Attosecond pulse production using resonantly-enhanced high-order harmonics

V.V. Strelkov*

A. M. Prokhorov General Physics Institute of the RAS, Moscow 119991, Russia

Moscow Institute of Physics and Technology (State University), 141700 Dolgoprudny, Moscow Region, Russia

We study theoretically the effect of the giant resonance in Xe on the phase difference between the consecutive high order resonantly-enhanced harmonics and calculate the duration of the attosecond pulses produced by these harmonics. For certain conditions resonantly-induced dephasing compensates the phase difference which is intrinsic for the off-resonance harmonics. We find these conditions analytically and compare them with the numerical results. This harmonic synchronization allows attosecond pulse shortening in conjunction with the resonance-induced intensity increase by more than an order of magnitude; the latter enhancement relaxes the requirements for the UV filtering needed for the attosecond pulse production. Using a two-color driving field allows further increase of the intensity. In particular, a caustic-like feature in the harmonic spectrum leads to the generation efficiency growth up to two orders of magnitude, however accompanied by an elongation of the XUV pulse.

PACS numbers: 42.65.Ky 32.80.Rm

Attosecond pulse production using high order harmonics generated by intense laser field [1, 2] is essentially based on the phase-locking of the harmonics. This phase-locking is well understood [3, 4] for the case when there are no resonances affecting the process. However, recently much attention has been paid to the role of resonances in high harmonic generation (HHG) in gases [5–8] and plasma plumes [9, 11] (for a review of earlier studies see also [12, 13]). It was shown that when the high harmonic frequency is close to the transition to an excited quasi-stable state of the generating particle the harmonic can be much more intense than the off-resonant ones. For the HHG in plasma plumes such enhancement can be as high as an order of magnitude of even more. The XUV generation efficiency enhancement due to the giant resonance in Xe was predicted in [14] and observed in [6, 7]. Namely, the XUV near 100 eV in the spectral region of about 20 eV is more intense than the lower-frequency XUV, and the enhancement near the center of the resonance is approximately an order of magnitude.

The broadband resonant enhancement potentially allows generating attosecond pulses using resonant harmonics. This approach is interesting not only because of the higher generation efficiency of the resonant HH, but also because it essentially reduces the requirements for harmonic filtering (the resonant region is naturally standing out). However, the phase-locking of resonant HH differs from the one of the non-resonant HH [10], so the attosecond pulse production in the former case is not straightforward. In this Rapid Communication we investigate this aspect of resonant HHG both numerically and analytically. We study the effect of the resonance on the phase difference between the neighbor harmonics and calculate the duration of the attosecond pulses produced by resonant harmonics.

The time-dependent three-dimensional Schrödinger equation (TDSE) is solved numerically for a single-active electron atom in an external laser field. The method of the numerical TDSE solution is described in [15]. The model atomic potential is (atomic units are used throughout):

\[ V(r) = -a_0 \exp\left(-\frac{r^2}{b_0^2}\right) + a_1 \exp\left(-\frac{(r-r_0)^2}{b_1^2}\right) \]

(1)

The first term is the binding potential of the atomic core, and the second one is the barrier providing a bound quasi-static state with positive energy. The potential is similar to the one used earlier in the resonant HHG calculations [16, 18]. Moreover, a double-barrier effective potential was found in [19] describing autoionization in time-dependent density-functional theory.

In the potentials that we used in [16, 18] the first term is a soft-Coulomb potential, whereas in the potential [1] it is Gaussian. Such potential does not have the Coulomb 'tails' and thus does not support Rydberg states; however, it provides more freedom to simulate the properties of the desired atom. Choosing the parameters \(a_0, a_1, b_0, b_1\), and \(r_0\) of the model potential we reproduce the ionization energy of Xe atom, frequency and width of the giant resonance so that the frequency and width of the resonantly enhanced region in the calculated HH spectrum is close to those observed in Ref. [9]. The parameters of the potential used in our calculations are \(a_0 = 7.1, b_0 = 1.0, a_1 = 4.5, b_1 = 0.5\) and \(r_0 = 1.23\) a.u. Throughout this paper we simulate only the shortest electronic trajectory (if not specially stated otherwise), suppressing the others with properly defined absorbing region in the numerical box, as it was done in [20].

