Time-frequency mapping of two-colour photoemission driven by harmonic radiation

Bruno Moio\textsuperscript{1,2,∗}, Gian Luca Dolso\textsuperscript{1,○}, Giacomo Inzani\textsuperscript{1,○}, Nicola Di Palo\textsuperscript{1,2,○}, Rocío Borrego-Varillas\textsuperscript{2,○}, Mauro Nisoli\textsuperscript{1,2,○} and Matteo Lucchini\textsuperscript{1,2,○}

\textsuperscript{1} Department of Physics, Politecnico di Milano, 20133 Milano, Italy
\textsuperscript{2} Institute for Photonics and Nanotechnologies, IFN-CNR, 20133 Milano, Italy

E-mail: bruno.moio@polimi.it

Received 22 June 2021, revised 30 July 2021
Accepted for publication 11 August 2021
Published 1 September 2021

Abstract
The use of few-femtosecond, extreme ultraviolet (XUV) pulses, produced by high-order harmonic generation, in combination with few-femtosecond infrared (IR) pulses in pump–probe experiments has great potential to disclose ultrafast dynamics in molecules, nanostructures and solids. A crucial prerequisite is a reliable characterization of the temporal properties of the XUV and IR pulses. Several techniques have been developed. The majority of them applies phase reconstruction algorithms to a photoelectron spectrogram obtained by ionizing an atomic target in a pump–probe fashion. If the ionizing radiation is a single harmonic, all the information is encoded in a two-color two-photon signal called sideband (SB). In this work, we present a simplified model to interpret the time-frequency mapping of the SB signal and we show that the temporal dispersion of the pulses directly maps onto the shape of its spectrogram. Finally, we derive an analytical solution, which allows us to propose a novel procedure to estimate the second-order dispersion of the XUV and IR pulses in real time and with no need for iterative algorithms.

Keywords: high-order harmonic radiation, extreme-ultraviolet, time-frequency distribution, two-colour photoemission

(Some figures may appear in colour only in the online journal)

1. Introduction
In recent years, ultrashort extreme-ultraviolet (XUV) pulses, produced by high-order harmonic generation (HHG) in gases, in combination with femtosecond infrared (IR) pulses in pump–probe experiments, have unlocked the possibility to study ultrafast dynamics in atoms, molecules and solids with unprecedented temporal resolution, down to the attosecond domain\textsuperscript{[1–5]}. The use of HHG is particularly interesting since it offers the possibility to generate ultrashort pulses, tunable in a broad spectral region from ultraviolet (UV) to soft x-rays. Important physical processes unfolding on a few-femtosecond timescale (e.g. conical intersections in molecules, dynamics of core and deep valence levels, etc) can be investigated by using XUV pulses, produced by filtering single harmonics (SHs) in a broad harmonic spectrum. This can be obtained, for example, by using a time-delay compensated monochromator \textsuperscript{[6, 7]}. Recently, experiments based on a two-colour pump–probe scheme with SH pulses where used to investigate the dissociate ionization in N\textsubscript{2} \textsuperscript{[8]} and the coupled electronic-nuclear dynamics in simple \textsuperscript{[9]} and complex molecules \textsuperscript{[10, 11]}, where the high time resolution, combined with energy tunability, offers new scenarios \textsuperscript{[12]}. In all these experiments a precise assessment of the properties under investigation stems from a proper and reliable characterization of the few-fs pulses. While this task may appear less demanding than the

∗ Author to whom any correspondence should be addressed.
characterization of attosecond pulses, its fulfilment in the XUV regime is not trivial [13]. The temporal properties of both the XUV and IR pulses can be extracted from a SH photoelectron experiment using phase retrieval algorithms [14]. However, assessing the pulses properties directly from the photoelectron spectra, without inversion procedures, would provide a method to intuitively interpret the experimental traces and would be more efficient in terms of computational time.

Here we analyze the time-frequency properties of a two-color, two-photon photoelectron signal, called sideband (SB), obtained by ionizing a target atom with an XUV SH radiation, in the presence of a suitably-delayed IR pulse [15]. Unlike previous works, we study the effect of the pulse spectral dispersion beyond the second order, highlighting how the group delay is directly imprinted in the SB time-frequency distribution. Starting from a model based on the strong field approximation (SFA), which neglects the influence of the atomic potential on the photoemitted electron, we will introduce the central momentum approximation (CMA) and the slowly-varying envelope approximation (SVEA), which will allow us to obtain a simplified expression of the photoemission probability, written as a Fourier transform. Finally, by applying a linear expansion we will obtain an analytical expression for the first SB signal which will allow us to show that it can be interpreted as a time-frequency distribution of the pulses involved in the photoionization experiment. This confirms that the SBs carry the temporal properties of the pulses in their shape [13], as for the case of a Gabor–Wigner transform for signal [16] and pulse analysis [17]. Furthermore, in the particular case of second-order dispersion, we will derive an analytical expression for the SB signal, providing a direct connection between the temporal properties of the radiation and the geometrical properties of the SBs, thus enabling a direct estimate of the temporal characteristics of the pulses.

