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John Heslar, Dmitry A. Telnov, and Shih-I Chu

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1 Enhancement of VUV and EUV generation by field-controlled resonance structures of 2 diatomic molecules

3 John Heslar,¹ Dmitry A. Telnov,^{2,*} and Shih-I Chu^{1,3,†}

4 ¹*Center for Quantum Science and Engineering, and Center for Advanced Study in Theoretical Sciences,
5 Department of Physics, National Taiwan University, Taipei 10617, Taiwan*

6 ²*Department of Physics, St. Petersburg State University,
7 7-9 Universitetskaya nab., St. Petersburg 199034, Russia*

8 ³*Department of Chemistry, University of Kansas, Lawrence, Kansas 66045, USA*

Below- and near-threshold harmonic generation provides a potential approach to achieve a high conversion efficiency of vacuum-ultraviolet and extreme-ultraviolet sources for the advancement of spectroscopy. Here we perform a time-dependent density functional theory study for the nonperturbative treatment of below- and near-threshold harmonic generation of CO and N₂ diatomic molecules subject to short near-infrared laser pulses and aligned parallel to the laser field polarization. We find that with the use of different driving laser pulse shapes we can control and enhance harmonic generation through the excited state resonance structures. Depending on the pulse shape, the enhancement can reach 5 to 7 orders of magnitude as compared to the reference sine-squared laser pulse of the same duration. The results for different driving laser intensities are also presented and discussed in detail.

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10 I. INTRODUCTION

11 High-order harmonic generation (HHG) has been the
12 enabling technology for ultrafast science in the vacuum-
13 ultraviolet (VUV) and extreme-ultraviolet (EUV) spec-
14 tral regions [1–3]. High conversion efficiency of HHG is
15 a goal that the experimentalists are trying to achieve in
16 order to create sources of intense VUV and EUV radi-
17 ation [4–13]. Lately, numerous advancements have been
18 made in VUV [4, 8, 10–12] and XUV [6–8] pulse gen-
19 eration that can be applied to many areas in ultrafast
20 science and technology [5, 9, 13]. However, achieving ef-
21 ficient VUV or EUV conversion of corresponding weak to
22 moderate driving pulses is challenging and lately major
23 attention has been focused in this area of research [4–
24 7]. Conversion efficiencies for moderate peak power driv-
25 ing lasers that have been achieved are orders of magni-
26 tude behind the values that have been demonstrated with
27 loose focusing strong peak power driving pulses [14–23].
28 Recently, Wang *et. al.* [7] demonstrated an enhanced
29 highly-efficient source of femtosecond EUV pulses where
30 there enhancements arose from both wavelength scaling
31 of the atomic dipole and improved spatio-temporal phase
32 matching.

33 The goal to boost the conversion efficiency can be
34 achieved in different ways. One approach is to increase
35 the repetition rates with weaker driving lasers, but here
36 one needs to make the laser focus tight to ensure high
37 enough intensity in the focus for efficient HHG by indi-
38 vidual atoms or molecules. However, making the laser
39 focus tight significantly worsens the phase matching con-
40 ditions in the macroscopic medium. Another approach to

41 enhance HHG is to increase the intensity of the driving
42 laser pulses. With stronger driving pulses, the laser focus
43 can be made loose, thus improving the phase matching
44 conditions, but in this case the conversion efficiency suf-
45 fers from low repetition rates. The third approach to
46 boost the conversion efficiency, which we follow in this
47 paper, is to enhance the HHG signal on the microscopic
48 level, for individual atoms or molecules.

49 As we have recently demonstrated along with the ex-
50 perimentalists [M. Chini *et al.*, Nat. Photonics 8, 437
51 (2014)], below-threshold harmonics represent one such
52 possibility, where phase matching in the argon medium
53 near atomic resonances enables enhancement of coherent
54 VUV line emissions. Such emissions can be controlled by
55 temporal structures of the few-cycle driving laser field
56 with an intensity of only $\sim 1 \times 10^{13}$ W/cm² [4], which is
57 achievable directly from few-cycle femtosecond oscillators
58 with nanojoule energy. In the present contribution, we
59 explore enhancement of VUV and EUV line emissions on
60 the microscopic level by field-controlled resonance struc-
61 tures of homonuclear (N₂) and heteronuclear (CO) di-
62 atomic molecules, which in return give narrow linewidth
63 VUV and EUV radiation. In this context, we will be fo-
64 cusing on the below- and near-threshold harmonics. In
65 the past, major attention was focused on the HHG regime
66 above the ionization threshold where the semiclassical
67 three-step model and strong field approximation are ef-
68 fective to explain the process. However, neglecting the
69 electronic structure of the target and interaction between
70 the electron and molecular core results in inadequate de-
71 scription in the below- and near-threshold HHG regime.

72 In this work, we present an all-electron time-dependent
73 density functional theory (TDDFT) with proper long-
74 range potentials to study the novel HHG regime of below-
75 and near-threshold harmonics in CO and N₂ molecules.
76 In this contribution, we identify and study excited state

* d.telnov@spbu.ru

† sichu@ku.edu

77 resonance structures in CO and N₂ molecules for differ-
 78 ent driving laser intensities. Furthermore, we investi-
 79 gate the conversion efficiency and show how to improve
 80 it using different types of driving laser pulses and pulse
 81 shapes. Our calculations reveal that a five orders of mag-
 82 nitude increase of the VUV line radiation in CO can be
 83 enabled on average, and the corresponding enhancement
 84 of EUV line emission in N₂ can reach up to seven or-
 85 ders of magnitude. Finally, we compare different types
 86 of the laser pulse and pulse shapes (we have tried five
 87 variants) and make a conclusion which one provides the
 88 best enhancement. We believe that the proposed method
 89 can be applied to other atomic and molecular systems to
 90 dramatically improve the conversion efficiency through
 91 the excited state resonance structures thus opening the
 92 door to the development of compact, high flux VUV and
 93 EUV light sources.

94 The organization of this paper is as follows. In Sec. II,
 95 we briefly describe the *all electron* TDDFT formalism
 96 for the general treatment of the multiphoton dynam-
 97 ics of heteronuclear and homonuclear diatomic molecular
 98 systems. In Sec. III, we analyze the below- and near-
 99 threshold resonance structures in the radiation spectra
 100 of CO and N₂ molecules and study the evolution of these
 101 resonance structures with different driving laser intensi-
 102 ties. In Sec. IV, we investigate the role of field-controlled
 103 enhancement of these resonance structures by applying
 104 five different laser pulse shapes and also clearly show
 105 the line emissions can be enhanced orders of magnitude.
 106 While the VUV or EUV emissions are greatly enhanced
 107 orders of magnitude, they still exhibit narrow linewidths.
 108 Sec. IV contains concluding remarks.

109 II. TIME-DEPENDENT DFT FOR 110 NONPERTURBATIVE TREATMENT OF 111 DIATOMIC MOLECULES IN ONE- AND 112 TWO-COLOR LASER FIELDS

113 The basic equations of TDDFT are the time-dependent
 114 one-electron Kohn-Sham equations [24] for spin orbitals
 115 $\psi_{i\sigma}(\mathbf{r}, t)$ which involve an effective potential $v_{\text{eff},\sigma}(\mathbf{r}, t)$
 116 (in atomic units),

$$i \frac{\partial}{\partial t} \psi_{i\sigma}(\mathbf{r}, t) = \left[-\frac{1}{2} \nabla^2 + v_{\text{eff},\sigma}(\mathbf{r}, t) \right] \psi_{i\sigma}(\mathbf{r}, t), \quad (1)$$

$$i = 1, 2, \dots, N_\sigma,$$

117 where $N_\sigma (= N_\uparrow \text{ or } N_\downarrow)$ is the total number of electrons
 118 for a given spin σ , and the total number of electrons in
 119 the system is $N = \sum_\sigma N_\sigma$. The time-dependent effective
 120 potential $v_{\text{eff},\sigma}(\mathbf{r}, t)$ is a functional of the electron spin-
 121 densities $\rho_\sigma(\mathbf{r}, t)$ which are related to the spin orbitals as
 122 follows:

$$\rho_\sigma(\mathbf{r}, t) = \sum_{i=1}^{N_\sigma} |\psi_{i\sigma}(\mathbf{r}, t)|^2, \quad (2)$$

123 (the summation includes all spin orbitals with the same
 124 spin). The effective potential $v_{\text{eff},\sigma}(\mathbf{r}, t)$ in Eq. (1) can
 125 be written in the following general form

$$v_{\text{eff},\sigma}(\mathbf{r}, t) = v_{\text{H}}(\mathbf{r}, t) + v_{\text{ext}}(\mathbf{r}, t) + v_{\text{xc},\sigma}(\mathbf{r}, t), \quad (3)$$

