Floquet calculation of harmonic generation from hydrogen molecular ions in strong monochromatic laser fields

Ts Tsogbayar and M Horbatsch

Department of Physics and Astronomy, York University, 4700 Keele Street, Toronto, Ontario, M3J 1P3, Canada
E-mail: marko@yorku.ca

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Abstract
We present Floquet calculations of high harmonic generation (HHG) for the lowest two electronic states of the $\text{H}_2^+$ ion by strong continuous-wave laser fields. We solve the non-Hermitian matrix problem to get accurate solutions to the periodic time-dependent Schrödinger equation (TDSE) by applying a pseudospectral representation combined with a complex absorbing potential method. This represents an alternative approach to direct TDSE solutions to obtain the harmonic spectra for the ion. We compare our HHG rates for the lower and upper states of $\text{H}_2^+$, which correspond to the gerade and ungerade ground states in the field-free case, with previously obtained results in the literature. We show that the enhancement of the ionization rates at the critical internuclear separation $R_c \approx 8$ au plays some role in the appearance of very strong harmonic orders $n = 5–11$ at $\lambda = 1064$ nm and $n = 5–9$ at $\lambda = 800$ nm and intensity $I = 10^{14}$ W cm$^{-2}$.

Keywords: hydrogen molecular ion, Floquet theory, harmonic generation, pseudospectral method, complex absorbing potential

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

1. Introduction

From both theoretical and experimental points of view, high-harmonic generation (HHG) is one of the most studied nonlinear phenomena for atoms and molecules interacting with an intense laser field, in which the system emits radiation at multiples of the laser frequency [1]. The physical mechanism of HHG is well understood for atoms using a three-step model [2–4]: (i) the electron is released by tunnel ionization from the atom core; (ii) the free electron is accelerated by the oscillating laser field, and later is driven back to the core; (iii) the electron can recombine with the core to emit a high-energy photon. This semiclassical formulation for the three-step model is based on the strong-field approximation by Lewenstein et al [4]. The model predicts a plateau in the harmonic spectra where many harmonics have similar strength, and it ends with a sharp cutoff. At the cutoff the maximum energy is well approximated by the simple and universal formula $I_p + 3.17U_p$, where $I_p$ is the ionization potential of the atom and $U_p$ is the pondermotive potential, defined as $U_p = (F/2\omega)^2$, with $F$ the laser electric field strength and $\omega$, the angular frequency in atomic units, respectively. The cutoff position can be estimated by $N_{\text{max}} = (I_p + 3.17U_p)/\omega$.

For symmetric diatomic molecules Kopold et al [5] extended the discussion of a (semi)-classical cutoff formula. They investigated two phenomena, which can become particularly important if one considers dissociating molecules, i.e., systems at large internuclear separation $R$. The so-called simpleman formula (1) is modified, since the ionized electron produced at nucleus $A$ upon re-collision can be re-combining either at nucleus $A$ or $B$. This can lead to a cutoff that is higher than the atomic one given in equation (1). In addition, there is the possibility that the field ionizes an electron at atom $A$, accelerates it, and recombination occurs directly at atom $B$. These classical cutoff positions have to be taken with a grain of salt, since they ignore the potential role of the Compton profile.
of the initial state, and the argumentation based on electron localization during ionization and recombination makes sense only at large \( R \), if the molecular orbital nature of the states is taken into account. Nevertheless, [5] serves to illustrate that the cutoff energies represent stationary points at which enhanced HHG should be observed. Evidence is presented in [5] from quantum calculations in a zero-range model potential for molecular cutoffs higher than equation (1), increasing the coefficient from 3.17 by up to a third for the cases considered.