This work is organized as follows: in section 2 we discuss the physical interpretation of a single-harmonic spectrogram (SHS), in section 3 we introduce a simplified model and analyze the properties of the SB signal. Finally, in section 4 we draw the conclusions.

2. Single-harmonic spectrogram (SHS)

If the XUV photon energy is sufficiently high, the effect of the atomic potential after ionization can be neglected and the photoionization of a rare gas by XUV radiation in the presence of a delayed IR pulse, can be described within the SFA by the following expression [18] (atomic units are used, unless otherwise specified):

\[
S(p, \tau) = \int_{-\infty}^{+\infty} dt X_{\alpha}(t + \tau) e^{i(p \cdot \vec{r})} e^{-\frac{(p \cdot \vec{r})^2}{2}}.
\]  

(1)

where \(X_{\alpha}(t)\) is the photoelectron wavepacket which, far from resonances and strong modulations of the species dipole moment [19, 20], equals the XUV temporal profile, \(E_{\text{XUV}}(t)\). \(I_p\) is the gas ionization potential, while \(p\) is the final electron momentum and \(\tau\) is the delay between the XUV and IR pulses.

In this simplified picture, the XUV pulse produces a photoelectron burst while the IR pulse, described by its vector potential, \(A_{\text{IR}}\), acts as an ultrafast phase modulator through the term:

\[
\phi(p, t) = -\int_{t}^{\infty} dt' (p \cdot A_{\text{IR}}(t') - A_{\text{IR}}(t')/2).
\]

(2)

We note that the model of (1) can fail in reproducing the photoelectron spectra if the XUV energy is comparable with \(I_p\). In that case, more precise calculations based on the time-dependent Schrödinger equation should be performed. Nevertheless, this regime goes beyond the scope of the present work as we are interested in the case where the specific target has no particular influence on the photoelectron distribution. This can always be achieved by employing XUV photons of enough energy. The collection of photoelectron spectra described by \(S(p, \tau)\), is called spectrogram and represents the photoemission probability as a function of \(p\) and \(\tau\). For big values of \(\tau\) the spectrogram coincides with the Fourier transform of the attosecond radiation (rescaled by \(I_p\)). The evolution of \(S\) at small delays depends, instead, on the temporal properties of \(E_{\text{XUV}}\) and \(A_{\text{IR}}\). In the case of an isolated attosecond pulse, the photoelectron center of mass follows the IR vector potential, and the spectrogram is called attosecond streaking trace [21, 22]. If the XUV radiation is composed by an attosecond pulse train obtained through the process of HHG [1], its spectrum is characterized by discrete peaks at the odd multiples of the generating IR driving frequency. At large delays the photoelectron spectra will thus show discrete peaks spaced by twice the IR photon energy and associated with the direct ionization by the odd harmonic photons. At small delays, the interaction with the IR field creates additional photoelectron peaks in between the harmonic signal, called SBs. In a photon picture, they can be explained in terms of the absorption of an harmonic photon followed by the additional absorption/emission of an IR photon [23, 24]. Considering a classical IR field as in (1), they can also be understood as the result of the interference among the streaking traces originated by each pulse composing the train [25, 26]. In this work we will concentrate on the spectrogram originated by a SH, which corresponds to a few-fs XUV pulse in time domain. Figure 1 displays an example of SHS calculated for transform-limited (TL) pulses (no spectral dispersion). The IR intensity is \(5 \times 10^{11}\) W cm\(^{-2}\) and IR and XUV bandwidths are 0.3 PHz and 0.4 PHz respectively, corresponding to a full-width-half-maximum in time of about 7.8 fs and 5.9 fs. The XUV pulse is centered at 35.65 eV (i.e. the 23rd harmonic of 800 nm radiation) and the noble gas is neon \((I_p = 21.6\) eV). At large delays \(S(p, \tau)\) presents a single peak corresponding to direct ionization by the harmonic, hereafter called main band (MB). Near the zero delay (i.e. the temporal overlap of the XUV and IR pulses), SBs appear above and below the MB. Previous works [13, 14] proved such SHS to be sensitive to the group delay dispersion of both XUV and IR pulses, demonstrating the applicability of phase reconstruction algorithms like the extended ptychographic iterative engine [14, 27, 28], to obtain their complete temporal characterization. In the following section we will concentrate on SHS and
show how high order dispersion terms are mapped in the delay-energy distribution of the first SB (labelled with SB$^+$ and SB$^-$ in figure 1).

