Carrier-Wave Rabi-Flopping Signatures in High-Order Harmonic Generation for Alkali Atoms

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We present a theoretical investigation of carrier-wave Rabi flopping in real atoms by employing numerical simulations of high-order harmonic generation (HHG) in alkali species. Given the short HHG cutoff, related to the low saturation intensity, we concentrate on the features of the third harmonic of sodium (Na) and potassium (K) atoms. For pulse areas of $2\pi$ and Na atoms, a characteristic unique peak appears, which, after analyzing the ground state population, we correlate with the conventional Rabi flopping. On the other hand, for larger pulse areas, carrier-wave Rabi flopping occurs, and is associated with a more complex structure in the third harmonic. These characteristics observed in K atoms indicate the breakdown of the area theorem, as was already demonstrated under similar circumstances in narrow band gap semiconductors.

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It is well known that semiconductors, when modeled as a two-level system, develop a periodic oscillation of the population inversion when interacting with constant light, a phenomenon predicted by Rabi in the 1930s, called Rabi flopping [1]. Rabi flopping has also been observed when using ultrafast optical pulses, e.g., in Refs. [2,3]. For these pulses, peculiar behavior emerges when the driven light intensity is so high that the period of one Rabi oscillation is comparable with that of one cycle of light. In this case, the area theorem has been shown to break down [3], and a new phenomenon, known as carrier-wave Rabi flopping (CWRF), emerges. These features can be schematically observed in the so-called Bloch sphere (for the definition and more details, see, e.g., Ref. [4]), presented in Fig. 1.

In particular, Fig. 1(a) depicts the conventional Rabi flopping on a Bloch sphere. For this case, the Rabi period is much larger than the driven light period and the Bloch vector, formed by the real ($u$) and imaginary ($v$) parts of the optical polarization and the population inversion ($w$) of a two-level system, spirals up starting from the south pole (corresponding to all the electrons in the ground state), reaches the north pole and returns to its initial position for the case of square-shaped pulses with an envelope area of $\Theta = 2\pi$. Here, optical oscillations are mapped to an orbit of the Bloch vector parallel to the $uv$ or equatorial plane. Additionally, oscillations of the population inversion are given by the motion in the $uw$ plane. The corresponding spectrum of the optical polarization would exhibit then two peaks centered around the two-level transition frequency.

On the other hand, Fig. 1(b) presents results for a much shorter pulse, where the Rabi period is equal to the driven light period. Even when the envelope area for this case is $\Theta = 4\pi$, it is clear that the Bloch vector does not return to the south pole, as may be expected. To the contrary, a more chaotic behavior is observed in the motion of the Bloch

FIG. 1 (color online). Sketch of the Bloch sphere showing the different regimes. (a) Schematic showing the travel of the Bloch vector for conventional Rabi flopping for a pulse with an envelope pulse area $\Theta = 2\pi$. (b) Same for carrier-wave Rabi flopping for a pulse with an envelope pulse area of $\Theta = 4\pi$. The bottom panels show the evolution of the population inversion $w$ (see the text for more details).
vector, resulting in a more complex shape in the spectrum of the optical polarization. Furthermore, the well-known area theorem of the nonlinear optics fails when this parameter regime is reached. Note that multipeak splitting of the resonance fluorescence spectrum by short pulses in the standard Rabi-flopping regime was predicted in Refs. [5–8], although this effect is due to a complex temporal interference effect, rather than CWRF.

Experiments on narrow band gap semiconductors have shown a clear signature of CWRF, which manifested itself as a split in the third harmonic of the emitted light into the forward direction [9]. Recently, Rabi flopping and the consequent coherent pulse reshaping has been experimentally observed in a quantum cascade laser [10], suggesting a new promising approach to short pulse generation. One of the advantages of atoms (relative to semiconductors) is the possibility to employ longer laser pulses and, as a consequence, to explore a broader range of laser parameters, as well as provide an alternative to carrier envelope phase (CEP) characterization. In addition, it has been shown that in semiconductors the Coulomb interaction of carriers in the bands gives rise to an enhancement of the external laser field and consequently the envelope pulse area by as much as a factor of two, considerably complicating the observed interpretation of CWRF phenomena [2,9,11].

