ABSTRACT: Raman scattering is a very powerful tool employed to characterize molecular systems. Here we propose a novel theoretical strategy to calculate the Raman cross-section in time domain, by computing the cumulative Raman signal emitted during the molecular evolution in time. Our model is based on a numerical propagation of the vibronic wave function under the effect of a light pulse of arbitrary shape. This approach can therefore tackle a variety of experimental setups. Both resonance and nonresonance Raman scattering can be retrieved, and also the time-dependent fluorescence emission is computed. The model has been applied to porphyrin considering both resonance and nonresonance conditions and varying the incident field duration. Moreover the effect of the vibrational relaxation, which should be taken into account when its time scale is similar to that of the Raman emission, has been included through the stochastic Schroedinger equation approach.

1. INTRODUCTION

One of the techniques mainly employed to investigate the vibronic structure of molecules is Raman spectroscopy,1−5 which has become a very powerful tool useful for the characterization of molecular systems.6 Due to the very short time scale of the process, the Raman signals are very sharp, and thanks also to the richness of information enclosed on the vibronic structure, they can constitute a fingerprint for molecules.7,8 What occurs in Raman scattering is a two-photon process9,10 that involves a first interaction with an incident radiation which makes the molecule explore a virtual state, namely a superposition of excited states, and the emission of a photon with energy that differs from the incident one by a molecular vibrational frequency.11,12 Considering an incident radiation in resonant conditions, the distinction is made between resonance Raman scattering and fluorescence emission that differ in time scale and spectral shape: Raman scattering is the early time emission (which generates spike lines in the spectrum) and the fluorescence is the late time emission (responsible for broad bands).13 In nonresonant conditions the Raman scattering and the fluorescence lay on a different range of the scattered energies and thus the distinction is unequivocal.

Raman scattering is distinguished from Rayleigh scattering, the elastic process, in which the scattered photon has the same energy of the incident one, or in other words the scattered photon brings the molecule back to the initial state. Raman process is related to the inelastic scattering of photons namely the scattered light has a different frequency with respect to the incident light. The bands generated by the Raman scattering are named Stokes when the scattered light has a frequency smaller than the incident light, otherwise they are called anti-Stokes. Differently from other spectroscopic techniques, the quantum mechanical description of the process is not trivial, and a great effort has been employed so far to give insight on the process and to theoretically reproduce the experimental results.

The theoretical methods employed to simulate Raman scattering start from the Kramers, Heisenberg, and Dirac (KHD)14−16 polarizability tensor and can be categorized in time-independent and time-dependent strategies. To the former belong the so-called sum-over-state methods based on an explicit expansion17−20 of the transition dipole moments ideally on all the molecular vibronic states.21−24 On the time-dependent side, the Lee and Heller12,13 theory gave a new...
picture of the Raman process as result of the propagation of a wavepacket on the excited-state potential energy surface. In this framework, the Raman intensity is determined by the half Fourier Transform of the overlap of initial and final states.\textsuperscript{25} Time-dependent strategies have been developed coupling Raman theory to DFT methods\textsuperscript{36–39} or employing Green’s function method to calculate the vibronic states overlap.\textsuperscript{40} On the other hand a real-time propagation can provide for the molecular response in the perturbation theory framework to retrieve the KHD polarizability.\textsuperscript{31,32} Other approaches developed in the time domain, based on the Fourier transform of the polarizability, have also increased the accuracy of results including the Herzberg–Teller vibronic coupling\textsuperscript{33–35} and the Duschinsky effect.

In this work we propose a time-dependent procedure able to give a view of the process that goes on in Raman scattering, based on a semiclassical approach, that is, an approach which combines the classical treatment of the electric field with a quantum description of the system through its vibronic wave function.\textsuperscript{36–42} On the basis of the time-dependent Raman scattering theory proposed by Lee and Heller,\textsuperscript{12} a real-time propagation of the system wave function has been coupled with an integration in time\textsuperscript{11} that results in the computation of the time-dependent Raman cross-section through the calculation of the second order wave function. Compared with the original treatment of Lee and Heller,\textsuperscript{12} and subsequent TD approaches building on that, the main novelties are (i) we are using a numerically defined excitation pulse shape, without assuming a monochromatic continuous wave excitation; (ii) we are not only calculating the total Raman scattered intensity, but we provide the time-dependent cumulative signal; (iii) we include relaxation effects (and potentially also decoherence ones) via the Stochastic Schroedinger equation technique. The latter point allows treating Raman and fluorescence on the same footing, both resulting (on different time scales) from the same simulation.

Employing a generic electric field shape, which could be even defined by points, makes the simulation closer to pulsed experimental setups.

The present approach, which is not only time-dependent but also time-resolved, allows having information on the time scale of the process and also on the way Raman spectra are built in time in comparison with the time scale of the incident radiation.

Within this procedure, phenomena of decoherence and relaxation can be included in the dynamics through stochastic methodologies to consider the interaction with a surrounding environment in an effective way.\textsuperscript{43–45} In this work, the stochastic Schroedinger equation has been employed in order to investigate the role of the vibronic relaxation on the Raman process. Vibronic relaxation becomes relevant in the process since it occurs within a picosecond time scale that is also the time scale of the dynamics here reported.

The method here proposed can be applied without further approximations both in resonance and nonresonance conditions between the pulse frequency and the electronic transition. It is worth mentioning that with this strategy both the Raman signal and the fluorescence emission are computed since both contributions, in the present semclassical formulation, come from the wave function at the second order with respect to the electric field even if the process is different. The scattered photon emitted in the Raman process causes the decay from a virtual state with energy determined by the pulse frequency to a vibronic state of the ground state, while the fluorescence emission needs, at first, to create a population on the excited state and so the emitted photon has energy close to the electronic transition. On this basis, the two phenomena can be distinguished when the two scattering signals are wide apart as in nonresonant conditions. Strategies to computationally tackle time-dependent frequency resolved spontaneous emission have been discussed in the past,\textsuperscript{46–48} as well as for time-dependent stimulated Raman;\textsuperscript{49,50} computational investigations of the time-development of spontaneous Raman spectra and its interplay with fluorescence received less attention.