The laser field intensity is switched on smoothly during 4 optical cycles, then it is constant during 4 cycles, and then decreases during 4 cycles; the shape of the laser pulse is described in [21]. We are using either a single-color driving field or a two-color one. The two-color field...
The driving laser intensities and wavelengths are shown in the graph; the two-color field is given by Eq. (2) with $\alpha = 1$ and $\phi = \pi/2$. The inset shows harmonic intensities near the resonance.

The resonant-induced delay of the XUV emission smoothly decreases with the increase of the detuning from the resonance. So, in the spectral region above the resonance the resonant harmonics are emitted later than they would be emitted in the absence of the resonance. This result is in agreement with the published experimental results for HHG in Sn$^+$ [10], as well as analytical and numerical studies [14, 18]. The found delay time for the harmonics near the center of the resonant line (68 as) is close to the lifetime of the quasi-stable state (77 as). This result can be well understood within the four-step model of the resonant HHG [10]: the resonant XUV emission is delayed with respect to the non-resonant one, and the delay time is the time which the system stays in the quasi-stable state after rescattering.

The resonant-induced delay of the XUV emission smoothly decreases with the increase of the detuning from the resonance. So, in the spectral region above the resonance the resonant harmonics are emitted later than they would be emitted in the absence of the resonance. This result is in agreement with the published experimental results for HHG in Sn$^+$ [10], as well as analytical and numerical studies [14, 18]. The found delay time for the harmonics near the center of the resonant line (68 as) is close to the lifetime of the quasi-stable state (77 as). This result can be well understood within the four-step model of the resonant HHG [10]: the resonant XUV emission is delayed with respect to the non-resonant one, and the delay time is the time which the system stays in the quasi-stable state after rescattering.

To do this we consider the chirped Gaussian attopulse:

$$F(t) = \exp(-i\Omega t) \int_{-\infty}^{+\infty} \exp\left(-2\ln(2) \left(\frac{\omega'}{\Delta\omega}\right)^2\right) \left(\frac{\omega'}{\Delta\omega}\right)^{\alpha} \exp\left(i\frac{1}{2}\omega'^2 + t_r(\Omega)\omega'\right) d\omega'$$

where $\Omega$ is the central frequency of the pulse and $t_r(\Omega)$ is the emission time of the pulse. Let us assume that the chirp of the pulse is only due to the variation of the emission frequency described by the simple-man model (below we denote this chirp as the 'free-motion-induced attochirp'). Thus the derivative of the spectral phase ($\varphi + \frac{1}{2}\omega'^2 + t_r(\Omega)\omega'$) over the frequency $\omega'$ is the classical electronic return time $t_r(\omega')$. So $K = \partial t_r/\partial \omega'$. From Fig. 1 we can see that $t_r$ is an almost linear function of $\omega'$ except for the lowest and the highest part of the plateau. This linear approximation can be found from...
the solution of the Newton’s equation for the electron in the simple-man model. We find that approximately

\[ K = \frac{1}{2\omega U_p} \quad (4) \]

where \( U_p \) is the ponderomotive energy. The duration of the chirped pulse (3) depends on its spectral width \( \Delta \omega \). The shortest duration is achieved when \( \Delta \omega = \sqrt{2 \ln(2)/K} \). Substituting (4) in the latter equation we find that

\[ \Delta \omega = 2\sqrt{2\ln(2)U_p\omega_l} \quad (5) \]

The duration (FWHM of intensity) of this pulse is

\[ \tau = 2\sqrt{\ln(2)/(U_p\omega_l)} \quad (6) \]

Estimates (5) and (6) agree very well with the numerical TDSE calculations for the off-resonant harmonics. Similar estimate of the shortest attosecond pulse duration was found in [25]. Note that equation (6) shows that \( \tau \propto \sqrt{\gamma I} \) where \( I \) is the laser intensity. Since the maximum laser intensity is practically limited by the target ionization, this equation shows that the minimum attopulse duration decreases with the laser frequency decrease.