3. First SB delay-energy distribution

At a given pulse duration, the number of SBs appearing in the SHS depends mainly on the chosen IR intensity. The first SB is formed by a two-color two-photon (XUV and IR) transition. As a result, its delay-energy distribution carries information on both the XUV and IR light temporal characteristics. Figures 2(a)–(c) show the effect of a finite IR second, third and fourth order dispersion (GDD, TOD and FOD, respectively) on the first upper SB of the spectrogram of figure 1. Figures 2(d) and (e) show the same analysis, but considering a transform limited IR pulse and a finite dispersion of the XUV pulse. It is worth noticing that the results we will discuss in the following do not depend on the amplitude and sign of the chosen dispersion parameters. In figure 2 we considered XUV and IR spectral dispersions of opposite signs in order to obtain similar SB$^+$ distributions and favour the direct comparison among the figure panels.

To assure complete control over the spectral phase of the pulses without changing their bandwidth, in these simulations both the IR and the XUV are defined in the frequency domain as follows:

$$
\hat{S} = S_0 \exp \left[ -\left( \frac{\omega - \omega_0}{\Delta \omega} \right)^2 + \sum_{n=2}^{\infty} \frac{D_n}{n!} (\omega - \omega_0)^n \right],
$$

where $S_0$ is the spectral amplitude, $\omega_0$ is the central frequency, $\Delta \omega$ is the spectral width and $D_n$ are the $n$th order dispersion terms. The temporal behavior of both pulses is then obtained by inverse Fourier transform. In general, we find the SB to closely track the group delay of either the IR or XUV pulses, marked with a black continuous curve. The effect of a second order dispersion has already been reported by Lucchini et al [28]: as a result of the linear distribution of the pulse frequencies along their temporal profiles, a finite GDD produces a linearly tilted SB in the delay-energy space. The effect of a finite TOD and FOD is instead reported here for the first time. We found the SB to be profoundly distorted by a finite TOD (figures 2(b) and (e)) or FOD (figures 2(c) and (f)), changing its structure in a characteristic manner. In particular, in the case of a pulse with a TOD, the higher and lower frequencies are shifted on the same side of the pulse envelope, with a quadratic distribution. This produces a c-shaped SB that follows a parabolic energy spreading, as expected from the associated group delay (black curve in the panels). With a FOD, instead, the SB exhibits an s-shaped distribution, tracking a cubic function. The good agreement between the SB delay-energy distribution and the group delay of the pulses can be easily explained with an intuitive picture. As the photoemission involves simultaneous interaction with the two fields, by changing the relative delay between them, we change the portion of pulses which actually overlap in time. Therefore, the time at which different frequencies appear in the pulse, namely the group delay, is mapped onto the final photoelectron distribution. In this regards, one of the pulses can be thought to behave as a gate for the other, as in the case of an ordinary time-frequency distribution [16].

3.1. Perturbative approach

To better understand the intuitive description of the first SB formation we discussed in the previous section, we can develop equations (1) and (2) to obtain a simplified model to describe the photoionization of a rare gas by a SH in the presence of an IR pulse.

Starting from (1), if we are far enough from atomic resonances and the rare gas dipole moment is sufficiently regular, we can apply the wave packet approximation [19], which states that the photoelectron wavepacket $X_e$ is very well approximated by the XUV electric field $E_{XUV}$ if the ionization cross section of the gas is constant. Given its relatively limited bandwidth, this approximation is justified for a SH. In (2), we can apply the CMA, replacing the final electron momentum $p$ by the average momentum of the wavepacket $p_c$. Moreover, we can assume the latter to be oriented along the IR field polarization, such that $p_c \times A_{IR} \approx p_c A_{IR}$. These considerations are common and widely used when employing the frequency-resolved optical gating for complete reconstruction of attosecond bursts (FROG-CRAB) approach [22] to characterize attosecond pulses. At this point, we perform the perturbative approximation, assuming that the IR vector potential has a low intensity ($\lesssim 10^{11}$ W cm$^{-2}$). As a result, (2) further simplifies and becomes:

$$
\phi(t) = -\int_t^{\infty} dt' p_c A_{IR}(t').
$$

If we express the IR vector potential in terms of its envelope and carrier,

$$
A_{IR}(t) = A(t) \cos(\omega_d t + \phi_0),
$$

we can apply the slowly varying envelope approximation (SVEA) while solving the integral of (4) and write the phase
\( \phi(t) \) as:
\[
\phi(t) \approx \frac{p_c}{\omega_0} E_{\text{IR}}(t),
\]
where \( E_{\text{IR}}(t) = -\frac{d}{dt} [A_{\text{IR}}(t)] \) is the IR electric field. In light of these approximations, we can reformulate (1) into the following:
\[
S(\omega, \tau) \approx \left| \int_{-\infty}^{+\infty} dt E_{\text{XUV}}(t + \tau) e^{i \frac{\omega}{\omega_0} E_{\text{IR}}(t)} e^{i \omega t} \right|^2, \tag{7}
\]
where we set \( \omega = p^2/2 + I_p \) to write \( S \) in the form of a Fourier transform.