126 where

$$v_{\text{H}}(\mathbf{r}, t) = \int \frac{\rho(\mathbf{r}', t)}{|\mathbf{r} - \mathbf{r}'|} d\mathbf{r}', \quad (4)$$

127 is the Hartree potential due to electron-electron Coulomb
 128 interaction and $\rho(\mathbf{r}, t)$ is the total electron density,

$$\rho(\mathbf{r}, t) = \sum_\sigma \rho_\sigma(\mathbf{r}, t). \quad (5)$$

129 $v_{\text{ext}}(\mathbf{r}, t)$ is the “external” potential due to the interac-
 130 tion of the electron with the external laser field and the
 131 nuclei. In the case of homonuclear or heteronuclear di-
 132 atomic molecules in a linearly polarized external laser
 133 field ($\mathbf{E}_1(t) \cdot \mathbf{r}$) or fields ($\mathbf{E}_1(t) \cdot \mathbf{r} + \mathbf{E}_2(t) \cdot \mathbf{r}$), we have

$$v_{\text{ext}}(\mathbf{r}, t) = -\frac{Z_1}{|\mathbf{R}_1 - \mathbf{r}|} - \frac{Z_2}{|\mathbf{R}_2 - \mathbf{r}|} \quad (6)$$

$$+ (\mathbf{E}_1(t) \cdot \mathbf{r} + \mathbf{E}_2(t) \cdot \mathbf{r}),$$

134 where \mathbf{r} is the electronic coordinate, $E_1(t)$ and $E_2(t)$ are
 135 the electric field amplitudes where the laser field is po-
 136 larized along the molecular axis, \mathbf{R}_1 and \mathbf{R}_2 are the co-
 137 ordinates of the two nuclei at their fixed equilibrium po-
 138 sitions, and Z_1 and Z_2 are the electric charges of the
 139 two nuclei, respectively. The internuclear separation R
 140 is equal to $|\mathbf{R}_2 - \mathbf{R}_1|$. Finally, $v_{\text{xc},\sigma}(\mathbf{r}, t)$ is the time-
 141 dependent exchange-correlation (xc) potential. Since the
 142 exact form of $v_{\text{xc},\sigma}(\mathbf{r}, t)$ is unknown, the *adiabatic* ap-
 143 proximation is often used [25–30]

$$v_{\text{xc},\sigma}(\mathbf{r}, t) = v_{\text{xc},\sigma}[\rho_\sigma] |_{\rho_\sigma = \rho_\sigma(\mathbf{r}, t)}. \quad (7)$$

144 When these potentials, determined by the time-
 145 independent ground-state density functional theory
 146 (DFT), are used along with TDDFT in the electronic
 147 structure calculations, both inner shell and excited states
 148 can be calculated rather accurately [31]. In this work, we
 149 utilize the improved van Leeuwen-Baerends LB α xc po-
 150 tential [32]. The LB α contains two empirical parameters
 151 α and β and has the following explicit form, in the adia-
 152 batic approximation,

$$v_{\text{xc},\sigma}^{\text{LB}\alpha}(\mathbf{r}, t) = \alpha v_{\text{x},\sigma}^{\text{LSDA}}(\mathbf{r}, t) + v_{\text{c},\sigma}^{\text{LSDA}}(\mathbf{r}, t)$$

$$- \frac{\beta x_\sigma^2(\mathbf{r}, t) \rho_\sigma^{1/3}(\mathbf{r}, t)}{1 + 3\beta x_\sigma(\mathbf{r}, t) \ln\{x_\sigma(\mathbf{r}, t) + [x_\sigma^2(\mathbf{r}, t) + 1]^{1/2}\}}. \quad (8)$$

153 Here, ρ_σ is the electron density with spin σ , and we
 154 use $\alpha = 1.19$ and $\beta = 0.01$ [27–30]. The first two
 155 terms in Eq. (8), $v_{\text{x},\sigma}^{\text{LSDA}}$ and $v_{\text{c},\sigma}^{\text{LSDA}}$ are the LSDA
 156 exchange and correlation potentials that do *not* have

157 the correct Coulombic asymptotic behavior. The last
 158 term in Eq. (8) is the nonlocal gradient correction with
 159 $x_\sigma(\mathbf{r}) = |\nabla\rho_\sigma(\mathbf{r})|/\rho_\sigma^{4/3}(\mathbf{r})$, which ensures the proper
 160 long-range Coulombic asymptotic potential $v_{xc,\sigma}^{\text{LB}\alpha} \rightarrow$
 161 $-1/r$ as $r \rightarrow \infty$. Note that if the conventional xc energy
 162 functional forms taken from local spin density approx-
 163 imation (LSDA) or generalized gradient approximation
 164 (GGA) [33, 34] are used, the corresponding xc potential
 165 $v_{xc,\sigma}(\mathbf{r}, t)$ will not possess the correct long-range asymp-
 166 totic ($-1/r$) behavior [35]. For the time-independent
 167 case, this exchange-correlation LB α potential has been
 168 found to be reliable for atomic and molecular DFT cal-
 169 culations [4, 27–30, 32, 36–38].

170 For the numerical solution of the TDDFT equations
 171 for diatomic molecules with proper long-range potential,
 172 we have recently developed a time-dependent generalized
 173 pseudospectral (TDGPS) method in prolate spheroidal
 174 coordinate system [27–30, 38–41]. The advantage of this
 175 method is that it allows *nonuniform* and optimal spa-
 176 tial grid discretization (denser mesh near each nucleus
 177 and sparser mesh at larger electron-nucleus separations).
 178 This improves greatly both the accuracy and the effi-
 179 ciency of the electronic structure and time-dependent
 180 calculations with the use of only a modest number of
 181 grid points. The time-dependent Kohn-Sham equations
 182 [Eq. (1)] are solved by means of the second-order split-
 183 operator technique in prolate spheroidal coordinates and
 184 in the *energy* representation [27, 42, 43] for the prop-
 185 agation of individual spin-orbitals. In this work, we
 186 extend this procedure to the numerical solution of the
 187 TDDFT calculations for the two-center homonuclear and
 188 heteronuclear diatomic molecular systems in the presence
 189 of moderate to intense laser fields.

190 Table I lists the MO energies calculated with the
 191 LB α potential, using 70 grid points in the pseudoradial
 192 spheroidal coordinate ξ and 30 grid points in the pseu-
 193 doangular spheroidal coordinate η . The agreement of
 194 the calculated valence MO energies with the experimen-
 195 tal data is well within 0.01 a.u. Also, since we will be
 196 focusing on the excited states for CO and N₂ molecules
 197 in Table II we list the vertical excitation energies for some
 198 excited states and compare with experimental data. To
 199 calculate the excited states in Table II we also use the
 200 same number of grid points as in Table I. When solv-
 201 ing Eq. (1), the pseudoradial coordinate is restricted to
 202 the domain from 0 to 40 a.u.; between 20 and 40 a.u.
 203 we apply an absorber which smoothly brings down the
 204 wave function for each spin orbital without spurious re-
 205 flections. For the time propagation, we use 4096 time
 206 steps per optical cycle (81920 steps for the whole pulse).

210 III. BELOW- AND NEAR-THRESHOLD 211 HARMONIC GENERATION: EXCITED STATE 212 RESONANCE STRUCTURES

212 After the time-dependent spin-orbitals $\psi_{i\sigma}$ are ob-
 213 tained, the induced dipole moment can be expressed as
 214 follows:

$$\mathbf{d}(t) = \sum_{i\sigma} \langle \psi_{i\sigma}(\mathbf{r}, t) | \mathbf{r} | \psi_{i\sigma}(\mathbf{r}, t) \rangle. \quad (9)$$

215 The spectral density of the radiation energy is given by
 216 the following expression:

$$S(\omega) = \frac{4\omega^4}{3\pi c^3} \left| \int_{-\infty}^{\infty} \mathbf{d}(t) \exp(-i\omega t) dt \right|^2. \quad (10)$$

217 Here ω is the frequency of radiation, c is the velocity of
 218 light. $S(\omega)$ has the meaning of the energy emitted per
 219 unit frequency range at the particular photon frequency
 220 ω .