This work is a prelude of the more challenging goal of computing the time-dependent spontaneous Raman scattering of molecules close to plasmonic nanoparticles, in order to simulate and predict surface-enhanced Raman scattering (SERS)\textsuperscript{31–33} experiments in a time-dependent perspective, combining the real-time dynamics of a molecule vibronic wave function with the propagation of the plasmonic nanostructures dielectric polarization.

The paper is organized as follows: In section 2 we summarized the theoretical methods divided into the formulation of the time-dependent Raman scattering cross-section, the model for the system vibronic wave function employed, the numerical propagation procedure, and the computational details of the calculations performed; the results obtained with our model and the related discussion are presented in section 3 and final remarks are reported in section 4.

2. METHODS

2.1. Time-Dependent Raman Scattering Formulation. In this section the main aspects of the theory we employed to describe Raman scattering are highlighted. The time-dependent picture of Raman scattering can be retrieved in the framework of perturbation theory through the two-photon formula. The first order correction to the system wave function allows a description of a single-photon process, such as absorption or emission. It is the result of the interaction between the time evolution of the unperturbed (initial) wave function and the electric field that induces a dynamics on an excited potential energy surface, which in atomic units reads as

\[ i\frac{\partial}{\partial t} \Psi(1)(t) = i \int_{-\infty}^{t} dt' e^{-i\hat{H}(t-t')} \hat{\mu} \cdot \vec{e}(t') e^{-i\hat{H}'t'} \Psi(0)(-\infty) \]

\[ \Psi(0)(-\infty) \] is the initial unperturbed wave function, considered here to be the ground state, with energy \( E_{0} \), \( \hat{\mu} \) is the transition dipole moment operator, \( \vec{e}(t') \) is the incident electric field interacting with the system at time \( t' \), and \( \hat{H} \) is nuclear plus electronic Hamiltonian in the Born–Oppenheimer (BO) approximation. Note that we are also departing from the original Lee and Heller treatment (see also ref 11) since there the propagation is done on electronic wave functions and \( \hat{H} \) refers to the electronic Hamiltonian of the proper potential energy surface. Instead, here the vibronic Hamiltonian (i.e., the total Hamiltonian written on the basis of the vibronic wave functions in the BO approximation) enters in the propagation of the wave function. The BO approximation limits our approach to molecules where the vertical excitation region of the lowest excited state is sufficiently far from avoided crossings, where it would break

\[ \text{Article} 

\begin{align*} 
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\text{https://doi.org/10.1021/acs.jpca.2c05245} 
\end{align*}
To account for spontaneous emission from the virtual state that strictly speaking would require a quantum electrodynamics (QED) treatment, we shall use the same technique proposed by Lee and Heller. In practice, a second classical field $\tilde{e}_S$ is considered, that is monochromatic at the frequency $\omega_0$, whose properties are then chosen to reproduce QED results. At the second order in the mixed $\tilde{e}_S$ and $\tilde{e}_I$ perturbation, neglecting the antiresonant term, one gets

$$|\Psi^{(2)}(t, \omega_0)\rangle = -\frac{1}{2} \int^t -\infty dt' \int^t -\infty dt'' e^{-i(\theta(t''-t')/\mu) - \tilde{e}_S e^{i\omega_0 t'}} e^{i\theta(t'-t'')/\mu} \tilde{e}_S e^{i\omega_0 t''} \psi^{(1)}(t') \psi^{(1)}(t'') \rangle$$

(2)

which differs from the original Lee and Heller theory for the missing antiresonant term. In eq 2 the system wave function, after a dynamics guided by the Hamiltonian $\hat{H}_I$, interacts with $\tilde{e}_S(t') = \tilde{e}_S e^{i\omega_0 t'}$ that brings the system at a low energy level, which then evolves under the vibronic Hamiltonian. The second electric field is not included explicitly in the dynamics since it is convenient to calculate its effect a posteriori. Neglecting the antiresonant term (that in which the field $\tilde{e}_S$ interacts with the molecule before $\tilde{e}_I$) is an approximation that limits the validity of the present treatment to resonant and preresonant conditions, while it would be unsuitable (in terms of quantitative accuracy for the Raman intensities) for long wavelength excitations. Nothing fundamental prevents the inclusion of this term in our approach, but to be calculated for a generic $\tilde{e}_S(t)$ it would require several numerical propagations, starting from all the possible excited vibronic states, representing an increase of the computational burden not needed in the present work.

Equation 2 can be expressed as a function of the first order wave function as

$$|\Psi^{(2)}(t, \omega_0)\rangle = \frac{i}{2} \int^t -\infty dt' e^{-i(\theta(t''-t')/\mu)} \tilde{e}_S e^{i\omega_0 t'} |\psi^{(1)}(t')\rangle$$

(3)

where the dynamics of $|\psi^{(1)}(t')\rangle$, expressed by eq 1, can be obtained numerically through a real-time propagation of the initial wave function interacting with any shape of the incident electric field.