In case of relatively long pulses (\( \gtrsim 3-4 \) optical cycles), (7) reproduces the photoelectron spectrogram with high degree of fidelity. To test its accuracy, we ran different simulations with an increasing level of approximation, as reported in figure 3, where we display the simulated first-upper SB. In figure 3(a) we show the simulations done with equation (1), i.e. no approximation applied, whereas figure 3(b) shows the effect of the CMA. Figure 3(c) adds the effect of the perturbative approximation. Finally, figure 3(d) shows the effect of all the approximations, up to the SVEA. The parameters of the simulations are the same as in figure 1, with in addition a third order dispersion of 500 fs\(^3\) on the IR. Such a case corresponds to an intense IR radiation with large dispersion and short TL. We chose the simulation parameters to stress the approximations and test their validity in a bad-case scenario. Despite the approximations of (7) seem very rough, the model still produces a very accurate trace even in this extreme case. In fact, the CMA does not have a large effect on the SB shape (only reduces its amplitude), as well as the perturbative approximation does not affect the structure and the features of the 2D signal. The SVEA is responsible for the main distortion, resulting in an up-down symmetrization of the SB. However, the distortion has a minor impact on the SB shape and how the temporal information is impressed on it. This proves that (7)
a proper tool for the description of the SB formation in a photoelectron experiment, even in cases where the approximations are at their limits of applicability.

To provide a further prove of this, we can expand the exponential in (7) in its Taylor series, that becomes:
\[ i \frac{2 \pi}{\omega_0} E_{\text{IR}}(t) = 1 + i \frac{P_c}{\omega_0} E_{\text{IR}}(t) + \frac{1}{2} \left( \frac{P_c}{\omega_0^2} E_{\text{IR}}(t) \right)^2 + \cdots \] (8)

If we concentrate solely on two-photon processes, we can neglect the higher-order terms and consider the first one only. The spectrogram is then represented by:
\[ S(\omega, \tau) \approx \frac{P_c^2}{\omega_0^2} \int_{-\infty}^{+\infty} dt E_{\text{XUV}}(t + \tau) E_{\text{IR}}(t) e^{i \omega t} \, \text{d}t. \] (9)

From a physical point of view, this expression describes the first SB formation as sum-frequency event, between an IR photon and an XUV photon [13]. If the photon energy is not evenly distributed in time, the final SB energy changes following the group delay of the two pulses as discussed above. This effect can also be explained from a mathematical point of view because (9) can be interpreted as a time-frequency distribution, such as the Wigner transform [17]. Equation (9) allows us to explain why the SVEA removes the up-down asymmetry in the SB signal (see figure 3). Without the SVEA, equation (9) should be rewritten as:
\[ S(\omega, \tau) \approx \frac{P_c^2}{\omega_0^2} \left| \int_{-\infty}^{+\infty} dt E_{\text{XUV}}(t + \tau) E_{\text{IR}}(t) e^{i \omega t} \right|^2. \] (10)

At a fixed delay, it corresponds to the modulus square of the convolution between the Fourier transform of the XUV pulse and the Fourier transform of the IR electric field divided by $\omega^2$. The division by $\omega^2$ (absent in (9)) is responsible for the observed up-down asymmetry of the SB signal, which disappears with the SVEA. We note that the symmetrisation caused by the SVEA does not affect the results discussed in the present work, unless the IR pulse duration is reduced to a few cycles.

### 3.2. Geometrical interpretation of the first SB

As shown in figure 2, a non-flat spectral phase of the XUV and IR pulses produces a characteristic distortion of the SB signal which varies with the dispersion order. For example, a quadratic phase dispersion (GDD) produces a tilted SB in the SHS. More precisely, chirped XUV is expected to produce SBs with a parallel delay-frequency tilt while a non-zero GDD of the IR field makes the SBs to assume a funnel shape (opposite sign of the induced tilt) [13]. The simplified description of (9) allows for an easy explanation of these effects.