The ‘resonantly-induced attochirp’ can be estimated taking into account the delay in the resonant XUV emission. As we discussed above, this delay is the lifetime of the resonance (denoted below as \( \Delta t \)) for the XUV in the center of the resonance, and it vanishes within the width of the resonance \( \Gamma \). So the resonantly-induced attochirp is \( K_{\text{res}} = -\Delta t/\Gamma \). Having in mind that \( \Delta t = 1/\Gamma \) we find that \( K_{\text{res}} = -\Gamma^{-2} \). From this equation and equation (4) we find that \( K = -K_{\text{res}} \) for

\[ \Gamma = \sqrt{2U_p\omega_l} \quad (7) \]

The conditions of our calculations for the single-color driving field were chosen so that the latter equation is approximately satisfied: we can see that in Fig. 2 the free-motion-induced attochirp is compensated with the resonantly-induced one, so the group of harmonics above the central frequency of the resonance have approximately the same phases. In contrast to this, the parameters of the two-color field used in our calculations lead to smaller free-motion induced attochirp, so the resonantly-induced one dominates.

In Figs. 3 and 4 we show the attosecond pulses calculated using XUV from different spectral regions. Namely, using the complex amplitudes of the microscopic response \( d(\omega) \) calculated via numerical TDSE solution we find the XUV intensity:

\[ I(t) = \left| \int_{\omega = -\infty}^{\infty} M(\omega)d(\omega)\exp(-i\omega t)d\omega \right|^2 \quad (8) \]

where the used spectrum mask \( M(\omega) \) is either a Gaussian \( M_G(\omega) = \exp(-2\ln(2)(\omega - \Omega)/(\Delta \omega)^2) \) or a step-like function: \( M_{\text{step}}(\omega) = \theta(\omega - \omega_{\text{low}})\theta(\omega_{\text{high}} - \omega) \). In Fig. 3 we present the attopulses formed by resonant harmonics below the resonance (calculated using \( M_{\text{step}} \) with \( \omega_{\text{low}} = 60 \text{eV} \) and \( \omega_{\text{high}} = 96 \text{eV} \)), above it (\( \omega_{\text{low}} = 96 \text{eV} \) and \( \omega_{\text{high}} = 130 \text{eV} \)), and all the resonant harmonics (\( \omega_{\text{low}} = 60 \text{eV} \) and \( \omega_{\text{high}} = 130 \text{eV} \)). We can see that the attosecond pulse formed by the harmonics above the resonance is much shorter than the one formed by those below the resonance. This is because the above-resonant harmonics are in phase, whereas those below the resonance have significant phase differences (see Fig. 2), as it was discussed above. In the same figure we show the attopulse
formed by the off-resonance harmonics calculated using $M_G$ with the central frequency $\Omega = 155$ eV and the width $\Delta \omega = 15.2$ eV. The latter is found numerically to minimize the pulse duration; the found width and duration are very close to the predictions of eq. (3) and (6), respectively. We can see that this pulse is slightly longer and much weaker than the one formed by the above-resonance harmonics. Moreover, if all the resonant harmonics are used to produce the attopulse, its duration does not increase dramatically, see dashed line in the Fig. 3.

Metal foils or multilayer mirrors are usually applied as spectral filters [1, 2, 27, 28] to obtain attosecond pulses; such filter, in particular, can transmit well all UV higher than certain frequency. To simulate the attosecond pulses obtained with such filter we use in equation (3) the step-like mask with $\omega_{\text{high}}$ which is much higher than the cut-off frequency. The results obtained using $\omega_{\text{low}}$ well-below and well-above the resonance are shown in Fig. 4. Again, in the latter case this number is chosen to minimize the duration of the attosecond pulse. In spite of this optimization, we can see that this attopulse using off-resonant harmonics is longer and more than an order of magnitude weaker than the resonant one. Note that the parameters of the attosecond pulse formed by the resonant harmonics are not very sensitive to $\omega_{\text{low}}$ as long as it is well-below resonance; this is natural because the off-resonant harmonics are much weaker than the resonant ones. This means that practically there is much freedom in choosing such filter as long as it transmits the resonant harmonics. Moreover, if the absorption edge of the filter is far from the resonance, the filter dispersion (which is usually pronounced only in the vicinity of the absorption edge) would not affect the attosecond pulse duration. So the duration of 165 as found in our calculations is close to the one which can be experimentally obtained using harmonics enhanced by the giant resonance in Xe.