221 We will focus first on the heteronuclear diatomic
 222 molecule carbon monoxide (CO). The HHG spectrum
 223 $S(\omega)$ is shown in Figure 1 for the CO molecule with a
 224 range of driving laser intensities $I_0 = (1 - 8) \times 10^{13}$
 225 W/cm² and a wavelength of 730 nm for a 20-optical-
 226 cycle sine-squared laser pulse [Eq. (14)]. In Fig. 1 we
 227 have clearly identified the excited state resonance peaks
 228 at photon energies 0.3931, 0.4306, and 0.4555 a.u. (all in
 229 the VUV region), which corresponds to the bound-bound
 230 transitions from $5\sigma - 6\sigma$, $5\sigma - 7\sigma$, and $5\sigma - 8\sigma$, respec-
 231 tively. These resonance peaks are similar to the atomic
 232 emission lines we recently observed and identified along
 233 with experimentalist for Ar atoms [4]. In Fig. 1(a)-(c) we
 234 study the evolution of the resonance peaks as a function
 235 of different driving laser intensities. In Fig. 1(a), we only
 236 observe the ($5\sigma - 7\sigma$) resonance peak in this intensity
 237 range. This is due to the fact the $5\sigma - 7\sigma$ resonance peak
 238 is near (almost embedded) to the non-resonance dipole
 239 allowed 7th order harmonic (H7), and also the 7σ excited
 240 state has largest value for the transition dipole (See Ta-
 241 ble III) of the three-excited states studied here. These
 242 are also the reasons that the $5\sigma - 7\sigma$ resonance peak is
 243 more intense than other resonance peaks [($5\sigma - 6\sigma$) and
 244 ($5\sigma - 8\sigma$)] in Fig. 1(a)-(c). Once the driving laser inten-
 245 sity is increased to $I_0 = 2.5 \times 10^{13}$ W/cm² we start to
 246 observe the $5\sigma - 6\sigma$ resonance peak (Fig. 1(b)). As the
 247 driving laser intensity is further increased to $I_0 = 4 \times 10^{13}$
 248 W/cm², the $5\sigma - 8\sigma$ resonance peak appears (Fig. 1(b)).
 249 In Fig. 1(c) all three excited state resonance peaks are
 250 clear and sharp peaks (narrow linewidths) up to a driving
 251 laser intensity of $I_0 = 8 \times 10^{13}$ W/cm². After increasing
 252 the driving laser intensity beyond $I_0 = 8 \times 10^{13}$ W/cm²
 253 (not shown here) the peaks become shifted and broad-
 254 ened and no resonance peaks can be clearly observed for
 255 intensities higher than $I_0 = 1 \times 10^{14}$ W/cm².

256 Now we will turn our attention to the nitrogen (N₂)
 257 homonuclear diatomic molecule. The HHG spectrum
 258 $S(\omega)$ is shown in Figure 2 for the N₂ molecule and has

TABLE I. Comparison of the field-free molecular orbital energy levels of CO and N₂, calculated with the LB α potential, and the experimental ionization potentials (in a.u.).

CO						
Orbital	1 σ	2 σ	3 σ	4 σ	1 π	5 σ
Expt. [44]	19.9367	10.8742	1.3964	0.7239	0.6247	0.5144
LB α	19.7721	10.7723	1.2601	0.7247	0.6276	0.5093
N ₂						
Orbital	1 σ_g	1 σ_u	2 σ_g	2 σ_u	1 π_u	3 σ_g
Expt. [45–47]	15.0492	15.0492	1.3708	0.6883	0.6233	0.5726
LB α	14.7962	14.7950	1.2162	0.6786	0.6199	0.5682

TABLE II. Vertical excitation energies, from the HOMO of the CO (5 σ) and N₂ (3 σ_g) molecules, calculated with the LB α potential, and the experimental values (in a.u.).

CO						
Orbital	5 $\sigma - 6\sigma$	5 $\sigma - 7\sigma$	5 $\sigma - 8\sigma$	5 $\sigma - 9\sigma$	5 $\sigma - 10\sigma$	5 $\sigma - 11\sigma$
Expt. [48]	0.3961	0.4188	0.4547	0.4623		
LB α	0.3836	0.4192	0.4523	0.4624	0.4689	0.4779
N ₂						
Orbital	3 $\sigma_g - 2\pi_u$	3 $\sigma_g - 3\sigma_u$	3 $\sigma_g - 4\sigma_u$	3 $\sigma_g - 5\sigma_u$	3 $\sigma_g - 6\sigma_u$	3 $\sigma_g - 7\sigma_u$
Expt. [48]	0.4745	0.4754	0.5277			
LB α	0.4770	0.4771	0.5273	0.5416	0.5507	0.5536

TABLE III. Excited state transition dipoles $\langle \Psi_f | z | \Psi_i \rangle$, from the HOMO (Ψ_i) of the CO (5 σ) and N₂ (3 σ_g) molecules at $t = 0$ (in a.u.).

CO			
Excited state (f)	6 σ	7 σ	8 σ
$ \langle \Psi_f z \Psi_{5\sigma} \rangle $	0.2195	0.4084	0.1889
N ₂			
Excited state (f)	3 σ_u	4 σ_u	5 σ_u
$ \langle \Psi_f z \Psi_{3\sigma_g} \rangle $	0.3890	0.1643	0.0194

the same range of driving laser intensities, wavelength
and time duration as calculated for the CO molecule in
Fig. 1. In Fig. 2 we have also clearly identified the N₂
molecules excited state resonance peaks at photon ener-
gies 0.4805, 0.5304, and 0.5660 a.u. (all in the EUV re-
gion), which corresponds to the bound-bound transitions
from 3 $\sigma_g - 3\sigma_u$, 3 $\sigma_g - 4\sigma_u$, and 3 $\sigma_g - 5\sigma_u$, respectively.
In Fig. 2(a)-(c) we study the evolution of the resonance
peaks as a function of different driving laser intensities.
The 3 $\sigma_g - 5\sigma_u$ resonance peak photon energy 0.5660 a.u.
is very close to the 3 σ_g ionization threshold (0.5682 a.u.).
Also, the 3 $\sigma_g - 5\sigma_u$ resonance peak is embedded in the
9th order harmonic (H9) which spans photon energies
both above and below the threshold. In Fig. 2(a), we
observe all three excited state (3 $\sigma_g - 3\sigma_u$, 3 $\sigma_g - 4\sigma_u$, and
3 $\sigma_g - 4\sigma_u$) resonance peaks at lower driving laser intensi-
ties ($I_0 = (1 - 2) \times 10^{13}$ W/cm²) as compared to the CO
molecule in In Fig. 1(a). In Table III the excited state
transition dipoles for the N₂ molecule are decreasing in
value the higher the excited state. This is also the trend
we observe in Fig. 2, hence, the resonance peak intensity

$S(\omega)$ has the following trend

$$(3\sigma_g - 5\sigma_u) < (3\sigma_g - 4\sigma_u) < (3\sigma_g - 3\sigma_u). \quad (11)$$

At larger driving laser intensities ($I_0 = (2.5 - 4) \times 10^{13}$
W/cm²) in Fig. 2(b) the 3 $\sigma_u - 5\sigma_u$ resonance peak starts
to broaden and as the driving laser intensity is increased
the peak is no longer distinguishable and totally embed-
ded in the 9th order harmonic at a driving laser intensity
of $I_0 = 4 \times 10^{13}$ W/cm². The 3 $\sigma_g - 3\sigma_u$ and 3 $\sigma_g - 4\sigma_u$ re-
sonance peaks in Fig. 2(b) still exhibit narrow linewidths
for these driving laser intensities. In Fig. 2(c) for stronger
driving laser intensities we observe the 3 $\sigma_g - 3\sigma_u$ and
3 $\sigma_g - 4\sigma_u$ resonance peaks start to broaden for a laser
intensity of $I_0 = 8 \times 10^{13}$ W/cm² and nearly vanish (very
broad) for laser intensities greater than $I_0 = 1 \times 10^{14}$ (not
shown here).