For simplicity, let us consider two Gaussian pulses of the following complex form:
\[ E_{\text{XUV}}(t) = \frac{E_x}{2} \exp \left[ -\frac{t^2}{\gamma_x^2} + i \left( \omega_x t + \eta_x \right)^2 \right] + \text{cc}, \] (11)
\[ E_{\text{IR}}(t) = \frac{E_0}{2} \exp \left[ -\frac{t^2}{\gamma_0^2} + i \left( \omega_0 t + \eta_0 \right)^2 \right] + \text{cc}, \] (11)

where $E_j$ and $\omega_j$ ($j = x, 0$) represent the fields amplitude and frequency. The parameter $\gamma_j$ is related to the TL width of the Gaussian envelope, $\sigma_j$, by the following equation:
\[ \gamma_j = \sigma_j \sqrt{1 + \frac{2 \beta_j^2}{\sigma_j^2}}, \] (12)

where $\beta_j$ represents the GDD, defined as the second derivative of the pulse spectral phase $\beta_j = \frac{d^2 \phi(\omega)}{d \omega^2} \big|_{\omega_j}$ . With this definition the actual Gaussian envelope full-width-half-maximum duration is given by FWHM = $\gamma_j \sqrt{2 \ln 2}$. The parameter $\eta_j$ in (11), called chirp rate, expresses the quadratic dependence of the instantaneous field frequency over time. For Gaussian pulses it is possible to show that this parameter is linked to $\beta_j$ and $\sigma_j$ by:
\[ \eta_j = \frac{2 \beta_j}{\sigma_j^2 + 4 \beta_j^2}. \] (13)

If we substitute the pulses in (11) into (9) it is easy to show that the first SB signal originates from the Fourier transform of the term:
\[ \exp \left\{ -\left( \frac{1}{\gamma_x^2} + \frac{1}{\gamma_0^2} \right) r^2 + i \left( \omega_x \pm \omega_0 \right) t + \left( \eta_x \pm \eta_0 \right) r^2 \right\}. \] (14)

While the plus sign corresponds to the absorption of an IR photon associated to the upper SB formation, the minus sign describes the emission of one IR photon which produces the lower SB (respectively SB$^+$ and SB$^-$ in figure 1). From (14) it is clear that a finite XUV chirp ($\eta_0 \neq 0$) has the same effect over the photoelectron central frequency for SB$^+$ and SB$^-$, thus inducing a parallel tilt, while the change of sign is responsible for the opposite funnel-like tilt induced by a non-zero IR chirp ($\eta_0 \neq 0$).

For the Gaussian pulses described in (9) and (11) has an analytical solution which allows to test our model from a quantitative point of view. For the sake of clarity, let us concentrate on the first upper SB only, SB$^+$, and redefine the frequency axis in order to be centered at nominal SB energy, by introducing $\omega' = \omega - (\omega_x + \omega_0)$. Following equation (9), the SB$^+$ signal can be approximated by:
\[ \text{SB}^+ (\omega', \tau) \approx S_0 \exp \left\{ -\frac{2(1-\Theta)}{\gamma_0^2} \tau^2 + 2 \Gamma^2 \Theta \omega_\tau - \frac{\gamma_0^2 \Theta}{2} \omega_\tau^2 \right\}. \] (15)

where we have introduced the quantities:
\[ \Gamma^{-2} = \gamma_x^{-2} + \gamma_0^{-2} \]
\[ H = \eta_x + \eta_0 \]
\[ \omega_\tau = \omega' - 2 \eta_0 \tau \]
\[ \Theta = \frac{\Gamma^2}{\gamma_0^2 (1 + 4 \Gamma^2 H^2)} \]
\[ S_0 = \frac{\pi P_c^2}{\omega_0 \gamma_0 \Gamma \sqrt{\Theta} \left( \frac{E_x E_0}{4} \right)^2}. \] (16)
The SB shape can be analysed by extracting the contours of the surface represented by equation (15). They can be expressed by the following formula, as a function of the pulse parameters in (11),

\[ A\tau^2 + B\tau\omega' + C\omega'^2 = \text{const}, \quad (17) \]

where the canonical form parameters are defined as follows:

\[ A = \gamma_2^2\eta_k^2 + \frac{\eta_k^2}{\gamma_0^2} + \frac{1}{\eta_k^2} + \frac{1}{\gamma_0^2} \]
\[ B = \gamma_2^2\eta_k - \gamma_0^2\eta_0 \]
\[ C = \frac{\gamma_2^2 + \gamma_0^2}{4}. \quad (18) \]