Practically, the wave function is expressed as a linear combination of vibronic eigenstates $|J\rangle$

$$|\Psi^{(2)}(t, \omega_0)\rangle = \sum_j C_j^{(2)}(t, \omega_0) |J\rangle$$

(4)

weighted by the time-dependent coefficients at the second perturbative order. Multiplying left and right of eq 4 by the eigenstate $\langle N |$ and substituting the definition of $|\Psi^{(2)}(t, \omega_0)\rangle$ given in eq 3, we obtain the time-dependent coefficients of the state $N$

$$\langle N | \Psi^{(2)}(t, \omega_0) \rangle = \frac{i}{2} \int^t -\infty dt' \langle N | e^{-i(\theta(t''-t')/\mu)} \tilde{e}_S e^{i\omega_0 t'} \psi^{(1)}(t') \psi^{(1)}(t'') \rangle$$

(5)

The first order wave function can be expressed as well as an expansion on vibronic states $|J\rangle$ weighted by the first order time-dependent coefficients, $|\psi^{(1)}(t')\rangle = \sum_j C_j^{(1)}(t') |J\rangle$, which can be computed through the dynamics of the system wave function, initially in the GS, interacting with an incident radiation (practical details later). Equation 5 can be rewritten as

$$C_j^{(2)}(t, \omega_0) = \frac{i\epsilon_j e^{-i\omega_0 t}}{2} \int^t -\infty dt' e^{i\omega_0 t'} \sum_j C_j^{(1)}(t') \langle N | \hat{\mu}_j | J \rangle$$

(6)

where $\epsilon_j = |e_j|$ is the monochromatic scattering field amplitude and $\hat{\mu}_j = (\hat{\mu}_j / \epsilon_j / \epsilon_s$ is the transition dipole moment operator along the direction of $\tilde{e}_S$.

The second order wave function coefficients of a state $N$ at the limit for $t$ that tends to infinity is exactly the Inverse Fourier Transform (IFT) with respect to the scattered frequency of the first order coefficients, that is directly related to the Raman scattering intensity. From another perspective, the IFT of the first order coefficients as in eq 6 considering a finite time $t$ collects the Raman scattering signal emitted until that moment, giving information on the ongoing Raman process. When the time dependence on the coefficients (i.e., without setting the limit to infinite) is kept, the Raman cross-section can be calculated as a function of time.

To get to this point the square modulus of the coefficients are computed as

$$|C_j^{(2)}(t, \omega_0)|^2 = \frac{\epsilon_j^2}{4} \int^t -\infty dt' e^{i\omega_0 t'} \sum_j C_j^{(1)}(t') \langle N | \hat{\mu}_j | J \rangle|^2$$

(7)

enclosing in the term $\int_L^{N}(t, \omega_0)|^2$ the integral over $t'$. Numerically, the integral is calculated by a fast Fourier transform including in the integrand a $\Theta(t - t')$ Heaviside function. The square modulus of the coefficients is the population of the molecular state $N$ created by the scattered radiation with frequency $\omega_0$ from upper vibronic states. The time dependence of the population gives information on the total photons emitted until the observation time $t$. When a specific polarization direction for the incident field ($\eta$) is chosen and the scattered signal produced by the scattered field oriented along $\lambda$ is selected, all the components $z_j^{N}(t, \omega_0)$ are calculated with indices $\lambda$, $\eta$ that run over the Cartesian directions. When the $z_j^{N}(t, \omega_0)$ terms for all the combinations of $\lambda$ and $\eta$ are collected, they can be combined in the same way as the polarizability tensor to compute the Raman intensity with a precise polarization of the incident and scattered radiation, and an isotropic average of the molecular orientation is assumed. In this situation, eq 7 can be rewritten as

$$\langle |C_j^{(2)}(t, \omega_0)|^2 \rangle_{\text{setup}} = \frac{\epsilon_j^2}{4} \int_{\text{setup}}^{N^{(2)}(t, \omega_0)|^2}$$

(8)

indicating with $\langle |C_j^{(2)}(t, \omega_0)|^2 \rangle_{\text{setup}}$ the population in the state $N$ after isotropic orientational average for a specific illumination-detection setup, and with $z_j^{N}(t, \omega_0)$ the combination of $z_j^{N}(t, \omega_0)$ terms proper for that setup. For example, for incident radiation with linear polarization
perpendicular to the detected scattered radiation and any
polarization for the scattered radiation, the terms combine as

$$z_{N,\text{setup}}^2(t, \omega_S) = \frac{45\alpha_N^2 + 7\gamma_N^2 + 5\delta_N^2}{45}$$

(9)

with

$$\alpha_N = \frac{z_{zz}^N + z_{yy}^N + z_{xx}^N}{3}$$

$$\gamma_N^2 = \frac{1}{2} \left[ (z_{xx}^N - z_{yy}^N)^2 + (z_{xx}^N - z_{zz}^N)^2 + (z_{yy}^N - z_{zz}^N)^2 \right]$$

$$\delta_N^2 = \frac{3}{4} \left[ (z_{xx}^N - z_{yy}^N)^2 + (z_{xx}^N - z_{zz}^N)^2 + (z_{yy}^N - z_{zz}^N)^2 \right.$$}

(10)

In order to compute the Raman cross-section as the ratio
between the scattered energy and the incident radiation
fluence we have to define these two quantities based on the
theory developed so far. The whole population of the states
involved in the scattering process is the sum of the square
modulus of the second order coefficients over all the \(N\)
vibronic states. The scattered energy is equal to the number of
photons emitted at frequency \(\omega_S\) multiplied by the emitted
energy \(\omega_S\) (in atomic units), where the number of scattered
photons is equal to the product between the second order
coefficients, representative of the states population achieved by
a specific scattered field, and the number of scattered fields
with frequency \(\omega_S\) and amplitude \(c_S\). It follows that the
scattered energy over a small \(d\omega_S\) frequency interval for a given
illumination detection setup \(dE_{\text{setup}}\) can be written as a function
of the second order coefficients as

$$dE_{\text{setup}}(t, \omega_S) = \sum_N \langle C_N^{(2)}(t, \omega_S) \rangle_{\text{setup}} \rho(\omega_S) \omega_S \, d\omega_S$$