Since we use an absorber when solving the time-
dependent Kohn-Sham equations (1), the normalization
integrals of the spin-orbitals decrease in time thus de-
scribing ionization. The ionization probability can be
calculated from the normalization of the wave function
at the end of the laser pulse:

$$P = 1 - \prod_{i\sigma} N_{i,\sigma}(T) \quad (12)$$

where

$$N_{i,\sigma}(T) = \langle \psi_{i,\sigma}(T) | \psi_{i,\sigma}(T) \rangle, \quad (13)$$

is the normalization (survival probability) of the $i\sigma$ -th
spin-orbital after the pulse. In Fig. 3, we show the
intensity dependence of the multiphoton total ioniza-
tion (MPI) probabilities at the end of the sine-squared

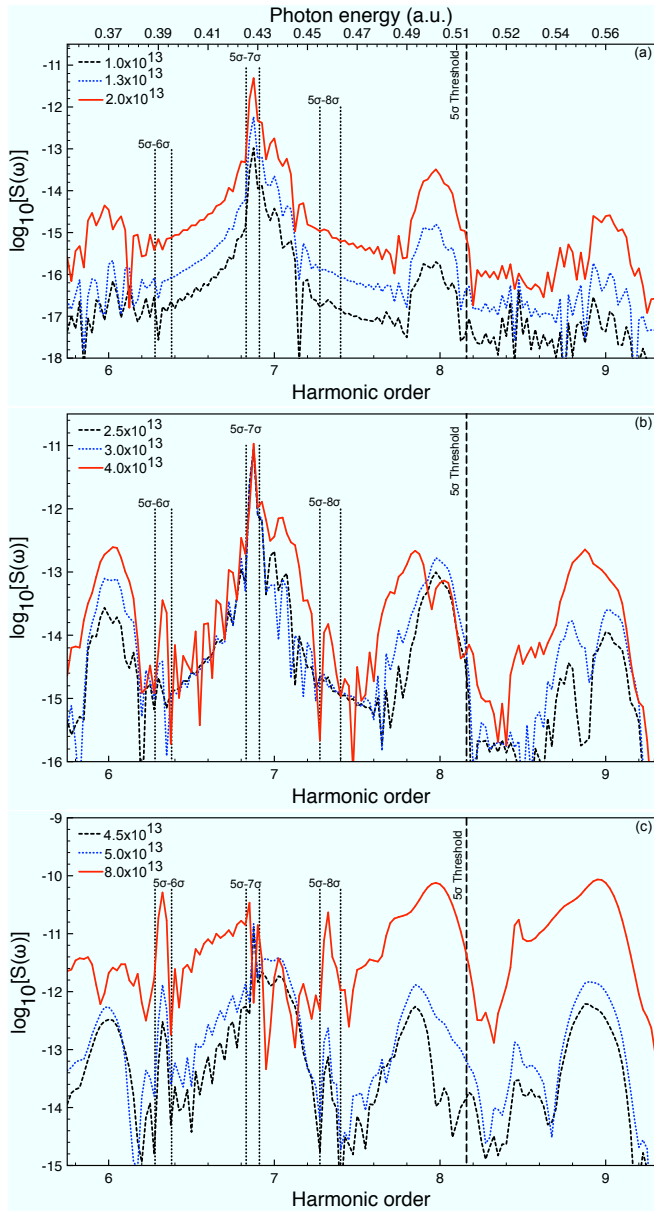


FIG. 1. (Color online) HHG spectrum $S(\omega)$ of the CO molecule in the \sin^2 laser pulse with a peak intensity of (a) $I_0 = (1 - 2) \times 10^{13}$ W/cm², (b) $I_0 = (2.5 - 4) \times 10^{13}$ W/cm², and (c) $I_0 = (4.5 - 8) \times 10^{13}$ W/cm². The laser pulse has a wavelength of 730 nm and a time duration of 20 optical cycles. Each excited state resonance peak is embedded within two vertical black dotted lines.

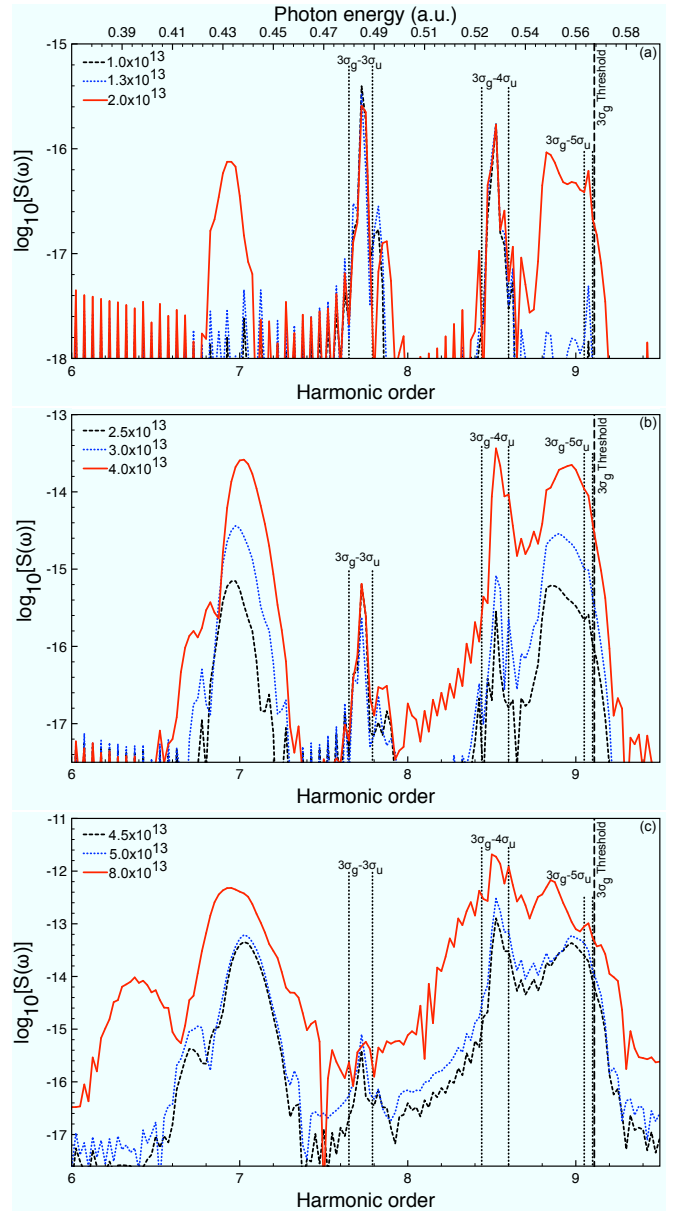


FIG. 2. (Color online) HHG spectrum $S(\omega)$ of the N₂ molecule in the \sin^2 laser pulse with a peak intensity of (a) $I_0 = (1 - 2) \times 10^{13}$ W/cm², (b) $I_0 = (2.5 - 4) \times 10^{13}$ W/cm², and (c) $I_0 = (4.5 - 8) \times 10^{13}$ W/cm². The laser pulse has a wavelength of 730 nm and a time duration of 20 optical cycles. Each excited state resonance peak is embedded within two vertical black dotted lines.

305 laser pulse ($t = T$). The degree of non-linearity
 306 [$d(\log P)/d(\log I_0)$ where P [Eq. (12)] is the total ioniza-
 307 tion probability] is close to 5. We note that the intensity
 308 range used in the calculations is beyond the applicability
 309 of the lowest-order perturbation theory, where the ioniza-
 310 tion probability P must be proportional to I_0^N , N being
 311 the minimum number of photons required for ionization
 312 (in the case of N₂ and CO molecules subject to 730 nm
 313 radiation, $N = 10$). Thus the calculated non-linearity

314 degree differs from that predicted by the lowest-order
 315 perturbation theory.

318 IV. FIELD-CONTROLLED RESONANCE 319 ENHANCED STRUCTURES

320 Hereafter, we will focus on the control and enhance-
 321 ment of the VUV excited state resonance peaks for the

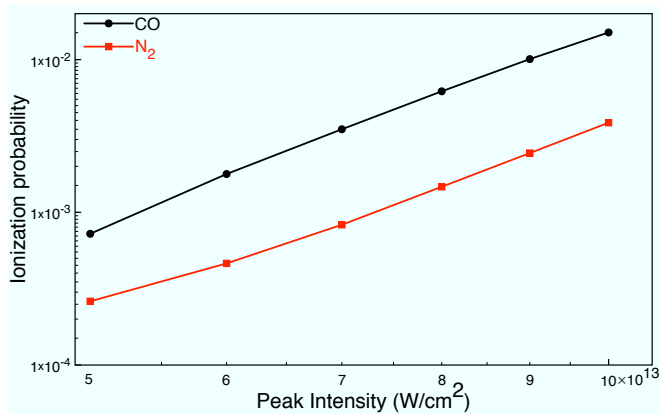


FIG. 3. (Color online) Ionization probabilities of CO and N₂ molecules versus the peak intensity of the sine-squared $[E_{LS}(t)]$ laser pulse.

CO molecule and the EUV excited state resonance peaks for the N₂ molecule. Our main goal is to enhance (increase the intensity of $S(\omega)$) the resonance structures orders of magnitude while still keeping the resonance peaks sharp and narrow. We have chosen to control and enhance the resonance structures by laser field-control, hence, with different laser pulse shapes. The NIR laser intensity will be $I_0 = 4 \times 10^{13}$ W/cm² for the CO molecule and $I_0 = 1.3 \times 10^{13}$ W/cm² for the N₂ molecule. At these laser intensities the CO and N₂ molecules excited state resonance peaks have narrow linewidths, and all three-excited states are populated and clearly visible in Fig. 1 for CO and Fig. 2 for N₂. We will try five different types of laser pulse shapes to control and increase the total ionization probability for the CO and N₂ molecules, therefore enhancing the excited state resonance structures.