This implicit equation, (17), defines a bundle of ellipses in the \((\omega', \tau)\) plane. However, the formal analogy is only partial, since the above mentioned space is not homogeneous, being a frequency-time space. To overcome this apparent limitation, we can redefine the axes as \(x_n = \tau/\sqrt{\gamma_2^2 + \gamma_0^2}\) and \(\omega_n = \omega/\sqrt{\gamma_2^2 + \gamma_0^2}\), such that the new \((\omega_n, \tau_n)\) space is dimensionless. The normalization parameter \(\sqrt{\gamma_2^2 + \gamma_0^2}\) is related to the duration of the energy-integral of the SB, as we will show in section 3.3. In this new space, the contours in (17) still represent ellipses, with the following new canonical parameters:

\[ A_n = A \left( \gamma_2^2 + \gamma_0^2 \right) \]
\[ B_n = B \]
\[ C_n = 1/4. \quad (19) \]

In figure 4 we show a comparison between the first SB, computed with the SFA model (figure 4(a)) with the same signal computed with equation (15), figure 4(b). In both cases, the 2D map is compared to the family of ellipses predicted by equation (17) (black curve). The XUV and IR transform-limit durations are 10 fs and 8 fs, while the chirp rates are \(\eta_k = 0.02 \text{ fs}^{-2}\) and \(\eta_0 = 0.01 \text{ fs}^{-2}\). The IR intensity is set to \(10^{11} \text{ W cm}^{-2}\). The results highlight the very good agreement between the SFA calculations and the approximation of equation (15). Apart from a small error in the amplitude, the geometrical features of SB+ are correctly retrieved.

Once the analogy between the two descriptions is proved, we can discuss in detail the effect of the pulse dispersion on the SB tilt. One way to do so, is to compute the local maxima of the lineouts, extracted at each delay or at each energy. These values can be calculated from equation (15), by computing the partial derivatives of the SB with respect to the frequency \(\omega',\) delay per delay, or with respect to the delay \(\tau,\) energy by energy. The result is a linear function in both cases, whose slope is represented by the parameters \(m^+_\omega\) and \(m^+_\tau\), as described in the following equations:

\[ m^+_\omega = -2\gamma_2^2\eta_k - \gamma_0^2\eta_0 \]
\[ m^+_\tau = -2\gamma_2^2\eta_k + \gamma_0^2\eta_0 + \gamma_k^2 - \gamma_0^2. \quad (20) \]

Here, the subscripts \(\omega\) or \(\tau\) denote the directions along which the maxima are calculated, whereas the superscript ‘\(^+\)’ indicates the first upper SB. If we consider the lower SB, the IR chirp rate changes its sign, meaning that the slopes are now represented by:

\[ m^-_\omega = m^+_\omega \]
\[ m^-_\tau = m^+_\tau \quad (21) \]

Figure 5 shows SB+ and SB− computed with the SFA model, compared to the functions of the maxima as extracted from the SB themselves (purple and green solid lines), and calculated with (20) and (21). Within the numerical accuracy of the calculations, the analytical prediction properly follows the numerical maxima.

Besides explaining the universal behaviour of the SB slope numerically observed in reference [13], the expressions we derived suggest a way to directly estimate the pulse GDDs from the experimental SB tilt, with very little prior information and with no need for iterative procedures. In fact, if we consider the slopes \(m^+_\omega\) and \(m^-_\omega\) and combine them to (13) and (12), we obtain the following relation for the pulses GDD:

\[ \beta_k = \frac{1}{2} \left( 1 + \frac{\sigma_k^2}{\sigma_0^2} \right) \left( \frac{1}{m^+_\tau} + \frac{1}{m^-_\tau} \right) \]
\[ \beta_0 = \frac{1}{2} \left( 1 + \frac{\sigma_0^2}{\sigma_1^2} \right) \left( \frac{1}{m^+_\tau} - \frac{1}{m^-_\tau} \right). \quad (22) \]
Therefore, by knowing the transform limited time duration of the pulses (equivalently, their bandwidths) and measuring the SB tilt, it is possible to quickly retrieve the spectral chirp of the pulses (equivalently, their bandwidths) and measuring the SB tilt, it is possible to quickly retrieve the spectral chirp of the pulses (equivalently, their bandwidths).

If we now consider the complex representation of the fields reported in equation (11) (i.e. obtained by neglecting the complex conjugate in their definition) by using the convolution property of the Fourier transform in (23), it is possible to demonstrate that for the photoelectron spectrum at zero time delay, \( S(\omega', 0) \), it holds:

\[
S(\omega', 0) \approx \frac{p^2}{\omega_0^2} \left| \mathcal{F}\{E_{\text{XUV}}(t) E_{\text{IR}}(t - \tau)\} \right|^2,
\]

where the symbol \( \ast \) denotes the convolution, while \( E_{\text{XUV}}(\omega) \) and \( E_{\text{IR}}(\omega) \) are the Fourier transforms of the complex fields, \( E_{\text{XUV}} \) and \( E_{\text{IR}} \).

