the figure 2(a) for the reconstruction of the attosecond pulse generated from the MgO. The CEP-dependent XUV spectra (figure 2(b)) are recovered after more than 40,000 iterations of the phase retrieval algorithm. The figure shows a π-CEP modulation with a slope similar to that in the figure 2(c). The reconstructed spectral amplitude and group delay of the XUV pulse generated from the MgO crystal are presented in figure 2(c).
by solid and dashed lines, respectively. The XUV spectrum has a two-peak structure due to a peak in the crystal absorption at 22 eV and this results in a positive delay of 5 fs.

Now, to conclude the paper we turn to simulation once again. We perform a one dimensional simulation of XUV generation from a MgO crystal by solving the time-dependent Schrödinger equation with a periodic potential [28–30]. We consider a single active electron moving along the laser field polarization for simplicity. The periodic structure of the crystal is treated by superposing regularly positioned 64 soft-core potentials with a lattice distance of 8\(a_0\) (Bohr radius). The spatial grid is composed of 2048 points with an extent of ±256\(a_0\), corresponding to a momentum space extent up to ±4\(\pi\)\(\hbar/a_0\) (atomic units of momentum) with a resolution of 2\(\pi/512\hbar/a_0\). First we calculate the field-free eigenenergies and their corresponding eigenenergies.

Figure 3(a) shows the energy levels of our crystal model in momentum space for the time-independent Hamiltonian. Energy level splitting is clearly seen in the figure 3(a). We choose the lattice distance and soft-core parameter to match the band gap energies of MgO between the valence band and the 1st and 3rd conduction bands, which are 7.8 eV and 22 eV, respectively. The band gap energy between the 2nd and 3rd conduction bands is 0.18 eV.

We then determine the time dependent evolution of the ground state electron wave function created by the laser field. For this, we apply a monocycle laser pulse with a wavelength of 1.8 μm and peak intensity of 10\(^{13}\) W cm\(^{-2}\) (both similar to our experimental conditions). The time-dependent dipole current is calculated by taking the expectation value of the momentum operator. Since we already have a complete set of electron wave functions from the ground state analysis, we project the time-dependent electron wave function to each band and decompose the dipole current into inter band contributions between the 3rd conduction band and valence band.

Figure 3(b) shows the spectrum and group delay of the calculated dipole current around the XUV photon energy of 22 eV, plotted by a blue solid line and red dashed line respectively. We find a strong absorption peak near 22 eV with 5.2 fs positive delay at the absorption energy that agrees with the corresponding absorption and delay observed experimentally, as depicted in the figure 2(c). The difference of the spectral amplitude and group delay between the experiment and calculation can be attributed to the reduced diffusion of the ionized electron wave packet for the long trajectory radiations due to the use of a single spatial dimension in the simulation. Further, instrumental broadening makes the spectrum widen and the absorption dip at the center be reduced. Due to the set up of our XUV spectrometer, the resolving power is 32, determined by our 500 μm-wide slit and the corresponding energy resolution is 0.69 eV at the photon energy of 22 eV. Therefore, we roughly estimate the linewidth of the absorption to be 0.2 eV, close to the band gap of 0.18 eV, by considering the measured bandwidth of 0.88 eV from the figure 2(c). The results show the attosecond pulse exhibits small group delay dispersion (i.e. the slope of the group delay curve is small).

The absorption and the delay shift of MgO are a quantum resonance phenomena. They arise from the existence of two pathways that lead to the creation of XUV photons (figure 3(c)). An electron initially in the ground state can be ionized by the laser field, indicated by the labelled black arrow. The laser field drives the electron in the conduction band where it approaches the next band. The electron then recombines, radiating an XUV photon with the energy of 22 eV, marked in magenta. This is the regular high harmonic generation process in atoms [1]. This direct pathway for the XUV photon generation is described in the equation (7) by a delta-function in the \(\tau\)-integration:

\[
X_{\text{handdrag}}(t) = \langle g | p | k - A(t) \rangle \int_0^t d\tau [i\hbar \delta(\tau) \\
+ \langle k - A(t) | \Gamma | k - A(t - \tau) \rangle] \\
\times \int_0^{t-\tau} d\tau' \langle k - A(t - \tau) | S | k - A(t - \tau - \tau') \rangle \\
\times \langle k - A(t - \tau - \tau') | T (t - \tau - \tau') | g \rangle. \tag{7}
\]

Figure 2. (a) CEP-dependent XUV spectra measured in experiment and (b) the reconstructed result by the phase retrieval method. (c) XUV spectrum and group delay of the attosecond pulse generated from the MgO crystal achieved by the reconstruction.
Figure 3. (a) Energy diagram of the 1D periodic potential model for simulating the MgO crystal. (b) XUV spectrum and group delay of the attosecond pulse calculated by the 1D TDSE periodic potential model. (c) Step-wise explanation for the resonance process in the MgO crystal at 22 eV. (d) Time-frequency distribution of the calculated attosecond pulse showing time-delayed tail at 22 eV after 2 fs.

Here $|g\rangle$ and $|k\rangle$ are the eigen functions in momentum space for the hole and electron, respectively. $T$ is the time-dependent operator representing tunnel ionization and $S$ is the time-evolution operator from $t - \tau - \tau'$ to $t - \tau$ with the vector potential for field acceleration. The last term in the time integration represents the recombination of the electron in the conduction band to the ground state creating the time-dependent dipole current. Alternatively, the free electron in the continuum state can make a transition to the next highest band by tunneling through the band gap for the energy of 0.18 eV and then return to the ground state. This is expressed in the equation (7) as the second term of the bracket in the $\tau$-integration, usually defined as an exponential decay with the decay time of $1/\gamma$ (here the band gap $\gamma$ is 0.18 eV). As a result, the time-relaxed tail of the dipole current is clearly seen in figure 3(d) with its time-frequency distribution at 22 eV after 2 fs by applying short-time Fourier transform. Due to the small contribution of long trajectory electron wave packets, negatively chirped XUV radiation above 22 eV is weak and the radiation due to short-trajectory is dominant under the decay at 22 eV.

In conclusion, we demonstrate that attosecond pulse measurement is possible using the CEP-dependence of XUV emission when using a few cycle driver. We have derived the principle equations for our method using the strong field approximation. Then, we measure the XUV emission from MgO around its band gap of 22 eV and observe a group delay shift in a region of strong absorption and retrieve the phase of the attosecond pulse using this method.

It is well known that the shaping and engineering of attosecond pulses is possible using material dispersion in solid crystals. Compression of a positively chirped attosecond pulse necessitates the pulse travel through a medium with negative dispersion around the spectral region of interest [31]. Similarly, solid materials with periodic structure also have negative dispersion around the band gap energy due to the strong absorption by the forbidden zone as shown in the figures 2(c) and 3(b). By choosing or designing a solid state medium with tailored absorption properties, this negative dispersion can be used to compensate the intrinsic positive chirp of attosecond pulses in a given spectral range. Thus, attosecond pulses can be optimized while they are being generated from solids by combining material science and semiconductor technology and tailoring solid media for attosecond pulse generation [32, 33].

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