When the atom was simplistically modeled as a twollevel system, conventional Rabi-flopping behavior and CWRF features were observed (see, e.g., Refs. [4,12] and references therein). However, it is well known that the two-level approximation breaks down when strong electric fields are applied, in particular in the CWRF regime. An important question emerges as to what extent CWRF could potentially be observed in real atoms. Prior work found Rabi-like oscillations in a model two-center molecule; however, this work presented no evidence that the CWRF regime was entered [13]. In this Letter, we demonstrate how the CWRF signatures show up in the high-order harmonic generation (HHG) spectra of real atoms. In particular, using a robust theoretical approach that accurately models both the ground and excited states of K atoms combined with realistic laser parameters, we observe clearly distinct features in the third harmonic and correlate them with the behavior of the ground state population. The latter approach is closely related to the description of semiconductors and atoms modeled as two-level systems. In order to further support our conclusions of CWRF-like behavior in K, we also compute HHG of Na atoms, in which the transition energy between the ground and first excited state is not resonant with the driven light, and, as a consequence, a conventional HHG spectra, i.e., single peaks at odd harmonics of the driven frequency, is observed. Our predictions can be tested experimentally using currently available ultrashort laser pulses of a Ti:sapphire laser with wavelengths centered in the range of 750–800 nm.

We start by describing our theoretical approach, putting special emphasis on the choice of the atomic potentials, such that our results for both ground and excited states are in excellent agreement with experimental measurements (see Table I). To find CWRF signatures, we focus on the HHG spectra and we stress that throughout the Letter no rotating wave approximation is made [4]. Since the HHG spectrum is proportional to the electron dipole moment, we can establish a one-to-one correspondence between matter, modeled as a collection of oscillators, and single atoms illuminated by a strong laser field (any macroscopic effect, such as phase matching, could be safely neglected, considering the low-order harmonic cutoff developed in alkali atoms, closely related to their low saturation intensity). To create the conditions for CWRF, we used an atomic system in which the period of a Rabi oscillation [14] (corresponding to the transition between the ground and the first excited states [15,16]) is similar to one period of the laser light. For the usual (Ti:sapphire) laser sources, such a system was provided by K atoms, with a transition energy between the ground and the first excited state of 1.61 eV (hence, close to the laser source photon energy of 1.55 eV).

Alkali atoms, due to their atomic structure (noble gas structure plus only one external electron), are well-suited to be described by the single active electron approximation (SAE). We therefore focus on such atoms to avoid the possible role of electron-electron correlations, which have been found to have an important, yet still poorly understood, role in HHG spectra [17]. Based on the SAE approximation, we use the atomic potential reported in Ref. [18] to describe K and Na atoms. Using a Hartree-Fock-based method we set the two parameters \(V_c\) and \(V_e\) in the following generic potential form: \(V_{K,Na}(r) = [V_e/(r + r_0)^2] - [V_c/(r + r_0)]\), where \(V_c\) accounts for the effect of the atomic core (nucleus plus all complete shells), \(V_e\) represents the external potential, and \(r_0 = (V_c/V_e)\). Using this method, we find the ground, \(3s\), and the first excited state, \(3p\), of K, as well as the \(4s\) and \(4p\) for Na, with precision of \(\Delta E \approx \pm 0.0084 E\) (for details, see Table I). In addition, we also compute numerically the element transition dipole \(d_{ns\rightarrow np} = \langle \psi_{ns} | z | \psi_{np} \rangle\) for both atoms \((n = 3 \text{ for Na and } n = 4 \text{ for K})\) and compare