Theoretical investigations for diatomic molecules, such as the \( \text{H}_2 \) molecule and the \( \text{H}_2^+ \) ion had been initially carried out by Krause et al [6] and Zuo et al [7, 8]. They performed a direct numerical solution of the time-dependent Schrödinger equation (TDSE) to obtain the HHG spectra. An alternative approach is the Floquet formalism which was employed successfully by Potvliege and Shakeshaft [10] to obtain the HHG spectra for \( \text{H} \) atoms using Sturmian basis functions. A treatment of HHG for complex atoms in intense laser fields based on \( R \)-matrix-Floquet theory has been given by Burke et al [11, 12]. Yet another method for time-periodic systems is the Floquet approach combined with complex rotation of the coordinate [13, 14].

For atomic hydrogen, the hydrogen molecule and molecular ion calculations of HHG spectra within the Floquet method combined with a complex rotated coordinate have been extensively investigated by Chu and his co-workers [15–19]. In those works a generalized pseudospectral approach was used for the spatial discretization of the resonant Hamiltonian, and a non-Hermitian split-operator technique was implemented before [20] to calculate the ionization rate for the \( \text{H}_2^+ \) ion had been initially carried out [21, 22]. The solution \( \Psi(t, r) \) of the TDSE for the \( \text{H}_2^+ \) ion was obtained by implementing before [20] to calculate the ionization rate for the lowest two electronic states of the ion by strong continuous laser fields in the low-frequency limit. We compare our results with those obtained in [8] and [18].

In the present article our goal is to report on HHG spectra for a linearly polarized intense laser field (with electric field aligned with the molecular axis) for the lowest two states of \( \text{H}_2^+ \), using the Floquet approach combined with a complex absorbing potential (CAP). This methodology was implemented before [20] to calculate the ionization rate for the lowest two electronic states of the ion by strong continuous laser fields in the low-frequency limit. We compare our results for the HHG with those obtained in [8] and [18]. Our method differs from that of [18] in that we do not use a time propagator, but assemble the wave function from the Floquet eigenstates.

The organization of this paper is as follows. In section 2.1 we start with the basic theoretical methodology to solve the TDSE for the \( \text{H}_2^+ \) ion, while making use of the time periodicity. In section 2.2 we provide details of how to get HHG spectra within the non-Hermitian Floquet approach. Section 3 discusses our results for HHG spectra for the lowest two states of the \( \text{H}_2^+ \), which is followed by conclusions.

2. Theory

2.1. The Floquet Hamiltonian

In the presence of an external field, for a diatomic molecule, such as \( \text{H}_2^+ \) the induced electronic motion happens on a faster time scale than the nuclear motion. Thus, we treat the dynamics in the Born–Oppenheimer approximation, in which the two nuclei are fixed (on the au time scale), and only the electronic motion is taken into account. The field-free electronic Hamiltonian of the \( \text{H}_2^+ \) molecule can be written in atomic units as

\[
H_0 = -\frac{1}{2} \nabla^2 - \frac{1}{|\mathbf{r} + \frac{e}{2}\mathbf{e}_z|} - \frac{1}{|\mathbf{r} - \frac{e}{2}\mathbf{e}_z|}, \tag{2}
\]

where \( \mathbf{r} \) is the electron position vector and \( R \) is the internuclear separation.

If we assume that the interaction of the electron with the external electric field \( V_F(t, r) \) is periodic in time with period \( T = 2\pi/\omega \), that is, \( H(t, r+T) = H(t, r) \), according to Floquet theory [21, 22], the solution \( \Psi(t, r) \) to the TDSE for the system

\[
i\frac{\partial}{\partial t} \Psi(t, r) = H(t) \Psi(t, r) = [H_0 + V_L(t, r)] \Psi(t, r), \tag{3}
\]

can be written as:

\[
\Psi(t, r) = e^{(-iE_F t)} \Phi(t, r), \tag{4}
\]

\[
\Phi(t, r) = \Phi(t + T) = \Phi(t), \tag{5}
\]

where \( E_F \) is called the Floquet quasi-energy, and the \( \Phi(n)(r) \) obey time-independent coupled-channel equations. The last expression in (5) represents the truncated ansatz used in practical calculations.