Making similar calculations for the HHG by the two-color field with the parameters $\alpha$ and $\phi$ considered above we find that the attopulse formed by above-resonant XUV is longer, and the one formed by below-resonant XUV is shorter than in the single-color field. This is the result of the attochirp behavior shown in Fig. 2 the absolute value of the total attochirp above the resonance is higher in the two-color case, below the resonance the relation is reversed. Note that this leads to smaller dephasing between the harmonics near the very center of the resonance. Since these harmonics are the most intense ones, this results in even shorter attosecond pulse than in the single-color field. Namely, the attosecond pulses formed by all XUV with frequency higher than $\omega_{\text{low}} = 60$ eV can be as short as 105 fs in the two-color case.

As it was shown both theoretically [15] and experimentally [29, 30] the harmonic yield rapidly decreases with the decrease of the driving wavelength. So the perspective of the generation efficiency increase using two-color field [31, 53] is especially important for the middle-infrared drivers considered here. To achieve maximum resonant harmonic intensity in the two-color field we chose the parameters $\alpha$ and $\phi$ of the latter so that the resonant frequency coincides with the caustic-like feature [7, 34] in the dependence of the returning electron energy on the return time. Due to this feature almost all detached electrons return back with the energy close to the one of the quasi-stable state. This leads to a further increase of the resonant harmonic generation efficiency. Fig. 5 shows the results calculated for the fundamental intensity $2.2 \times 10^{14}$ W/cm$^2$, $\alpha = 1/3$, $\phi = 1.0$ (here we take into account all the electronic trajectories in the TDSE). We compare the HHG efficiency in the two-color field with that in the single-color field having the intensity equal to the sum of the intensities of the fundamental and the second harmonic in the two-color case. Fig. 5 shows that the gain from using the two-color field with the proper parameters can be about two orders of magnitude. Together with the resonance-induced enhancement this provides the level of conversion which can be interesting for using such harmonics as an efficient source of coherent XUV in the range of 100 eV. However, the inset in Fig. 5 shows that the generation efficiency increase using such caustic-like feature leads to a loss of the attosecond nature of the emitted XUV: the calculated XUV pulse is of approximately 2 fs duration. This value is close to the time interval when the classical electrons return with energy close to the resonant one.

Thus in this paper we find conditions for which the free-motion-induced attochirp can be compensated by the resonantly-induced attochirp, leading to phase synchronization of a group of resonant harmonics. It is
shown that attopulses with duration of 165 as can be obtained using resonantly-enhanced harmonics generated in Xe. This duration is smaller than the minimal duration of the attosecond pulse formed by the off-resonant harmonics; it can be further reduced down to almost hundred attoseconds using the two-color driver. Resonant HHG enhancement leads to an increase of the attopulse intensity by more than an order of magnitude and relaxes the requirements for the XUV filtering: only harmonics much lower than the resonance should be suppressed by the filter. Using two-color field with specific parameters providing ‘caustic-induced’ enhancement of the resonant harmonics provides further (almost two-orders of magnitude) increase of the XUV intensity at the cost of increasing the pulse duration to above 1 fs.

Note that the giant resonances are observed also in other atoms, ions and molecules. The detailed investigation of their applicability for attosecond pulse production can be a natural development of the present study. However, our findings, in particular, the estimate of the laser field parameters required for the phase synchronization of the resonant harmonics given by eq. (7) should be applicable for other resonances as well.

This study was funded by RSF (grant N 16-12-10279).