In a similar way, using the integration property of the Fourier transform in (23), we can relate the SB lineout at \( \omega' = 0 \), i.e. \( \omega = \omega_1 \pm \omega_0 \), to the time convolution of the two complex light pulses:

\[
S(0, \tau) \approx \frac{p^2}{\omega_0^2} \left| \mathcal{F}\{E_{\text{XUV}}(t) E_{\text{IR}}(t)\} \right|^2.
\]

It is worth mentioning that (24) and (25) are derived from the mathematical properties of the Fourier transform and thus they are valid in general, regardless of the particular shape of the pulses.

Besides some particular lineouts, it might be interesting to evaluate the SB integrals over delay and frequency. The energy integral of the first upper SB in (15) yields:

\[
S_{\omega} = \int_{-\infty}^{+\infty} S(\omega, \tau) d\omega = A_{\varepsilon} \exp\left(-\frac{2\tau^2}{\gamma_2 + \gamma_0}\right),
\]

where \( A_{\varepsilon} \) is the amplitude, expressed as:

\[
A_{\varepsilon} = \sqrt{\frac{2\pi}{\omega_0^4}} \left( \frac{E_{\varepsilon} E_0}{4} \right) \left( \frac{1}{\gamma_2 + \gamma_0} \right)^{1/2}.
\]

The integral is therefore a Gaussian function, whose amplitude and standard deviation depend on the actual durations of the pulses. In particular, the width of \( S_{\omega} \) is equal to the width of the convolution of the real field intensity profiles \( I_{\text{IR}}(t) = |E_{\text{IR}}(t)|^2 \) and \( I_{\text{XUV}}(t) = |E_{\text{XUV}}(t)|^2 \), explaining why the energy integral of the SB signal scales with the convolution law [13].

The integral of the SB \( + \) with respect to the delay axis gives instead:

\[
S_{\tau} = \int_{-\infty}^{+\infty} S(\omega, \tau) d\tau \propto \exp\left(-\frac{\omega^2}{2\left(\frac{1}{\sigma_\omega^2} + \frac{1}{\sigma_\tau^2}\right)}\right).
\]

If we define the spectral intensities \( I_{\text{IR}}(t) = |E_{\text{IR}}(t)|^2 \) and \( I_{\text{XUV}}(t) = |E_{\text{XUV}}(t)|^2 \), \( S_{\tau} \) has the same width as their convolution. Figure 6 compares the SB integrals performed numerically over the SFA calculations of figure 4(a) (red solid curves), and estimated with convolution of the pulses temporal and spectral intensities (black dashed curves). The results show good agreement both for the energy integral (figure 6(a)) and the delay integral (figure 6(b)), proving that the SB integrals...
encode the convolutions of the pulses, both in the temporal and spectral domain.

4. Conclusions

We investigated in detail the effect of high-order dispersion terms upon the SB formation in a two-color photoemission process. By means of SFA simulations we showed that the energy-delay distribution of the SB signal qualitatively follows the GDD of the pulses as it generally happens in time-frequency distributions. Indeed, we showed that within the SVEA and the perturbative approximation, the first SB signal can be written as the Fourier transform of the time product of a signal and a gate as in a generalized Gabor transform. In the particular case of Gaussian pulses with a quadratic dispersion, we derived an analytical description of the SB signal whose contour lines are represented by ellipses. This allowed us to put the geometrical properties of the SB signal in direct relation with the temporal properties of the pulses and explain, for example, the different effect of the IR and XUV chirp on the SB tilt. In this view, we showed that one can use the information about the tilt of the upper and lower SBs to simultaneously extract the dispersion of the XUV and IR pulses. Moreover, we showed that the SB profiles at zero pump-probe delay or at the SB central energy, correspond, respectively, to the temporal and frequency convolution of the IR and XUV complex pulses, whereas the energy and delay integrals correspond to the convolution of the intensity temporal and spectral profiles. By fully deriving the chirp dependence of the SB tilt, our results explain why SHSs can be used in a FROG-CRAB approach fully deriving the chirp dependence of the SB tilt, our results explain why SHSs can be used in a FROG-CRAB approach.

![Figure 6. Comparison between the SB integral (red solid curve) and the pulse intensity convolutions (black dashed curve), in the temporal, (a), and spectral domain, (b). The remarkable agreement between the curves shows that the SB integrals encode the pulse convolutions.](image-url)

**Funding**

This project has received funding from the European Research Council (ERC) under the European Union’s Horizon 2020 research and innovation programme (Grant Agreement No. 848411 title AuDACE). ML and GI further acknowledge funding from MIUR PRIN aSTART, Grant No. 2017RKWTMY. MN acknowledges funding from MIUR PRIN, Grant No. 20173B72NB. We acknowledge support from Laserlab-Europe EU-H2020 GA No. 871124.