Substitution of the solution ansatz (4) into the Schrödinger equation (3) leads to a time-dependent eigenvalue problem:

\[
H_F(t, r) \Phi(t, r) = E_F \Phi(t, r), \tag{6}
\]

where the Floquet Hamiltonian \( H_F(t, r) \) is defined as

\[
H_F(t, r) = H(t, r) - i\frac{\partial}{\partial t}. \tag{7}
\]

In this work we assume that the external field is provided by a linearly polarized monochromatic laser whose electric field is aligned with the internuclear axis of the \( \text{H}_2^+ \) ion, and that the dipole approximation is valid. Then the interaction \( V_L(t, r) \) in length gauge takes the form

\[
V_L^{(l)}(r, t) = F_z \cos \omega t, \tag{8}
\]

where \( F \) is the laser field strength. The length gauge is indeed more appropriate for low-frequency fields [20], and we employ this gauge for \( \omega = 0.0428, 0.05695, \) and 0.08565 au in the present work, which correspond to wavelengths \( \lambda = 1064, 800 \) and 532 nm, respectively.

For the solution of the Floquet (steady-state) Hamiltonian (7), the time variable \( t \) is treated in analogy to a coordinate variable, and the Schrödinger equation (6) is solved as for the stationary states of the time-independent Schrödinger equation. Once we find \( \Phi(t, r) \) from the steady-state Schrödinger equation (6), we obtain the solution \( \Psi(t, r) \) to the TDSE (3) via equation (4).

We choose prolate spheroidal coordinates to deal with the \( \text{H}_2^+ \) ion (as described in [20]), in which the Born–Oppenheimer treatment (equation (2)) gives an analytic solution to the Schrödinger equation [23–25].

The final time-independent coupled equations for the non-Hermitian matrix problem in length gauge to be implemented are
As shown by Telnov and Chu [18] in Floquet theory the HHG forms. one uses the dipole operator (12) or the velocity or acceleration rates $/Gamma_1$ versus the value from Telnov and Chu [18]; (b) plot for more harmonics than shown in (a); here $N_{\text{max}} \approx 15; (c) I = 1 \times 10^{14}$ W cm$^{-2}$, and $N_{\text{max}} \approx 17$. The green line in (b), (c) connects the data of [18] while the red line connects the present data. The blue error bars are based on the calculations with different $\mu_c$. The vertical dashed lines indicate the semi-classical cutoff values $N_{\text{max}}$.

$$[H_0(\mu, v) - i\eta^{(0)} W(\mu) ] \phi_n(\mu, v) + \frac{1}{2} F z [\phi_{n-1} + \phi_{n+1}]$$

$$= (E_F - n\omega) \phi_n(\mu, v) + \frac{1}{2} F [\phi_{n-1} + \phi_{n+1}];$$

$$n = 0, \pm 1, \pm 2, \ldots, \pm N_F.$$  \hspace{1cm} (9)

Details of the discretization of this equation and calculation of the resonance parameter $E_F$ in equation (9) are given as

$$H_0 = -\frac{1}{2} R^2 (\mu^2 - \nu^2) \frac{\partial}{\partial \mu} \left[ (\mu^2 - 1) \frac{\partial}{\partial \mu} \right]$$

$$+ \frac{\partial}{\partial \nu} \left[ (1 - \nu^2) \frac{\partial}{\partial \nu} \right] + \frac{\mu^2 - \nu^2}{\mu^2 - 1} (1 - \nu^2) \frac{\partial^2}{\partial \nu^2}$$

$$- \frac{4\mu}{(\mu^2 - \nu^2)};$$

and

$$W(\mu) = \Theta(\mu - \mu_c)(\mu - \mu_c)^2;$$

where $\Theta$ is the Heaviside step function, $\eta$ is a small positive parameter, and $\mu_c$ determines the ellipse outside of which the CAP dampens the outgoing wave in the asymptotic region.