We first consider the NIR sine-squared laser pulse $E_{LS}(t)$ which has the following form

$$E_{LS}(t) = F_L \sin^2 \frac{\pi t}{T_L} \sin \omega_L t, \quad (14)$$

where $T_L = 2\pi/\omega_L$ and ω_L denote the pulse duration and the carrier frequency [here we choose the laser wavelength as 730 nm ($\omega_L = 0.0624$ a.u.)], respectively; F_L is the NIR peak field strength. The pulse has a duration of 20 optical cycles (~ 49 fs) and is shown along with its Fourier transform in Fig. 4. These are the same laser parameters we used in Figs. 1 and 2. Next, the XUV (XUV will be labeled for the laser pulse, and we will use EUV for the excited state resonance peaks) laser pulse has the form

$$E_X(t) = F_X \sin^2 \frac{\pi t}{T_X} \sin \omega_X t, \quad (15)$$

where $T_X = \frac{2\pi}{\omega_X}$ and ω_X denote the pulse duration and the carrier frequency [here we choose ω_X equal to the HOMO ionization potential for CO ($\omega_X = 0.5093$ a.u.) and N₂ ($\omega_X = 0.5682$ a.u.) molecules.] Here F_X is the

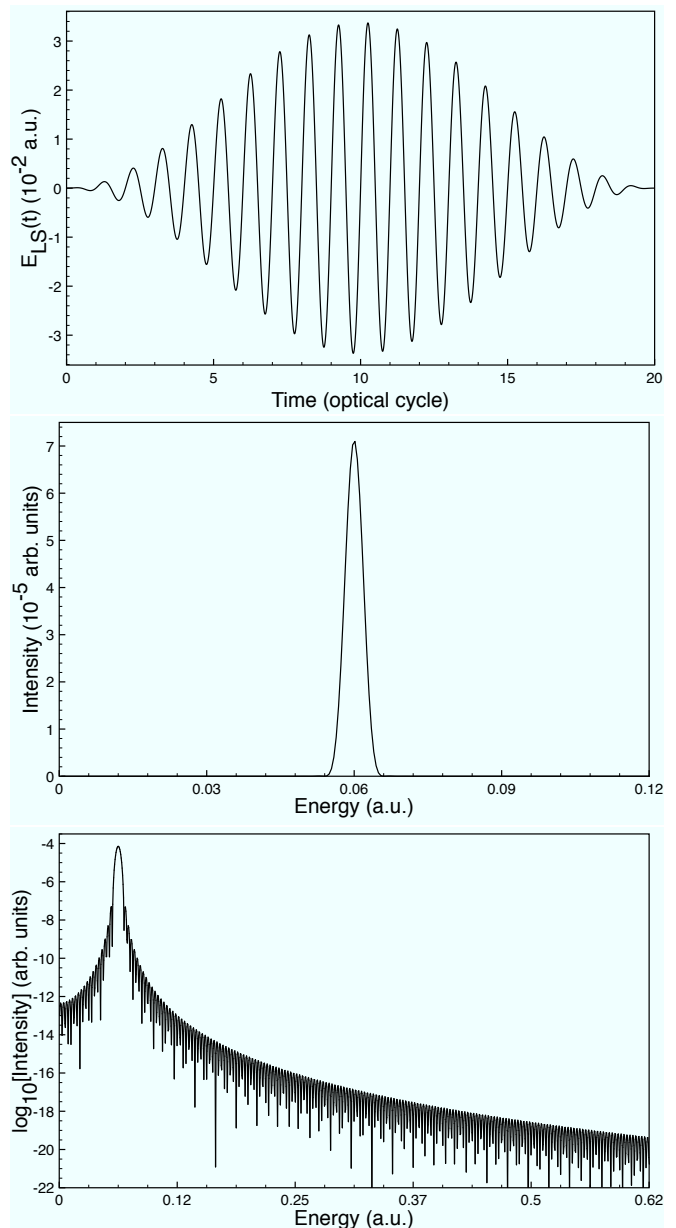


FIG. 4. (Color online) The NIR sine-squared laser pulse $E_{LS}(t)$ [Eq. (14)] as a function of time (upper panel). The laser has a peak intensity of $I_0 = 4 \times 10^{13}$ W/cm² and wavelength 730 nm. The Fourier transform of the $E_{LS}(t)$ laser pulse in linear (middle panel) and log scale (lower panel).

peak field strength of the XUV pulse, and with a pulse duration of 2 optical cycles [shown in Fig. 5 along with its Fourier transform] (which is ~ 600 as for the CO molecule and ~ 535 as for the N₂ molecule). The XUV peak intensity is 1×10^{10} W/cm². The next pulse shape we try is a flat-top NIR laser pulse $E_{LF}(t)$, which has

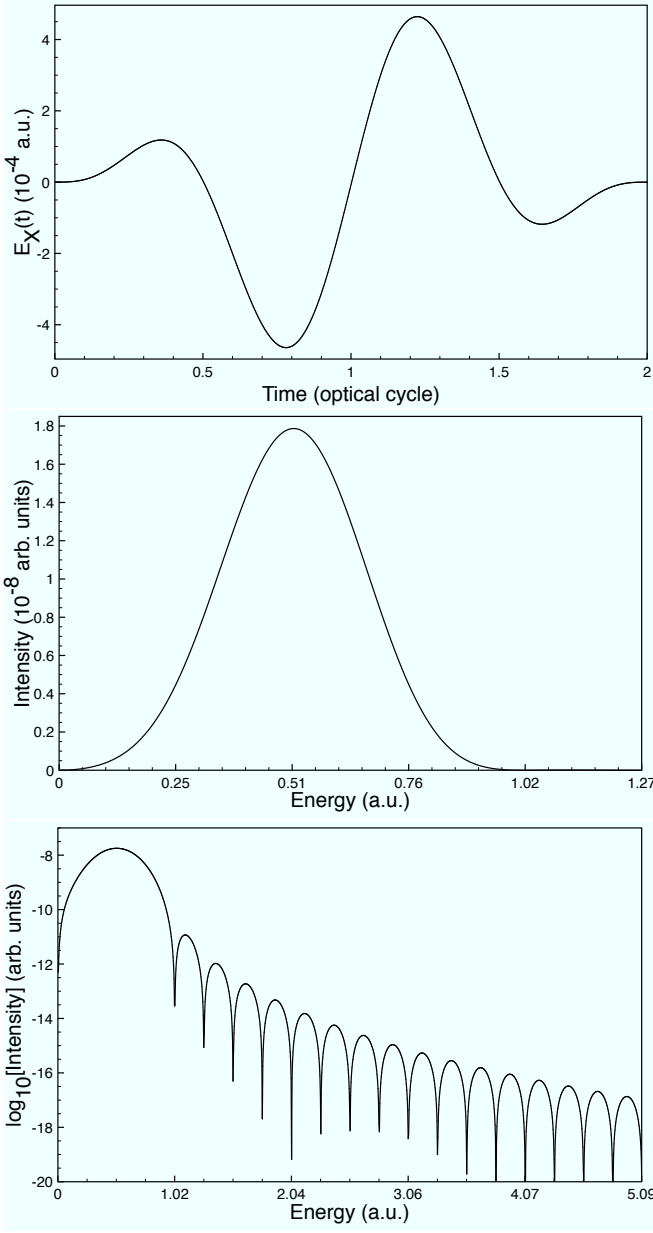


FIG. 5. (Color online) The XUV sine-squared laser pulse $E_X(t)$ [Eq. (15)] as a function of time (upper panel). The laser has a peak intensity of $I_X = 1 \times 10^{10}$ W/cm² and frequency $\omega_X = 0.5093$ a.u. The Fourier transform of the $E_X(t)$ laser pulse in linear (middle panel) and log scale (lower panel).