(11)

where \(\rho(\omega_S)\) is the density of states of the scattered field
with frequency \(\omega_S\) and amplitude \(c_S\). As mentioned, the number of emitted photons can be easily computed from eq 11 by
dividing for the photon energy \(\omega_S\). Substituting the square of
the field amplitude with \(c_S^2 = \frac{8\omega_S}{\nu^2}\), the density of states of the
scattered field with \(\rho(\omega_S) = \frac{\nu^2}{(2\pi)^3} d\Omega\), and the second order
coefficients as in eq 8, we end up with the scattered energy per
unit of frequency and per unit of solid angle

$$\frac{dE_{\text{setup}}(t, \omega_S)}{d\omega_S d\Omega} = \frac{\alpha_N^4}{4\pi^2 c} \sum_N z_{N,\text{setup}}^2(t, \omega_S)$$

(12)

On the other hand, the fluence of the incident field depends on
its integral over time as

$$\mathcal{F}_I = \frac{c}{4\pi} \int_{-\infty}^{\infty} dt |\mathcal{E}_I(t)|^2$$

(13)

Any field shape can be used in principle, even a known-by-
points electric field. In this case we considered a Gaussian
enveloped sinusoidal field with equation

$$\mathcal{E}_I(t) = \mathcal{E}_{I,0} \sin(\omega_0 t) e^{-(t-t_0)^2/(2\sigma^2)}$$

(14)

centered in \(t_0\) with amplitude \(\mathcal{E}_{I,0}\), width \(\sigma\) and frequency \(\omega_0\).

Integrating the incident electric field over time leads to the
fluence of the incident field

$$\mathcal{F}_I = \frac{c \sigma}{8\sqrt{\pi}} |\mathcal{E}_{I,0}|^2$$

(15)

Dividing the scattered energy of eq 12 by the incident field
fluence in eq 15, the cumulative time-dependent cross-section
per unit of solid angle and per unit of scattered frequency is
computed as a sum over the contribution given by each\vibronic state \(N\):

$$\sigma_{\text{setup}}^{\text{flu}}(t, \omega_S) = \frac{2\omega_S^2}{\pi \sigma |\mathcal{E}_{I,0}|^2} \sum_N z_{N,\text{setup}}^2(t, \omega_S)$$

(16)

With eq 16 we can not only recollect the overall Raman cross-
section for a given illumination-detection setup, that can be
also computed through strategies based on the transition
polarizability tensor, but also the Raman signal as a function of
time. In other words, this approach lets one know how progressively the Raman signal is accumulated and which is the time scale of the process.

2.2. Vibronic Wave Function. To obtain the Raman signal it is worthwhile to consider a proper wave function in the dynamics, dressing the electronic states with vibrational levels. The wave function, expanded on vibronic states, is considered within the BO approximation to separate electronic and nuclear motions. In this framework each vibronic eigenstate \(|\psi\rangle\) used in the previous section can be expressed as a product between an electronic \(|\psi_e\rangle\) and a vibrational \(|\psi_v\rangle\) wave function \(|\psi\rangle = |\psi_e\rangle|\psi_v\rangle\). Based on this approximation, the transition dipole moment between two vibronic states is expressed as

$$\mu_{ij} = \langle \psi_i | \langle \psi_f | \mu_e | \psi_f \rangle | \psi_i \rangle = \langle \psi_i | \hat{\mu}_e | \psi_i \rangle$$

(17)

with \(|\psi_i\rangle\) and \(|\psi_f\rangle\) being the initial and final vibrational levels which belong respectively to the ground \((|\psi_i\rangle)\) and the excited \((|\psi_f\rangle)\) electronic states. The vibrational levels are expanded
within the normal mode approximation; e.g. anharmonic terms are neglected. Transition dipole moments between vibronic states that belong to the same electronic state are also neglected.

At this level the electronic and vibrational parts are
decoupled so the Herzberg–Teller (HT) coupling has been included to recover for the transition dipole moment dependence on the normal mode coordinates, even if in an approximated way. Introducing the HT coupling allows inclusion of the dependence on nuclear coordinates in terms of a perturbative expansion of the electronic transition dipole due to the presence of a vibrational manifold. Therefore, the electronic transition dipole can be written as a Taylor expansion in the nuclear displacements around the nuclear equilibrium configuration \(Q_0^e\)

$$\hat{\mu}_e = (\hat{\mu}_e)^0 + \sum_k \left( \frac{\partial \hat{\mu}_e}{\partial Q_k} \right)^0 Q_k + o(Q^3)$$

(18)

where \(k\) runs over the normal mode coordinates. The first term of eq 18 is the electronic transition dipole moment calculated at the nuclear equilibrium position, while in the second term appears the transition dipole moment derivative with respect to
the normal mode coordinates evaluated at the ground state equilibrium geometry. When eq 18 is included in eq 17, the new expression for the transition dipole moment between two generic vibronic states reads as
\[ \tilde{\mu}_j = \langle \chi^i \mid \chi^i \rangle \langle \mu \rangle_0 + \sum_k \langle \chi^i \mid Q_k \chi^i \rangle \left( \frac{\partial \langle \mu \rangle_k}{\partial \tilde{Q}_k} \right)_0 \] (19)

The first term of eq 19 includes the Franck–Condon integral calculated between the final and initial vibronic states when the electric dipole moment is considered at the equilibrium normal mode coordinates. The second term accounts for the Herzberg–Teller contribution including a first order correction of the electric dipole moment with respect to the normal mode coordinates.