2.2. Calculation of HHG spectra from the non-Hermitian Floquet approach

Once we have found the time-dependent Floquet wave function $\Psi(\mathbf{r}, t)$ via equation (4), we can compute the time-dependent dipole moment along the internuclear axis, $d(t)$, as

$$d(t) = \langle \Psi(\mathbf{r}, t)|z|\Psi(\mathbf{r}, t) \rangle.$$  \hspace{1cm} (12)

Following [18, 26], the $n$th-order harmonic generation (HG) rates $\Gamma_n$ (the number of photons with frequency $n\omega$ emitted per unit time) are calculated by the Larmor formula

$$\Gamma_n = \frac{4\pi^3 c^3 |d_n|^2}{3e^3},$$

where $c$ is the speed of light, and $d_n$ is the Fourier transform of the time-dependent dipole moment (12) as

$$d_n = \frac{1}{T} \int_0^T dt \exp(in\omega t) d(t).$$  \hspace{1cm} (14)

As shown by Telnov and Chu [18] in Floquet theory the HHG rates are obtained to the same accuracy irrespective of whether one uses the dipole operator (12) or the velocity or acceleration forms.

3. Results and discussion

3.1. HHG rates for the equilibrium separation $R = 2$ au

We attempted first to obtain the HHG spectra previously reported by Telnov and Chu [18]. In our grid representation we have two main parameters, and the results for the HG spectra appear to be sensitive to them, implicitly via the Floquet wave function $\Psi(\mu, v, t)$. One of two (artificial) parameters which control the wave function is $\mu_c$, which determines the region where the CAP starts. Another one is the absorbing strength parameter $\eta$. Ideally, the results ought to be insensitive to these two parameters. It is obvious that the value of $\mu_c$ should be larger than the quiver radius of a free electron $\alpha_0 = F/\omega^2$ (in atomic units), because the main contribution to the HG spectra comes from the free electron driven back to its parent ion or two-centre core.

In analogy to the $\eta$-trajectory in the calculation of the resonance parameter $E_F^{(0)}$ in [27], we initially obtain the HHG rate $\Gamma_n$ for varying $\eta^{(0)}$. Within a certain range of the $\eta^{(0)}$-trajectory, namely where the complex eigenenergy value $E_F^{(0)}$ stabilizes, the resonance wave function is accurate, and in turn, it should yield accurate HHG spectra there.

In panel (a) of figure 1 we show the HHG rate $\Gamma_{15}$ versus $\eta^{(0)}$ for the lower state of the $H^+$ ion at the equilibrium separation ($R = 2$ au) in the field of intensity $I = 5 \times 10^{13}$ W cm$^{-2}$ and wavelength $\lambda = 532$ nm. In panel (a) we show results for $\Gamma_{15}$ for $\eta^{(0)} \geq 0.05$ (at $\eta^{(0)} \approx 0.05$ the computation is inaccurate). We chose four different $\mu_c$ values, in the range $13.5 \leq \mu_c \leq 16.5$ au, which are much larger than $\alpha_0 = 5.15$ au. The bottom green line shows the result obtained by Telnov and Chu [18] for comparison (the dependence on the complex scaling parameter is not given in that work). Our results for $\Gamma_{15}$ are higher by up to a factor of 2.5, as compared to with the value $\Gamma_{15} = 4.17 \times 10^{-22}$ au given in [18]. In panel (b) we show the HHG spectrum. We note that for orders $n > 15$ the agreement with the results of [18] is good and is independent of the chosen value of $\mu_c$. In all plots of the HHG rate we do not show results for order $n = 1$, because they are usually much higher. The cutoff position is around $n = 15$, which is indeed consistent with our result. For each harmonic order we use four different values of $\mu_c$ to compute $\Gamma_n$, and
I \times 10^{14} \text{ W cm}^{-2}

\begin{align*}
\text{Harmonic order} & \quad 10^{-10} \\
\text{Harmonic order} & \quad 10^{-12} \\
\text{Harmonic order} & \quad 10^{-14} \\
\text{Harmonic order} & \quad 10^{-16}
\end{align*}