362 the following form

$$E_{LF}(t) = 0, \quad 0 \leq t < 5T_L, \quad (16)$$

$$E_{LF}(t) = F_L \sin \omega_L t, \quad 5T_L \leq t \leq 15T_L, \quad (17)$$

$$E_{LF}(t) = 0, \quad 15T_L < t \leq 20T_L, \quad (18)$$

363 where this pulse has a five-optical-cycle field-free propa- 368
 364 gation at the leading ($0 \leq t < 5T_L$ [Eq. (16)]) and trail- 369
 365 ing ($15T_L < t \leq 20T_L$ [Eq. (18)]) edges and has a flat-top 370
 366 central part of constant peak field strength for 10 optical 371
 367 cycles [Eq. (17)] of the total pulse duration of 20 optical 372

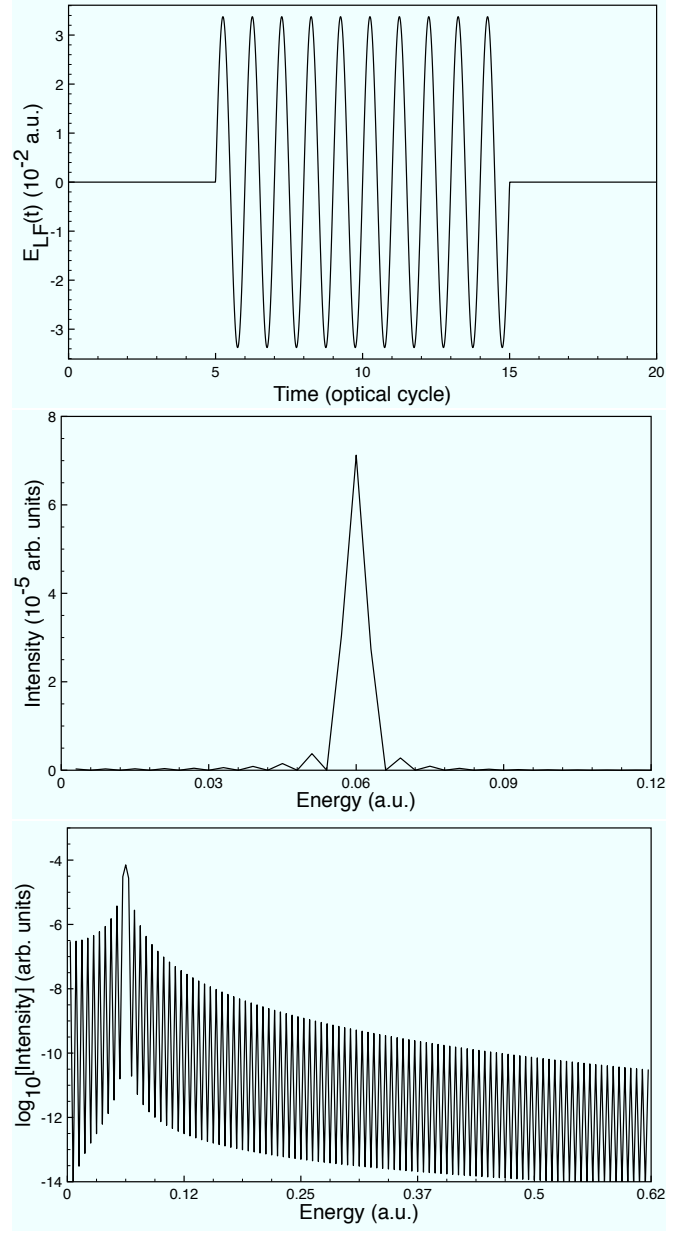


FIG. 6. (Color online) The flat-top NIR laser pulse $E_{LF}(t)$ [Eqs. (16)–(18)] as a function of time (upper panel). The laser has a peak intensity of $I_0 = 4 \times 10^{13}$ W/cm² and wavelength 730 nm. The Fourier transform of the $E_{LF}(t)$ laser pulse in linear (middle panel) and log scale (lower panel).

368 cycles. The flat-top NIR laser pulse is shown along with 369
 370 its Fourier transform in Fig. 6. Here, the laser param- 371
 372 eters (F_L , ω_L , and T_L) are the same as for the $E_{LS}(t)$ 372
 373 laser pulse in Eq. (14). The last type of pulse we try is 373
 374 a sine-squared ramped NIR laser pulse $E_{LR}(t)$, which has

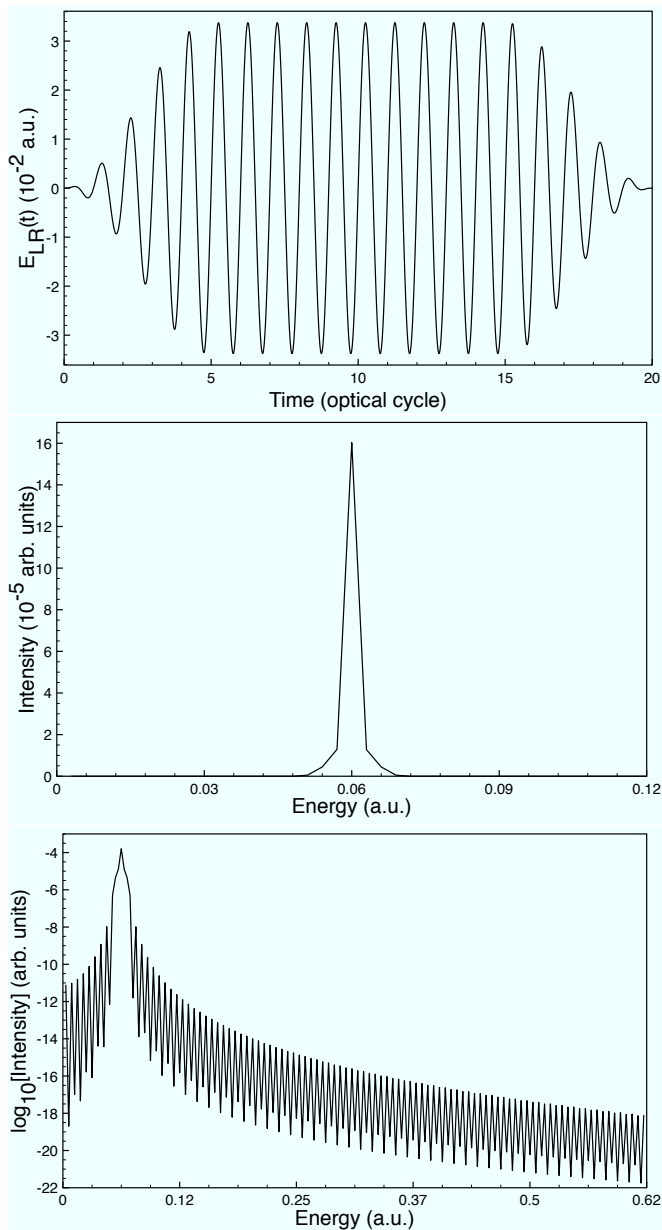


FIG. 7. (Color online) The sine-squared ramped NIR laser pulse $E_{LR}(t)$ [Eqs. (19)–(21)] as a function of time (upper panel). The laser has a peak intensity of $I_0 = 4 \times 10^{13}$ W/cm² and wavelength 730 nm. The Fourier transform of the $E_{LR}(t)$ laser pulse in linear (middle panel) and log scale (lower panel).

374 the form

$$E_{LR}(t) = F_L \sin^2 \frac{\pi t}{T_L} \sin \omega_L t, \quad 0 \leq t < 5T_L, \quad (19)$$

$$E_{LR}(t) = F_L \sin \omega_L t, \quad 5T_L \leq t \leq 15T_L, \quad (20)$$

$$E_{LR}(t) = F_L \sin^2 \frac{\pi t}{T_L} \sin \omega_L t, \quad 15T_L < t \leq 20T_L, \quad (21)$$

375 where this pulse has five-optical-cycle sine-squared ramps
 376 at the leading ($0 \leq t < 5T_L$ [Eq. (19)]) and trailing
 377 ($15T_L < t \leq 20T_L$ [Eq. (21)]) edges and has a sine-

378 squared central part of constant peak field strength for 10
 379 optical cycles [Eq. (20)] of the total pulse duration of 20
 380 optical cycles. The sine-squared ramped NIR laser pulse
 381 is shown along with its Fourier transform in Fig. 7. Also,
 382 for this pulse shape the laser parameters (F_L , ω_L , and
 383 T_L) are the same as for the $E_{LS}(t)$ laser pulse in Eq. (14).
 384 In Table IV we provide the values of $E_p = \int E^2(t)dt$ (pro-
 385 portional to the total pulse energies) for different pulse
 386 shapes shown in Figs. 4–7. As one can see, the total en-
 387 ergies differ by a factor of two at most. Certainly, this
 388 difference cannot explain a several orders of magnitude
 389 enhancement of the resonance peaks revealed by our cal-
 390 culations. It is not the total pulse energy but the pulse
 391 shape effects that play a major role here. We also note
 392 that the different pulses used here have narrow frequency
 393 distributions as seen in Figs. 4–7. In this case, the pulse-
 394 shape control can be achieved in the time domain rather
 395 than in the frequency domain. Discussion of possible ex-
 396 perimental techniques is, however, beyond the scope of
 397 the present paper.