2.3. Numerical Propagation. As mentioned above, within our approach the dynamics of the first order wave function is computed numerically. Practically the coefficients of the vibronic expansion are propagated through a second-order Euler algorithm as
\[ C_N(t + dt) = C_N(t - dt) - 2i dt (H_0 - \tilde{\mu} \cdot \tilde{\xi}(t)) C_N(t) \] (20)
in which dt is the time step chosen for the propagation, \( H_0 \) is the time independent Hamiltonian that returns the energy of the state \( N \) while \( \tilde{\mu} \cdot \tilde{\xi}(t) \) accounts for the interaction with the incident electric field. The incident radiation can have in principle any profile since it is numerically defined. In this work we consider a Gaussian enveloped sinusoidal equation for the incident field as said before.

The greater part of the calculations has been performed considering the molecule as a closed quantum system. However, some others have been performed including the vibrational relaxation from the upper vibrionic states to the lower level of the same electronic state. To this purpose the system wave function has been propagated through the Stochastic Schroedinger Equation \((\text{SSE})\) including non-radiative decay rate from the upper vibronic states toward the lower level of the electronic excited state. The SSE in a Markovian regime reads
\[ \frac{d}{dt} \Psi(1)(t) = -i\mathcal{H}(t)\Psi(1)(t) - \frac{1}{2} \sum_q \tilde{S}_q(t) \tilde{S}_q(t) \Psi(1)(t) \] (21)
where \( q \) runs over the number \( M \) of decay channels of the system and \( I_q(t) \) is a white noise function associated with the Markov approximation needed to include the fluctuation induced by the environment. The first term in the r.h.s. of eq 21 represents the deterministic term related to the energy of the system and the interaction with the incident field, while the second and third terms are responsible for the decay processes guided by operators \( \tilde{S}_q(t) \). According to previous works, relaxation has been included by the operator
\[ S_q^{\text{rel}} = \sqrt{\Gamma_q | \phi_q \rangle \langle \phi_q |} \chi^i | \phi_q \tan{\theta} \] (22)
which is responsible for the decay of the vibronic state \( | \phi_q \rangle \chi^i \) population on the state \( | \phi_q \rangle \chi^i \) where \( q \) is the vibrational level and \( \epsilon \) the electronic state. The decay rate \( \Gamma_q(t) \) can be taken either as phenomenological parameters (as in the present case, see Computational Details) or estimated by ab initio simulations. Practically the coefficients dynamics are computed through the second-order Euler algorithm as in eq 20 while the stochastic terms of SSE are included by means of a quantum jumps algorithm. This algorithm is based on discontinuous trajectories. At each time step during the dynamics, the probability of the jump to occur associated with the decay operator \( \tilde{S}_q(t) \) and given by \(| \Psi(1)(t) \rangle \tilde{S}_q \langle \Psi(1)(t) \rangle | dt \) is estimated. Whether the jump actually occurs or not is decided by a Monte Carlo algorithm. When the jump happens, the wave function \( | \Psi(1)(t) \rangle \) is replaced by \( \tilde{S}_q | \Psi(1)(t) \rangle \) and normalized; on the contrary it is only normalized. The density matrix evolution is obtained by averaging over several of these trajectories. For further details see ref 43.

The coefficients dynamics of each trajectory has been employed to compute the second order coefficients. Varying the polarization of the incident field and the scattered radiation, we computed all the terms \( z_{ej}^{(i)}(t, \omega_b) \) needed to calculate, for a given illumination-detection setup and for each trajectory, the time-dependent Raman cross-section \( \sigma^{\text{setup, traj}}(t, \omega_b) \) and finally the average on all the realizations has been taken as
\[ \sigma^{\text{setup}}(t, \omega_b) = \frac{1}{N_{\text{traj}}} \sum_{\text{traj}} \sigma^{\text{setup, traj}}(t, \omega_b) \] (23)
The Raman cross-section is a measurement of the amount of population generated in each final vibronic state by scattered fields with different frequencies \( \omega_b \). The population computed from a single trajectory is meaningless in itself but when the average over a number of realizations is taken, the SSE converges to the Lindbland master equation results.

2.4. Computational Details. The approach presented in the previous sections has been applied to porphyrin, as a test case. The quantum chemical calculations have been performed through DFT methods at the B3LYP/6-31G level of theory using Gaussian. The ground state and first excited state were optimized, and then the vibrational calculation was performed on both potential energy surfaces. To obtain a proper wave function for the system, the two electronic states have been dressed with 109 vibrational states each (the ground vibronic state and the first vibronic level for each normal mode). We computed the FC and HT integrals of the molecule between all the vibronic states belonging to different electronic states through FCclasses code. The vibronic analysis has been carried out within the adiabatic Hessian model which takes into account the Duschinsky mixing and the frequency changes between ground and excited state. The FC and HT integrals have been employed to compute the transition dipole moments as in eq 19. The system wave function is approximated by including only one excited electronic state, thus when in nonresonance conditions there could be significant differences on the spectra shape and intensity compared with the experimental results. However, the focus of this work is on the time scale of the process and on the interplay with fluorescence.

The dynamics of the first order system wave function has been computed through the WaveT code. During the propagation the system interacts with an incident electric field shaped as a Gaussian enveloped sinusoidal signal, as
mentioned above. Three different simulations for each field setup have been computed varying the incident field direction along \( x, y, \) and \( z \) axes in order to have all the information needed for the calculation of the term \( z_{N,\text{setup}}^{2}\left(t, \theta_{k}\right) \) as in eq 9.