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\text{Harmonic order} & \quad 10^{-12} \\
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\text{Harmonic order} & \quad 10^{-12} \\
\text{Harmonic order} & \quad 10^{-14} \\
\text{Harmonic order} & \quad 10^{-16}
\end{align*}

use them to define an average value with standard deviation. Panel (c) shows the same plot for the HHG rate for the doubled intensity, $I = 2 \times 10^{14} \text{ W cm}^{-2}$. The cutoff law is clearly obeyed around $n = 17$ by both the present and previous [18] results. As compared to [18] our HHG rates are higher above the cutoff but also less certain.

In figure 2 data are shown for increased laser intensities. In panel (a) of figure 2 in the HHG spectra the cutoff position moves up to 19, but the calculated spectrum extends the plateau to higher orders. Our result agrees well with that obtained by Telnov and Chu [18]. In panel (b) of figure 2 we show the same plot for $I = 5 \times 10^{14} \text{ W cm}^{-2}$. In this case although the general features of the obtained HHG spectrum follow those of [18], it does show significant deviation at certain harmonic orders, namely for $n = 5, 11$ and 27. To check our answer carefully, we gradually increased the number of Floquet channels, because the HHG rates at higher harmonic orders require a higher number of photon couplings. In panel (c) of figure 2 we display the same plot as shown in panel (b), but the number of Floquet channels $N_F$ is 72, 82 and 86, respectively. Thus we know that the results are converged at $N_F = 86$ in the HHG order range presented in our plots. The data in figure 2(c) are based on a matrix diagonalization with $N_F = 86$, with the specified truncation imposed in equation (4) when computing (12).

Next we continue with HHG rates while moving towards the low-frequency limit. In figure 3 we show the rates for the lower (red) and upper (blue) states of the $\text{H}_2^+$ ion for laser fields of $I = 1 \times 10^{14} \text{ W cm}^{-2}$ and $\lambda = 1064 \text{ nm}$. All HHG calculations are carried out with $\eta(0) = 0.25$. Panel (a) shows the spectrum at internuclear separation $R = 5 \text{ au}$, while panels (b), (c) demonstrate corresponding results at $R = 10 \text{ au}$. Given that $\omega = 0.0428 \text{ au}$, the cutoff positions given by the classical formula (1) are found around $n = 45$ and $n = 43$ at $R = 5 \text{ au}$ and $R = 10 \text{ au}$, respectively, and are shown by vertical dashed lines for the lower state in figure 3 (for the upper state, the cutoff position is close to it, since both states have almost the
same ionization potential at large internuclear separations). As discussed by Bandrauk and co-workers in [7, 8], these classical cutoff positions are referred to as the atomic plateau. They argue that a first plateau region can be identified as a molecular plateau: its cutoff occurs at harmonic order

\[ n_M = \frac{2\Omega_R}{\omega} = \frac{2d_0F}{\omega} \approx \frac{RF}{\omega} \]  

(15)

where \( \Omega_R \) is the Rabi frequency for driving transitions between the \( 1\sigma_g \) and \( 1\sigma_u \) states and the transition dipole moment \( d_0 \) grows towards \( R/2 \) with increasing \( R \) (for details cf [8]). According to this model in which the two lowest states are driven resonantly, since \( \omega \geq \epsilon_{1\sigma_g} - \epsilon_{1\sigma_u} \), the values of the cutoff positions can be found at \( n_M = 5 \) for \( R = 5 \) au, and \( n_M = 11 \) for \( R = 10 \) au, respectively. These calculated values of \( n_M \) can be observed in the data given in figure 3. The harmonics for \( n < n_M \) are produced predominantly by the bound parts of the wave function. One can interpret the molecular plateau process as localized electron removal in one well, acceleration by the laser field towards the other well followed by recombination. Yu et al [9] have recently performed TDSE and Maxwell–TDSE calculations to demonstrate the feasibility of attosecond pulse generation in this HG regime.