| TABLE I. Experimental and theoretical values of the energy gap between the ground and first excited states, joint with the transition dipole both for Na and K. |
|-----------------|-----------------|-----------------|
|                  | Sodium          | Potassium       |
|                  | Experimental (NIST) | Numerical (present) |
|                  |                  |                  |
| \(3s\)           | 5.139 eV        | 5.135 eV        |
| \(3p\)           | 3.036 eV        | 3.038 eV        |
| Transition dipole| 2.49 au         | 2.40 au         |
|                  |                  |                  |
| \(4s\)           | 4.340 eV        | 4.347 eV        |
| \(4p\)           | 2.730 eV        | 2.725 eV        |
| Transition dipole| 2.92 au         | 2.79 au         |
them with the experimental values reported in Ref. [19] (see Table I). As can be deduced, our theoretical dipole values show excellent agreement with experimental measurements, indicating that the potential given above accurately represents K and Na atoms. For finding optimal laser parameters to experimentally observe CWRF in atoms, an accurate dipole moment is essential since the Rabi-flopping frequency is linearly proportional to the dipole moment as well as the electric field strength [10,14]. In addition, HHG spectra, which are the focus of our investigation, are believed to be particularly sensitive to the details of electron dynamics inside atoms and molecules. As a consequence a reliable atomic potential is crucial both to accurately describe the ground state (as many atomic potentials in the literature do), and, in the case of resonant transitions due to Rabi flopping, the excited state as well [17,20].

The main features of the CWRF are expected to appear when a two-level system is used to model an atom coupled with light [21], but the finer details of the HHG spectrum would be certainly missing, considering the atomic potential is pivotal in order to simulate adequately both a real atom and the emitted harmonic radiation. To this end, we compute HHG spectra by numerically solving the three-dimensional time-dependent Schrödinger equation (3D TDSE) in the length gauge and using the atomic potential $V_{K,Na}(r)$, given above for K and Na atoms. The harmonic yield from a single atom is then proportional to the Fourier transform of the dipole acceleration of its active electron and can then be obtained from the electronic wave function after time propagation. Our code is based on an expansion of spherical harmonics $Y_n^m$ and takes advantage of the cylindrical symmetry of the problem (hence, only the $m = 0$ terms need to be considered). The time propagation is based on a Crank-Nicolson method implemented on a splitting of the time-evolution operator that preserves the norm of the electronic wave function. The coupling between the atom and the laser pulse in the length gauge, linearly polarized along the $z$ axis, is written as $V(z,t) = E(t)z$, where $E(t)$ is the laser electric field defined by $E(t) = E_0 \sin^2(\omega_0 t/2N) \sin(\omega_0 t + \phi)$. $E_0$ is the laser electric field peak amplitude ($E_0 = \sqrt{I/I_0}$ with $I_0 = 3.5 \times 10^{16}$ W/cm$^2$), $\omega_0 = 0.0596$ a.u. ($\lambda = 765.1$ nm), $N$ the total number of cycles in the pulse, and $\phi$ the CEP. Furthermore, the laser period is $T = 2\pi/\omega_0 \approx 2.5$ fs. In the simulations presented here, we consider the case $N = 20$, corresponding to an intensity envelope of full width half maximum (FWHM) of 0.36$\sqrt{NT}$, (7.2 optical cycles $\approx 18$ fs FWHM), and $\phi = 0$ (see below for details).

As discussed in the introduction, we set the input laser frequency to the same value (in atomic units) as the energy corresponding to the transition $4s \rightarrow 4p$ of K in order to observe a CWRF-like behavior. In order to compare with a conventional situation (meaning the usual conditions for HHG), we use the same input laser parameters with Na atoms, for which the transition energy from the ground to the first excited state, $3s \rightarrow 3p$, corresponds to 2.10 eV, and is therefore nonresonant with the laser frequency.

In Fig. 2, we show the harmonic spectra computed from the 3D TDSE for both K [Figs. 2(a), 2(b), and 2(c)] and Na [Figs. 2(d), 2(e), and 2(f)] atoms. We have chosen the laser parameters to cover three different regimes, namely, for Figs. 2(a) and 2(d) the envelope pulse area, $\Theta_{K,Na}$, is close to $2\pi$. The envelope pulse area is defined as $\Theta_{K,Na} \approx d_{K,Na} E_0 \Delta t$, where $d_{K,Na}$ is the dipole transition matrix for K or Na (see Table I) and $\Delta t$ the FWHM pulse duration ($\Delta t \approx 18$ fs, i.e., $\Delta t \approx 750$ a.u.). In particular, for a laser intensity $I = 3.158 \times 10^{11}$ W/cm$^2$ ($E_0 = 0.003$ a.u.), $\Theta_K \approx 2\pi$ and $\Theta_{Na} \approx 5.4$ (for comparison, see the values used in semiconductor GaAs [9]). Panels 2(c) and 2(f) correspond to values of $\Theta_K$ and $\Theta_{Na}$, respectively, close to $4\pi$ (11.7 for K and 10.1 for Na), obtained by keeping constant the pulse duration and now using a laser intensity $I = 1.108 \times 10^{12}$ W/cm$^2$ ($E_0 = 0.0056$ a.u.). Panels 2(b) and 2(e) were chosen to have an intermediate value of intensity $I = 5.6144 \times 10^{11}$ W/cm$^2$ ($E_0 = 0.004$ a.u.), corresponding to pulse envelope areas of 8.4 and 7.2 for K and Na, respectively.