The dynamics length is 24 ps in almost all the calculations and the time step considered is 0.024 fs. When it is not differently specified, the field has a Gaussian enveloped sinusoidal shape (see sec. 2.3) with \( \sigma = 256 \) fs, \( \epsilon_{10} = 5.14 \times 10^{5} \) V/m, centered at \( t_{0} = 968 \) fs. In nonresonant conditions, the excitation energy is \( \alpha_{0} = 2.04 \) eV, which is 0.24 eV less than the vertical transition frequency \( S_{1} \leftarrow S_{0} \). After the calculation of the coefficients dynamics, the cumulative Raman cross-section was computed at different times, through eq 16. The convergence of the spectrum can be checked as explained in ref 21.

In the simulation with the vibrational relaxation, 50 repetitions of the dynamics were computed with vibrational relaxation equal to 1.8 ps.\(^70\) Convergence with respect to the number of employed trajectories when including vibrational relaxation in resonance conditions has been confirmed comparing the Raman spectra at 2.4 ps computed with either 50 or 100 trajectories. The result is reported in Figure S1 of the Supporting Information.

### 3. RESULTS AND DISCUSSION

The procedure developed to compute Raman scattering cross-section has been applied to porphyrin, laying in the \( xy \) plane as Figure 1 shows. The calculation of the cumulative time-dependent Raman cross-section has been performed by employing two different field frequencies to study the nonresonance and resonance conditions with respect to the electronic excitation energy. Moreover also the field duration and the presence of the vibrational relaxation have been exploited.

To validate the numerical strategy, we compared porphyrin Raman spectrum computed through our TD method in nonresonant conditions with the spectrum computed with a time independent strategy, by employing the FCclasses code.\(^21,68\) The results, reported in Figure S2, show a good agreement, therefore confirming the correctness of our strategy.

Moreover a comparison with the experimental Raman spectrum of porphyrin \(^1\) is reported in Figure S3. Apart from an overall rescaling of the frequencies, which is expected for harmonic spectra,\(^21,73\) the agreement is good. The remaining discrepancies (beside the frequency rescaling), may be due to the different environments (the experiment is conducted in CH\(_4\)Cl\(_2\), while the calculation is performed in vacuum), to anharmonic effects beside the rescaling and to inherent limitation of the DFT xc functional employed.

#### 3.1. Nonresonance Conditions

In all the calculations reported in this section, the first order coefficients of the system wave function have been computed when the system is initially in the ground electronic state and then interacts with an incident electric field nonresonant with the electronic transition as explained before. After the computation of the coefficients dynamics varying the incident field direction along \( x, y, \) and \( z \) axes, three different scattering directions (along \( x, y, \) and \( z \) axes) have been considered for each incident field direction to compute the time-dependent Raman cross-section in atomic units as in eq 16. Since the electronic transition dipole moment \( S_{1} \leftarrow S_{0} \) has the major contribution along the \( x \) and \( y \) axes, the terms with incident and/or scattered field along the \( x \) and \( y \) axes give the largest contribution to the Raman cross-section.

**Neglecting Vibrational Relaxation.** The Raman spectrum as a function of time has been reported in Figure 2a for six different time delays with respect to the beginning of dynamics.

The results show how the Raman intensity is built during and after the interaction with the incident field. Before 484 fs, far from the maximum intensity of the electric field, the Raman intensity is negligible, but before reaching the center of the field at 968 fs, the Raman signal has already generated due to the short time taken by the process. After 1451 fs from the beginning of the dynamics quite all the Raman intensity is recovered, even if the pulse has not switched off completely. The last spectrum here reported, after 2.4 ps from the beginning of the dynamics, when the pulse is approximately null, is totally equal to the final spectrum computed at 24 ps: all the Raman contribution to emission is achieved in the first hundreds femtoseconds from the interaction with the incident field.

These observations are confirmed by the second column of Table 1 which reports the integral of the peak at 738 cm\(^{-1}\) as a function of time, representing the amount of Raman signal collected until the observation time from the beginning of the dynamics without the dependence on the frequency spread signal. The integrals highlight the fast increasing intensity in the first few hundreds of femtoseconds around the maximum field intensity followed by a constant value, a signal that the maximum emitted intensity in this region has been achieved. One possible way to interpret the results is that Raman scattering persists as long as the wave function coefficients have memory of the pulse frequency or, from a more physical viewpoint, until the vibronic excited states are transiently perturbed by the presence of the incident field. After the end of the interaction with the field no more Raman signal is collected since the process, very quickly, is exhausted.

**Increasing Field Time Duration.** Another calculation has been performed increasing the field time duration by modifying the width of the pulse with \( \sigma = 496 \) fs and moving the center of the pulse at 2419 fs. In this case the pulse is larger so it is even closer to a monochromatic pulse than the previous one. The Raman spectrum at different time delays from the beginning of the dynamics is reported in Figure 2b. The vibronic states that participate in the Raman spectrum are the same as in the previous case, while the time scale of the signal generation is longer and comparable to the width of the incident pulse employed here. The signal peaks are narrower and well separated due to the incident field shape, closer to a

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**Figure 1.** Structure of porphyrin molecule and orientation employed for the calculations.
monochromatic radiation than in the previous one. As a consequence, these results highlight the dependence of the spectrum line shape on the electric field shape and, in particular in this case, on the duration of the pulse. At last the intensity of the Raman signal in time can be compared with the previous one by removing the frequency dependence through the integration of the peak at 738 cm$^{-1}$ as reported in Figure 2c and in the fourth column of Table 1. These results show that the amount of scattered radiation is quite the same at the end of the dynamics although the time of the process is longer in the case of a longer incident pulse. On the other hand in the fluorescence emission region (around the transition $S_1 \leftarrow S_0$ energy) the emitted signal has a lower intensity when a longer lasting field is employed, as represented in Figure 2d, because the smaller frequency amplitude of the electric field can excite (nontransiently, as needed for fluorescence emission) less amount of the excited states population.