Panel (b) of figure 3 shows the HHG spectrum calculated by equation (14) for the lower state of the ion and its comparison with that obtained in [8] (green). We note that the agreement between the Floquet result and the calculation for a finite 30-cycle pulse is excellent up to order 39. Beyond this order the harmonics for the finite pulse with square envelope continue to be strong, while the Floquet results fall off. The comparison of \( N_F = 82 \) and \( N_F = 86 \) calculations indicates that the present Floquet calculations for \( \lambda = 1064 \) nm are converged.

### 3.2. Harmonic generation rates as a function of internuclear separation

In this subsection we present HG rates for moderate orders \( n \), i.e., in the molecular plateau region at the intensity \( 1 \times 10^{14} \) W cm\(^{-2} \) as a function of internuclear separation \( R \). In figure 4 we show the ionization rates for the lower and upper states of the ion for two wavelengths of the laser field, 1064 nm and 800 nm.

In figure 5 we show the contour plots of the localized probability density for the lower and upper states of the ion for three internuclear separations. The states are quite distinct for \( R = 2, 4 \) au, but acquire a very similar character due to the resonant mixing at \( R = 8 \) au. Plots of Floquet Siegert-state probability densities for atomic hydrogen were obtained recently for similar laser fields [29]. In the present work the use of a CAP suppresses the outgoing wave, and therefore we cannot confirm the conclusions of [29] about the shift in location where the ionized electron is produced. In terms of large \( R \) the \( H^+ \) ionization rates (figure 4) show structures which may be caused by resonances with dressed excited Floquet states as in the atomic limit [29].

Since the physical interpretation for the enhancement of the ionization rates and its connection with the Floquet probability density demonstrated in figures 4 and 5 respectively are discussed in [30–33] and later in [20, 27], we do not repeat it here. Our goal is to demonstrate how the enhancement of the ionization rate in certain \( R \)-regions affects the HG rates within the region of the molecular plateau, i.e., for \( n \leq n_M \) (15).

In figure 6 we show the rates \( \Gamma_n \) as functions of \( R \) for the lower (red) and upper (dashed blue) states of the ion using a common linear scale. As can be seen in figure 6(c), when \( R \) varies from the equilibrium separation 2 to 12 au, the fifth
Harmonic for the lower and upper states has very similar large peaks around $R \approx 5$ au, where the first enhancement of the ionization rate (figure 4(a)) occurs. Note, however that the ionization rate patterns for the two states are not as similar as those of $\Gamma_5$. Observing $\Gamma_7$ and $\Gamma_9$ in panels (d), (e) we find a shift in the peaks towards larger $R$. For the lower state a large peak in the ionization rate appears around $R \approx 9$ au (figure 4(a)), and $\Gamma_9$ also displays a peak there (figure 6(e)). Meanwhile, the upper-state $\Gamma_9$ rate deviates for $R > 8$ au, somewhat in accord with its decrease in ionization rate.

The HG rates for higher orders of $n$ (beyond $n = 11$) become smaller, thus we do not show them here. Together figures 4(a) and 6 demonstrate that the enhancement of the ionization rate for the lower and upper states of the hydrogen molecular ion can be linked to an enhancement of the harmonic generation rates in certain $R$-ranges. This happens for harmonic orders within the molecular plateau region $n < n_M$.

As a further demonstration of the correspondence we show the HG rate as a function of $R$ for $\lambda = 800$ nm in figure 7, while
the ionization rates are given in figure 4(b). The upper state shows a more prominent harmonic generation peak around \( R = 8 \text{ au} \).