From the HHG spectra of K atoms, we observe a drastic change around the third harmonic, $\omega/\omega_0 = 3$, as the pulse envelope area increases. Our results are directly analogous to the manifestation of CWRF behavior observed in semiconductors in Ref. [9]. Note the marked contrast to the HHG spectra of Na, shown in panels 2(d), 2(e), and 2(f), where a characteristic peak is present in the third harmonic.
regardless of the envelope pulse area. The onset of this more complex behavior in K atoms for sufficiently large envelope pulse areas is in agreement with conclusions in Ref. [3], where the onset of CWRF behavior and the consequent breakdown of the area theorem was predicted for a two-level resonant system when the pulse envelope area significantly exceeds $2\pi$.

To get further insight into the physical mechanism behind the complex structure of the HHG spectra in K atoms, in Fig. 3 we present the time dynamics of the ground state population for all the cases depicted in Fig. 2. For a two-level system, used as a prototypical model for a semiconductor or a simplistic picture for a real atom, the electron dynamics due to interaction with laser light can be represented on a Bloch sphere (for details, see, e.g., Refs. [4,9]). In this case, the ground state populations and the regular Rabi oscillations can be depicted as moving along the surface of the sphere [see Fig. 1(a)]. When the CWRF regime is reached, clear signatures, corresponding to the breakdown of the area theorem, should occur on the Bloch sphere as well [see Fig. 1(b)].

Following an analogy with a two-level system, we can distinctly observe CWRF-like behavior in Figs. 3(b) and 3(c) and the corresponding counterpart in the HHG spectra [Figs. 2(b) and 2(c)]. On the contrary, a conventional behavior in the third harmonic [Figs. 2(a) and 3(a)], can be correlated with (i) ordinary Rabi oscillations for the case of K [Fig. 2(a)], i.e., the ground state is completely depopulated, even though the laser intensity is low and this would be analogous to a travel of the Bloch vector from the south to the north pole [9], and (ii) normal behavior of atoms in the strong field for all the Na cases [Figs. 2(d)–2(f)], i.e., gradual depopulation of the ground state due to laser ionization [Figs. 3(d)–3(f)].

In conclusion, we find the signatures of CWRF in real atoms, by studying the third harmonic of alkali atomic species. Analogous to the case of semiconductors, we can correlate this feature with the complex dynamics of the ground state population. Our model uses accurate values for the atomic wave function of both ground and excited states (as is evidenced by the excellent agreement between the calculated states energies with experimental values) as well as accurate laser parameters, easily achievable with the current laser technology. In particular, a Ti:sapphire laser provides laser pulses with wavelengths centered in the range 750–800 nm, very close to the 765 nm value corresponding to the transition energy $4s \rightarrow 4p$ in K. As a consequence, the experimental confirmation of our results appears straightforward. Moreover, the CWRF phenomenon in atoms could emerge as a robust alternative for CEP characterization for long pulses as can be seen in Fig. 4, where we show HHG spectra for K [Fig. 4(a)] and Na [Fig. 4(b)] and for two different values of the CEP. Note that in the case of Na (out of resonance), the third harmonic does not present appreciable differences. However, the third harmonic of K is strongly affected, in spite of the fact that the driving laser is rather long (20 cycles) in total duration. It is well known that in a conventional situation, the HHG spectra are only sensitive to the CEP changes when the driving laser field is a few-cycle pulse. Furthermore, the CWRF could be used to control the laser-induced ionization, known as a crucial ingredient for harmonic propagation, via manipulation of the ground state population.
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