398 In Fig. 8 we compare the excited state resonance peaks
 399 ($S(\omega)$) for the CO molecule as a function of laser pulse
 400 shapes. The red line (a) is the sine-squared reference
 401 pulse in Eq. (14), which is the pulse shape used in Figs. 1
 402 (a, b, and c) to calculate the HHG spectrum for a range
 403 of peak intensities. In Fig. 8 all NIR pulse shapes [Fig. 8
 404 (lines a–e)] applied to produce the HHG spectrum use a
 405 peak intensity of 4×10^{13} W/cm² and the XUV pulse
 406 [Fig. 8 (lines b and c)] has a peak intensity of 1×10^{10}
 407 W/cm². The black line (b) in Fig. 8 makes use of the sine-
 408 squared NIR laser pulse in Eq. (14) and a XUV attosec-
 409 ond laser pulse Eq. (15) to give a two-color (NIR+XUV)
 410 laser field process. The 600 as XUV laser pulses ($E_X(t)$)
 411 is turned on at the end (48.4 fs) of the 49 fs NIR pulse
 412 ($E_{LS}(t)$). Usually, this type of two-color laser pulse
 413 ($E_{LS}(t) + E_X(t)$) is used for pump-probe measurements
 414 where in this process [Fig. 8 (line b)] the NIR pulse would
 415 be the pump and the XUV pulse would be the probe.
 416 When a CO molecule absorbs a photon from the XUV
 417 field with the intensity 1×10^{10} W/cm², it can be excited
 418 with a substantial probability from the HOMO (5σ) to
 419 one of many singly excited states ($5\sigma - n\sigma$) or to the
 420 continuum, depending on the energy of the XUV photon
 421 absorbed.

422 In Fig. 8 (line b), when the NIR pulse comes first, it
 423 excites the HOMO (5σ) to the three dominant excited
 424 states and at the end of the NIR pulse when the XUV
 425 pulse is turned on, it populates more of the HOMO (5σ)
 426 to many excited states. When the XUV pulse comes
 427 at the very end of the NIR pulse [Fig. 8 (line b)] both
 428 pulses can be used as a “pump” pulse. Here, since both
 429 the XUV and NIR pulses are used to “pump” the ground
 430 state (5σ) to the excited states ($n\sigma$), we observe an en-
 431 hancement (~ 3 orders of magnitude larger) in the reso-
 432 nance peaks in Fig. 8 (line b) compared to the reference
 433 sine-squared NIR pulse [Fig. 8 (line a)]. In Fig. 8 (line c),
 434 we have the opposite to (line b), now the XUV pulse and
 435 the NIR pulse are turned on at the same time. The XUV

436 pulse only has a time duration of 600 as which is very
 437 short compared to the NIR pulse of 49 fs. When the XUV
 438 pulse populates the ground state to many excited states
 439 and is then turned off, the NIR pulse is strong for these
 440 already populated excited states and further one-photon
 441 (NIR) transitions can occur or the CO molecule can be
 442 ionized. Also, we see splitting of the excited state resonance
 443 peaks into subpeaks in Fig. 8 (line c). It is understood
 444 that such additional peaks arise due to excitation
 445 of other molecular levels by the combination of the XUV
 446 and NIR fields, which in fact, we recently observed and
 447 identified the mechanism for peak-splitting in He [49] and
 448 Ar [50] atoms in two-color two-photon (XUV+NIR) pro-
 449 cesses. Here, we have identified when the XUV pulse is
 450 turned on at the beginning of the calculation [Fig. 8 (line
 451 c)] along with the NIR pulse the excited state resonance
 452 peaks split into subpeaks. Therefore, this two-color laser
 453 pulse process in Fig. 8 (line c) can not produce narrow
 454 linewidth VUV line emissions for the CO molecule.

455 Figure 8 (line d) makes use of a flat-top single-color
 456 NIR laser pulse given by Eqs. (16)–(18). The flat-
 457 top pulse has a central part with a constant peak field
 458 strength of 4×10^{13} W/cm² for 10 optical cycles of the
 459 total 20 optical cycles duration. The flat-top pulse in-
 460 creases the total ionization probability to 5×10^{-5} , as
 461 compared to the reference sine-squared pulse [Fig. 8 (line
 462 a)] where the total ionization probability is equal to
 463 5×10^{-6} . Since the flat-top laser pulse increases the total
 464 ionization probability compared to sine-squared pulse at
 465 the same peak field strength (4×10^{13} W/cm²) the in-
 466 tensity of the HHG spectrum $S(\omega)$ should also increase.
 467 We see this enhancement (increase in $S(\omega)$) in Fig. 8
 468 (line d) where all resonance structures are increased ~ 5
 469 orders of magnitude as compared to the reference pulse
 470 [Fig. 8 (line a)]. While the $6\sigma - 8\sigma$ excited state resonance
 471 peaks are increased orders of magnitude [Fig. 8 (line b)],
 472 they still exhibit VUV emissions with narrow linewidths.
 473 Also, we see in Fig. 8 (line d) the flat-top laser pulse has
 474 the largest enhancement of the HHG spectrum intensity
 475 $S(\omega)$ compared to any of the other pulses or pulse shapes.
 476 In

477 Fig. 8 (line e) we use a sine-squared ramped laser pulse
 478 given by Eqs. (19)–(21) which also increases the total
 479 ionization probability (2×10^{-5}) compared to the sine-
 480 squared laser pulse (5×10^{-6}), since the sine-squared
 481 ramped laser pulse has a longer time duration with peak
 482 field strength (10 optical cycles) than that of the sine-
 483 squared pulse. Again, we observe an increase in total
 484 ionization probability so therefore the HHG spectrum in-
 485 tensity $S(\omega)$ is increased [Fig. 8 (line e)] and both the 6σ
 486 and 8σ excited state resonance peaks are enhanced by the
 487 exact order of magnitude as with use of the flat-top pulse
 488 [Fig. 8 (line d)]. The 7σ excited state resonance peak is
 489 also enhanced with use of the ramped pulse [Fig. 8 (line
 490 e)], but not as much as with the flat-top pulse [Fig. 8
 491 (line d)]. In Fig. 8 (lines a-e) we can clearly see that
 492 the flat-top and ramped laser pulses produce a narrow
 493 linewidth enhanced resonance VUV radiation for the CO

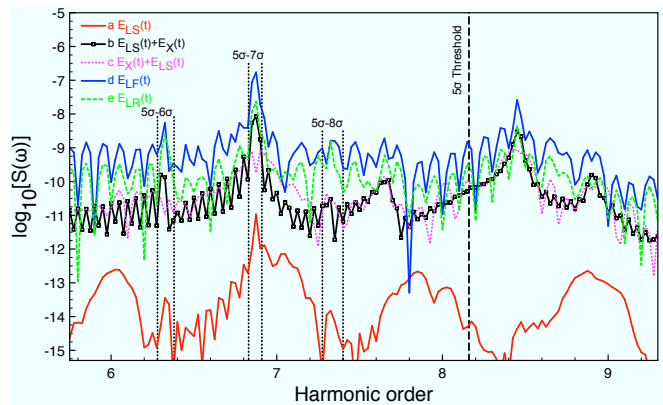


FIG. 8. (Color online) HHG spectrum $S(\omega)$ of the CO molecule using five different types of laser pulses. All NIR laser pulse shapes (lines a-e) and XUV pulses [lines b (square symbol solid black) and c (dotted magenta)] have a peak intensity of $I_0 = 4 \times 10^{13}$ W/cm² and $I_X = 1 \times 10^{10}$ W/cm², respectively. Here, for the CO molecule the XUV laser pulse $E_X(t)$ has a carrier frequency of $\omega_X = 0.5093$ a.u. and a pulse duration of ~ 600 as (2 optical cycles). a) (lower solid red line) $E_{LS}(t)$ is the \sin^2 NIR laser pulse [Eq. (14)] with a wavelength of 730 nm and a time duration of 20 optical cycles [same laser parameters as in Fig. 1(b)]. b) is the same parameters for the NIR $E_{LS}(t)$ pulse [Eq. (14)] as in a) with an additional XUV pulse $E_X(t)$ [Eq. (15)] that is turned on at the end (~ 19 th optical cycle of the NIR $E_{LS}(t)$ pulse) of the calculation. c) is the same parameters for the NIR $E_{LS}(t)$ pulse as in a) with an additional XUV pulse $E_X(t)$ [Eq. (15)] that is turned on at the beginning (0th optical cycle of the NIR $E_{LS}(t)$ pulse) of the calculation. d) (upper solid blue line) $E_{LF}(t)$ is the \sin^2 NIR flat-top laser pulse [Eqs. (16)–(18)] and e) (dashed green line) $E_{LR}(t)$ is the ramped \sin^2 NIR laser pulse [Eqs. (19)–(21)]. Laser pulses d) and e) have the same peak intensity, wavelength, and total time duration as in a). Each excited state resonance peak is embedded within two vertical black dotted lines.