Including Vibrational Relaxation. Another calculation of porphyrin Raman cross-section has been carried out including vibrational relaxation. To this point, field features have been kept as in the first calculation reported in this section (nonresonant electric field with $\sigma = 256$ fs), and the vibrational relaxation has been included from upper vibronic states to the lowest level of the correspondent electronic state with the phenomenological lifetime of 1.8 ps. 70 trajectories have been computed through SSE equation using in each case three incident field directions (along $x, y, z$ axes) and the dynamics of each simulation is 4.8 ps long. For each trajectory the time-dependent Raman cross-section has been computed, and then they have been averaged to get the mean Raman cross-section. The time evolution of the Raman cross-section including vibrational relaxation, reported in Figure 3, looks very close to the results in which it is neglected (Figure 2a) as expected since the time scale of the process seems faster than the

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**Figure 2.** Time dependent Raman cross-section in atomic units at six different time delays with respect to the beginning of dynamics, with incident pulse in nonresonant conditions with $\sigma = 256$ fs (a) and with $\sigma = 496$ fs (b). (c) Time dependent Raman cross-section integral of the peak at 738 cm$^{-1}$ in the case of field with $\sigma = 256$ fs and with $\sigma = 496$ fs. (d) Scattering signal at the end of dynamics in the resonance region when the system interacts with a nonresonant field in which $\sigma$ is 256 fs or 496 fs. The inset is a magnification of the emitted signal generated with an incident field that has $\sigma = 496$ fs.

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**Table 1. Cross-Section Integrals of the Peak at 738 cm$^{-1}$ as a Function of Time in Three Cases: with $\sigma = 256$ fs (from Figure 2a), with $\sigma = 496$ fs (from Figure 2b), and with $\sigma = 256$ fs Including also Vibrational Relaxation (from Figure 3)**

| Time (fs) | Cross-section integral ($\sigma = 256$ fs) | Cross-section integral ($\sigma = 496$ fs) | Cross-section integral (vib. relaxation) |
|-----------|------------------------------------------|------------------------------------------|------------------------------------------|
| 484       | 6.7250 × 10^{-16}                        | 4.3633 × 10^{-16}                        | 6.7236 × 10^{-16}                        |
| 726       | 1.7272 × 10^{-14}                        | 1.6738 × 10^{-14}                        | 1.7277 × 10^{-14}                        |
| 968       | 9.9813 × 10^{-13}                        | 4.9304 × 10^{-14}                        | 9.9807 × 10^{-14}                        |
| 1209      | 1.8585 × 10^{-13}                        | 1.0129 × 10^{-13}                        | 1.8582 × 10^{-13}                        |
| 1451      | 2.0457 × 10^{-13}                        | 1.5386 × 10^{-13}                        | 2.0456 × 10^{-13}                        |
| 1936      | 2.0538 × 10^{-13}                        | 1.8756 × 10^{-13}                        | 2.0536 × 10^{-13}                        |
| 2428      | 2.0538 × 10^{-13}                        | 2.0538 × 10^{-13}                        | 2.0536 × 10^{-13}                        |
| 24289     | 2.0538 × 10^{-13}                        | 2.0538 × 10^{-13}                        | 2.0536 × 10^{-13}                        |
vibrational decay. This result is confirmed by the time-dependent integral of the peak at 738 cm\(^{-1}\), the values of which, reported in column six of Table 1, totally resemble those of the deterministic dynamics (without vibrational relaxation) reported in the second column of the same table. As a conclusion, the introduction of vibrational relaxation is not needed, at least in nonresonant conditions, due to the different time scale of the processes.

3.2. Resonance Conditions. The results reported in this section are related to calculations carried out with the incident field frequency that matches the porphyrin vertical excitation energy (\(\omega_I = 2.28\) eV), while the maximum amplitude is equal to \(5.14 \times 10^5\) V/m, the pulse width has been varied as in the calculations of section 3.1. The time-dependent scattering cross-section has been computed on the basis of the system dynamics when the incident field is polarized alternately along \(x, y,\) and \(z\) axes. For these calculations in resonant conditions, we refer to the results as to scattering cross-sections, rather than to Raman cross-sections given that the Raman scattering and the fluorescence emission lay in the same spectral range. Thus, in principle the two contributions may be mixed and the scattering cross-section is no more the result of the pure Raman contribution.

Neglecting vibrational relaxation. In the first calculation the width of the pulse is \(\sigma = 256\) fs and it is centered at \(t_0 = 968\) fs, as in the first calculation in nonresonance conditions. In Figure 4a the time-dependent scattering cross-section accumulated at six different times from the beginning of dynamics has been reported. Differently than in nonresonance conditions, in this case the scattering cross-section does not achieve its maximum after 1451 fs but at 2419 fs the intensity is further increasing, as an effect of the fluorescence emission. Moreover the vertical transition matches a few vibronic states in the excited state that give the largest contribution to the scattering spectrum. Also the speed at which the peaks’ intensity increases is different, and it is larger for the peak at 156 cm\(^{-1}\), in which the fluorescence emission prevails over the Raman scattering signal, as also Figure 4b shows in terms of the integral of the scattered signals at 156 cm\(^{-1}\) and 738 cm\(^{-1}\).

Increasing Field Time Duration. As it has been done in nonresonant conditions, we tested the influence of a longer pulse on the time-dependent scattering cross-section in resonance conditions, by using a larger width of the pulse (\(\sigma\)
The cumulative scattering cross-section for six different times, reported in Figure 5a, shows thinner peaks than the previous simulation and a longer time scale as expected. Moreover the peak at 156 cm$^{-1}$ is enhanced also in this case.