As can be observed from figure 7(b), \( \Gamma_7 \) displays strong peaks around \( R \approx 4 \text{ au} \) in a region for both states, where the ionization rates are not strong. This is particularly true for the lower state. This behaviour, thus, must come from bound-state populations [8]. However, a major peak can be observed in \( \Gamma_7 \) around \( R \approx 8 \text{ au} \) (figure 7(d)) for the upper state, where a strong enhancement of the ionization rate does occur (figure 4(b)). A significant feature around \( R \approx 8 \text{ au} \) is also visible in \( \Gamma_\text{\text{Gamma1}} \) for the upper state (figure 7(e)).

Compared to the \( \lambda = 1064 \text{ nm} \) case figure 7(f) shows a much weakened \( \Gamma_\text{\text{Gamma1}} \) rate. This is a consequence of the molecular cutoff law (15). Naively, one might think that a reduced wavelength \( \lambda \) (increased \( \omega \)) will simply push the cutoff to higher \( R \). However, at \( R > 10 \text{ au} \) the ionization rates drop (cf figure 4). One reason why the dynamics change at large \( R \) is that the upper and lower states become nearly degenerate, i.e., \( \omega \gg \epsilon_{1\sigma} - \epsilon_{1\sigma} \) and the strong-coupling regime [8] is reached.

From the behaviour of the \( n < n_M \) rates \( \Gamma_n \) shown in figures 6, 7 it is obvious that the dipole moment equation (12) is a complicated periodic function of time when \( R \gg 2 \text{ au} \). It is no longer dominated by the fundamental frequency \( \omega \) as is usual for atomic HG spectra.

4. Conclusions

We have presented a non-Hermitian Floquet calculation of HHG for the lowest two \( \text{H}_2^+ \) eigenstates in monochromatic strong laser fields using the length gauge. A pseudospectral representation of the Hamiltonian was applied, and the CAP method was implemented to avoid the calculation of an oscillatory tail in the coupled-channel resonance wave function and to get accurate resonance parameters. In this approach even though we needed to solve a large non-Hermitian matrix problem to get the solution for the TDSE, we avoided problems that are associated with time-stepping algorithms, particularly the accumulation of phase errors.

The results for the HHG rates for the lower state of \( \text{H}_2^+ \) in strong laser fields were compared with previous calculations in the literature [8, 18]. The cutoff positions in the HHG spectra were examined and compared with the classical cutoff formula given in [4]. Good agreement was found for \( I \leq 10^{14} \text{ W cm}^{-2} \) and some deviations at higher intensities for high harmonic orders.

In the region of the molecular plateau, \( n < n_M \), (cf equation (15)), while considering separations \( R = 4–10 \text{ au} \) it is shown that the enhancement of the ionization rate for the lower and upper states at \( R \approx 8 \text{ au} \) causes an enhancement of the harmonic generation rate in the following way: \( \Gamma_7 \) for both states and \( \Gamma_\text{\text{Gamma1}} \) for the lower state of \( \text{H}_2^+ \) in a field of \( I = 10^{14} \text{ W cm}^{-2} \) and \( \lambda = 1064 \text{ nm} \) show peaks; similarly \( \Gamma_7 \) for the upper state of \( \text{H}_2^+ \) in a field of \( I = 10^{14} \text{ W cm}^{-2} \) and \( \lambda = 800 \text{ nm} \) displays peaks at \( R \approx 8 \text{ au} \). The lower bound of this enhanced HG region \( R = 4 \text{ au} \) is characterized by near-resonant coupling \( \omega \approx \epsilon_{1\sigma} - \epsilon_{1\sigma} \). It is bounded at \( R \approx 10 \text{ au} \) by the decrease in ionization rate (cf figure 4). On the other hand, for \( \lambda = 800 \text{ nm} \) \( \Gamma_7 \) dominates the HG spectra around \( R \approx 4 \text{ au} \) for the upper and lower states which cannot be associated with an enhanced ionization rate, but with strong resonant coupling, as explained in [8].

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