[1] Paul P. M. et al. Science 292 1689 (2001)
[2] Tzallas P. et al. Nature 426 267 (2003)
[3] Ph. Antoine, A. L’Huillier, M. Lewenstein, Phys. Rev. Lett. 77, 1234 (1996)
[4] Salieres P. et al., Science 292 902 (2001)
[5] S. Gilbertson, H. Mashiko, Ch. Li, E. Moon, and Z. Chang, Appl. Phys. Lett. 93, 111105 (2008).
[6] A. D. Shiner, et al., Nature Phys. 7, 464 (2011).
[7] D. Facciala, S. Pabst, B. D. Bruner, A. G. Ciriolo, S. De Silvestri, M. Devetta, M. Negro, H. Soifer, S. Stagira, N. Dudovich, and C. Vozzi, Phys. Rev. Lett. 117, 093902 (2016)
[8] J. Rothhardt, S. H’adrich, S. Demmler, M. Krebs, S. Fritzsch, J. Limpert, and A. T’unnermann Phys. Rev. Lett. 112, 233002 (2014)
[9] R. A. Ganeev, T. Witting, C. Hutchison, et al. Phys. Rev. A 88, 033838 (2013)
[10] S. Haessler, et al., New J. Phys. 15, 013051 (2013)
[11] N. Rosenthal and G. Marcus, Phys. Rev. Lett. 115, 133901 (2015)
[12] R. A. Ganeev, J. Mod. Opt. 59, 409 (2012).
[13] R. A. Ganeev, High-Order Harmonic Generation in Laser Plasma Plumes (Imperial College Press, London, 2012).
[14] M. V. Frolov, N. L. Manakov, T. S. Sarantseva, M. Y. Emelin, M. Y. Ryabikin, and A. F. Starace, Phys. Rev. Lett. 102, 243901 (2009).
[15] V. Strelkov, A. Sterjantov, N. Shubin, V. Platonenko, J. Phys. B 39, 577 (2006)
[16] V. Strelkov, Phys. Rev. Lett. 104, 123901 (2010)
[17] M. Tudorovskaya and M. Lein Phys. Rev. A 84, 013430 (2011)
[18] V.V. Strelkov, M.A.Khokhlova, N.Yu Shubin, Phys Rev A 89, 053833 (2014)
[19] V. Kapoor, Phys. Rev. A 93, 063408 (2016)
[20] V. V. Strelkov, M. A. Khokhlova, A. A. Gonorovsk, I. A. Gonoskov, and M. Yu. Ryabikin, Phys. Rev. A 86, 013404 (2012)
[21] V. V. Strelkov, Phys. Rev. A 74, 013405 (2006)
[22] Y. Mairesse et al., Science 302, 1540 (2003)
[23] P. B. Corkum, Phys. Rev. Lett. 71, 1994 (1993)
[24] K. J. Schafer, B. Yang, L. F. DiMauro, and K. C. Klander, Phys. Rev. Lett. 70, 1599 (1993)
[25] M. A. Khokhlova and V. V. Strelkov, Phys Rev A 93, 043416 (2016)
[26] V. T. Platonenko, V. V. Strelkov, Quantum Electron. 27, 779 (1997)
[27] R. López-Martens, et al., Phys. Rev. Lett. 94, 033001 (2005)
[28] Goulielmakis E et al. Science 320 1614 (2008)
[29] P. Colosimo et al. Nature Phys. 4 386 (2008)
[30] A. D. Shiner et al. Phys. Rev. Lett. 103 073902 (2009)
[31] H. Eichmann, A. Egbert, S. Nolte, C. Momma, B. Wellegehausen, W. Becker, S. Long, and J. K. McIver, Phys. Rev. A 51, R3414 (1995)
[32] I. J. Kim, C. M. Kim, H. T. Kim, G. H. Lee, Y. S. Lee, J. Y. Park, D. J. Cho, and C. H. Nam, Phys. Rev. Lett. 94, 243901 (2005).
[33] A. S. Emelina, M. Yu. Emelin, R. A. Ganeev, M. Suzuki, H. Kuroda, and V. V. Strelkov, Opt. Express 24, 13971 (2016)
[34] O. Raz, O. Pedatuz, B. D. Bruner and N. Dudovich, Nature Photonics 6, 170 (2012)