**Data availability statement**

The data that support the findings of this study are available upon reasonable request from the authors.

**ORCID IDs**

Bruno Moio [https://orcid.org/0000-0001-9649-7362]
Gian Luca Dolso [https://orcid.org/0000-0002-7441-2660]
Giacomo Inzani [https://orcid.org/0000-0002-0864-5976]
Nicola Di Palo [https://orcid.org/0000-0001-8386-6990]
Rocio Borrego-Varillas [https://orcid.org/0000-0002-4499-0558]
Mauro Nisoli [https://orcid.org/0000-0003-2309-732X]
Matteo Lucchini [https://orcid.org/0000-0001-6476-100X]

**References**

[1] Krausz F and Ivanov M 2009 Rev. Mod. Phys. 81 163–234
[2] Cirelli C et al 2018 Nat. Commun. 9 955
[3] Calegari F et al 2014 Science 346 336–9
[4] Vos J, Cattaneo L, Patchkovski S, Zimmermann T, Cirelli C, Lucchini M, Kheifets A, Landsman A S and Keller U 2018 Science 360 1326–30
[5] Nisoli M, Decleva P, Calegari F, Palacios A and Martín F 2017 Chem. Rev. 117 10760–825
[6] Poletto L, Villaressi P, Benedetti E, Ferrari F, Stagira S, Sansone G and Nisoli M 2007 Opt. Lett. 32 2897–9
[7] Poletto L, Villaressi P, Frassetto F, Calegari F, Ferrari F, Lucchini M, Sansone G and Nisoli M 2009 Rev. Sci. Instrum. 80 123109
[8] Eckstein M, Yang C-H, Kubin M, Frassetto F, Poletto L, Ritzé H-H, Vrakking M J J and Kornilov O 2015 J. Phys. Chem. Lett. 6 419–25
[9] von Conta A, Tehlar A, Schletter A, Arasaki Y, Takatsuka K and Wörner H J 2018 Nat. Commun. 9 1–10
[10] Reitsma G, Hummert J, Dura J, Borrego-Varillas R, Willis T, Prin X, Lepine F and Kornilov O 2019 J. Phys. Chem. A 123 3068–73
[11] Hervé M et al 2021 Nat. Phys. 17 327–31
[12] Lucchini M, Murari M, Lucarelli G D, Frassetto F, Poletto L, Mendive-Tapia D, Köppel H and Nisoli M 2019 Ultrashort fast relaxation processes in ethylene cation investigated by sub-15 fs extreme-ultraviolet pulses 2019 Conf. Lasers and Electro-Optics Europe European Quantum Electronics Conf. (CLEO/Europe-EQEC) p 1
[13] Lucchini M, Lucarelli G D, Murari M, Trabattoni A, Fabris N, Frassetto F, De Silvestri S, Poletto L and Nisoli M 2018 Opt. Express 26 6771–84
[14] Murari M, Lucarelli G D, Lucchini M and Nisoli M 2020 Opt. Express 28 10210–24
[15] Glover T E, Schoenlein R W, Chin A H and Shank C V 1996 Phys. Rev. Lett. 76 2468
[16] Pei S-C and Ding J-J 2007 IEEE Trans. Signal Process. 55 4839–50
[17] Hong K-H, Kim J-H, Kang Y H and Nam C H 2002 Appl. Phys. B 74 s231–6
[18] Kitzler M, Miliosевич N, Scrinzi A, Krausz F and Brabec T 2002 Phys. Rev. Lett. 88 173904
[19] Yakovlev V S, Gagnon J, Karpowicz N and Krausz F 2010 Phys. Rev. Lett. 105 073001
[20] Borrego-Varillas R and Lucchini M 2021 Opt. Express 29 9711–22
[21] Itatani J, Quéré F, Yudin G L, Ivanov M Y, Krausz F and Corkum P B 2002 Phys. Rev. Lett. 88 173903
[22] Mairesse Y and Quéré F 2005 Phys. Rev. A 71 011401
[23] Paul P M 2001 Science 292 1689–92
[24] Muller H G 2002 Appl. Phys. B 74 s17–21
[25] Lucchini M, Ludwig A, Kasmi L, Gallmann L and Keller U 2015 Opt. Express 23 8867–79
[26] Cattaneo L, Vos J, Lucchini M, Gallmann L, Cirelli C and Keller U 2016 Opt. Express 24 29060–76
[27] Spangenberg D, Rohwer E, Brügmann M H and Feurer T 2015 Opt. Lett. 40 1002–5
[28] Lucchini M, Brügmann M H, Ludwig A, Gallmann L, Keller U and Feurer T 2015 Opt. Express 23 29502–13