498 molecules excited states ($6\sigma - 8\sigma$).

496 In Fig. 9 we compare the excited state resonance peaks
 497 (HHG spectrum $S(\omega)$) for the N₂ molecule as a func-
 498 tion of laser pulse shapes as we did for the CO molecule
 499 in Fig. 8. All pulse shapes are the same as described
 500 previously for the CO molecule in Eqs. (14)–(21). For
 501 the N₂ molecule we chose the NIR peak laser intensity
 502 (I_0) to be 1.3×10^{13} W/cm² and all other NIR laser pa-
 503 rameters (ω_L , T_L and total pulse duration) are the same
 504 as for the CO molecule and given previously throughout
 505 Eqs. (14)–(21). Here the XUV peak laser intensity (I_X)
 506 is 1×10^{10} W/cm² with a time duration of ~ 535 as, and
 507 $\omega_X = 0.5682$ a.u. is its central frequency. The details
 508 of the different pulses [Fig. 9 (lines a-e)] are described
 509 previously when discussing Fig. 8 for the CO molecule.

510 Now let us turn to the discussion of the enhanced res-
 511 onance EUV line structures for the N₂ molecules excited
 512 states ($3\sigma_u - 5\sigma_u$). In Fig. 9 all pulses or pulse shapes
 513 that we applied enhanced the resonance excited state
 514 peaks at least 7 orders of magnitude as compared to the

515 sine-squared reference pulse Fig. 9 (line a). The NIR
 516 flat-top laser pulse [Eqs. (16)–(18)] enhances the reso-
 517 nance peaks [Fig. 9 (line d)] more than any other pulses
 518 or pulse shapes in Fig. 9 compared to the reference pulse
 519 [Fig. 9 (line a)]. By using the NIR flat-top laser pulse
 520 [Fig. 9 (line d)] the resonance peaks are enhance 8 orders
 521 of magnitude as compared to the reference pulse
 522 [Fig. 9 (line a)]. Again, since the flat-top laser pulse
 523 increases the total ionization probability (1.3×10^{-5})
 524 as compared to the sine-squared pulse total ionization
 525 probability (3.0×10^{-6}), therefore the resonance peaks
 526 $S(\omega)$ value will be increased. In descending order of
 527 resonance enhancement, the next pulse to enhance the
 528 resonance peaks 7 orders of magnitude is the NIR sine-
 529 squared laser pulse in Eq. (14) and a XUV attosecond
 530 laser pulse Eq. (15) used together [Fig. 9 (line b)]. For
 531 this two-color laser pulse case in Fig. 9 (line b) the XUV
 532 laser pulse is turned on near the end ($\sim 19th$ optical
 533 cycle) of the NIR pulse and the XUV laser field has a
 534 total pulse duration of ~ 535 as. As we have discussed
 535 above for the CO molecule case for this type of two-color
 536 laser pulse case, when the XUV pulse comes near the end
 537 of the NIR pulse, both pulses (NIR and XUV) are acting
 538 as “pump” pulses. Again meaning for the case here,
 539 the N_2 molecules HOMO ($3\sigma_g$) is dominantly populated
 540 (pumped) to the excited states ($3\sigma_u - 5\sigma_u$) with the NIR
 541 pulse, and then when the XUV pulse is turned on ($\sim 19th$
 542 optical cycle) it further “pumps” the HOMO ($3\sigma_g$) elec-
 543 tron to many singly excited states or to the continuum,
 544 depending on the energy of the XUV photon absorbed.
 545 Next, in descending order of resonance enhancement is
 546 the sine-squared ramped laser pulse given by Eqs. (19)–
 547 (21) which also increases the total ionization probability
 548 (9.6×10^{-6}) compared to the sine-squared laser pulse
 549 (3.0×10^{-6}) and hence, increases the resonance peaks 7
 550 orders of magnitude [Fig. 9 (line e)].

551 Lastly, we look at Fig. 9 (line c) where the NIR sine-
 552 squared laser pulse in Eq. (14) and a XUV attosecond
 553 laser pulse Eq. (15) are used together, where for this case
 554 the XUV pulse is switched on at the beginning of the cal-
 555 culation, along with the NIR pulse. As explained above
 556 for the CO molecule this case (for the N_2 molecule) starts
 557 to cause splitting of the excited state resonance peaks
 558 into subpeaks in Fig. 9 (line c) and a dominant narrow
 559 linewidth EUV line emission is not observed for the N_2
 560 molecule. In Fig. 9 (lines a-e) we can clearly see that
 561 the flat-top and the (NIR+XUV) pulse [Fig. 9 (line b)]
 562 would have advantage in producing a narrow linewidth
 563 enhanced EUV radiation through the resonance excita-
 564 tion of the $3\sigma_u - 5\sigma_u$ electronic states in N_2 molecules.

565 V. CONCLUSION

566 In this article, we have presented a detailed investiga-
 567 tion and analysis of a novel regime of below- and near-
 568 threshold HHG, characterized by narrow linewidth reso-
 569 nance structures, which can be controlled by the tempo-

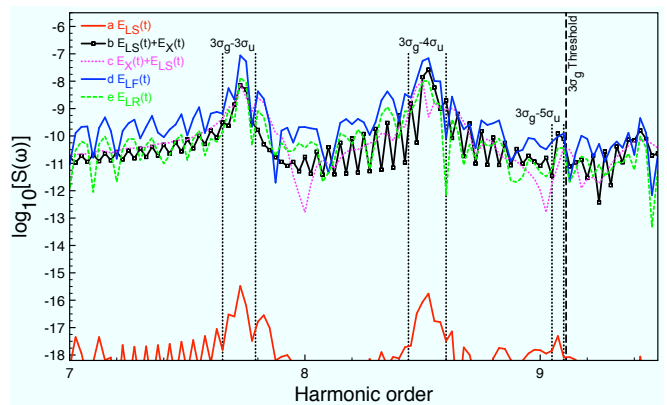


FIG. 9. (Color online) HHG spectrum $S(\omega)$ of the N_2 molecule using five different types of laser pulses or pulse shapes. All NIR laser pulse shapes (lines a-e) and XUV pulses (lines b and c) have a peak intensity of $I_0 = 1.3 \times 10^{13}$ W/cm² and $I_X = 1 \times 10^{10}$ W/cm², respectively. Here, for the N_2 molecule the XUV laser pulse $E_X(t)$ has a carrier frequency of $\omega_X = 0.5682$ a.u. and a pulse duration of ~ 535 as (2 optical cycles). Lines a-e pulse shapes are the same as in Fig. 8. Each excited state resonance peak is embedded within two vertical black dotted lines.

TABLE IV. Values of $E_p = \int E^2(t)dt$ for different pulse shapes for the peak intensity 4×10^{13} W/cm² and wavelength 730 nm (in a.u.).

Laser pulse $E(t)$	E_p
$E_{LF}(t)$	11.5
$E_{LS}(t)$	8.6
$E_{LR}(t)$	15.8
$E_X(t) + E_{LS}(t)$	8.6

570 ral behavior of a few-cycle driving laser field. Here, we
 571 demonstrate the enhancement of the VUV line radiation
 572 for the CO molecule and the EUV line radiation for the
 573 N_2 molecule with different types (NIR and XUV) and
 574 shapes of laser pulses. We make use of TDDFT with a
 575 correct asymptotic long-range ($-1/r$) potential to ensure
 576 that the individual spin-orbitals have the proper ioniza-
 577 tion potentials. Our analysis of the different laser pulses
 578 used in the calculations reveals that the best results for
 579 enhancement of the resonance structures are achieved
 580 with the flat-top, ramped, and NIR + XUV laser pulses.
 581 When using these three types of laser pulses, the VUV
 582 line emissions for the CO molecule are increased on aver-
 583 age by 5 orders of magnitude and the EUV line emissions
 584 for the N_2 molecule are increased on average by 7 orders
 585 of magnitude compared with the reference sine-squared
 586 pulse. We believe that the proposed *all-electron* TDDFT
 587 method for the study of below- and near-threshold reso-
 588 nance structures in the HHG spectra can be applied to
 589 other atomic and molecular systems and help to search
 590 for the conditions to improve the conversion efficiency in
 591 generation of coherent VUV and EUV radiation.

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