Figure 5. (a) Time-dependent scattering cross-section in atomic units at six different time delays with respect to the beginning of dynamics, with incident pulse in resonance conditions and $\sigma = 496$ fs. The inset shows a magnification of the part of the spectrum with lower intensity. (b) Time-dependent scattering cross-section integral of the peak at 738 cm$^{-1}$ in the case of field with $\sigma = 256$ fs and with $\sigma = 496$ fs. (c) Scattering signal at the end of dynamics in the resonance region when the system interacts with a resonant field which $\sigma$ is 256 fs or 496 fs.

= 496 fs). The cumulative scattering cross-section for six different times, reported in Figure 5a, shows thinner peaks than the previous simulation and a longer time scale as expected. Moreover the peak at 156 cm$^{-1}$ is enhanced also in this case.

The comparison of these results with those obtained with a shorter pulse in terms of the integral of the peak at 738 cm$^{-1}$ are shown in Figure 5b. In this case, differently than in the nonresonant case, the duration of the pulse has an effect on the...
total scattered signal since the cross-section increases with the longer pulse. This is due to the sum of the two processes that are going on: on one hand the pure Raman scattering, that should remain constant with the pulse duration as in nonresonant conditions, and the fluorescence intensity that strongly depends on the amount of population created on the excited state.

In Figure 5c the convoluted scattering cross-section at 24 ps has been reported for the two simulations with different field widths. In this case the trend as a function of the field duration is opposite with respect to the one in nonresonance conditions since, as mentioned before, the longer is the resonant field the more intense is the scattered signal in the emission region in which both the Raman and fluorescence contributions are mixed. Moreover, on the basis of the results, here the fluorescence contribution prevails.

Practically, in resonant conditions, a relevant population on vibronic states that belong to the excited electronic state, is generated, and it persists even after the incident field is switched off giving a non-negligible contribution to the fluorescence emission. Due to the longer time scale of the process, the fluorescence emission at 24 ps is not exhausted as the Raman scattering which is a faster process as it does not need to create a population on the excited state to occur, but it is the result of the quantum superposition of vibronic states that belong to two different electronic states. By computing a longer dynamics for the first order coefficients, all the emission intensity can be retrieved, but that is beyond the aim of this work.

Including vibrational relaxation. Finally, in resonant conditions, a simulation has been carried out including the vibrational relaxation from upper vibronic states to the lower level of the corresponding electronic state. To this point 50 trajectories have been computed for a dynamics 4.8 ps long, considering three directions of the electric field along x, y, and z axes. After computing the time-dependent cross-section for each trajectory, the average of them has been taken. The resulted scattering cross-section, reported in Figure 6, shows a profile close to the one in absence of vibrational relaxation. However, the maximum intensity is lower. This can be better appreciated by the integral of the peak at 738 cm⁻¹ (Figure 7a) which shows a flat trend after the maximum emission has been achieved while without including relaxation the emission is still increasing. Moreover the convoluted emission spectrum at 4.8 ps shows a lower intensity when the vibrational relaxation is included, as Figure 7b shows, since the population of the vibronic state with frequency equal to 156 cm⁻¹ (which gives the largest contribution to the scattering spectrum) has decayed.

4. CONCLUSION

A computational procedure to calculate Raman scattering cross-section as a function of time has been presented, based on the first order coefficients of the wave function computed after the interaction with an incident electric field that can have any possible shape. The main advantage of this approach is to give a time-dependent picture of the Raman process, not only because it is based on the time-dependent strategies developed so far, but especially for the possibility to compute the cumulative Raman signal at different times of the process. Giving a time-dependent picture of the process allows explicit inclusion of phenomena that have the same time scale of the Raman scattering, such as the interaction with an incident pulse with femtoseconds to picoseconds duration and any possible profile, and the vibrational relaxation, which is particularly relevant in resonant conditions to simultaneously simulate Raman and fluorescence emission. Moreover, in a future perspective, this method allows inclusion of the mutual interaction with a nearby plasmonic nanoparticle, that not only affects the intensity of the Raman signal but may change also the shape and the time scale of the process due to the local field felt by the molecule. Based on this premise, our approach will be useful also to simulate results of SERS experiments in a time-dependent fashion.

This procedure has been applied to porphyrin to calculate the time-dependent Raman cross-section in resonant and nonresonant conditions exploiting also the effect of the electric field duration and the presence of vibrational relaxation. In nonresonant conditions the Raman scattering is perfectly distinguishable from the fluorescence emission as they appear in different spectral regions, and so as well the vibrational relaxation does not affect the Raman signal since it occurs on a longer time scale. On the side of the pulse shape, the field duration does not alter the total Raman signal gathered at the end of the process, while the fluorescence region is strongly affected by it.

On the other hand, when in resonant conditions the Raman scattering is rapidly followed by the fluorescence emission which is a more intense phenomenon that occupies the same spectral region, so the distinction between the two...
contributions is not trivial. Also the spectrum shape is very
different than the one in nonresonant conditions due to the
great enhancement of the peaks generated from the excited
vibronic states around the vertical excitation energy. In these
conditions, the presence of the vibrational relaxation is needed
since it occurs in the time scale of the fluorescence emission,
and indeed the spectrum is strongly influenced by the decay
process. Our approach gives information on the interplay
between Raman scattering and fluorescence emission in terms
of relative weight of the two processes on the generation of the
spectrum line shape even if the two contributions are not
quantitatively distinguishable.

■ ASSOCIATED CONTENT

Supporting Information
The Supporting Information is available free of charge at
https://pubs.acs.org/doi/10.1021/acs.jpca.2c05245.
Convergence of Raman spectrum with 50 trajectories,
comparison between porphyrin Raman spectrum
computed with the present TD method and with a
time-independent strategy, comparison between por-
phyrin Raman spectrum computed with the present TD
approach and the experimental one (PDF)

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■ REFERENCES

(1) Rostron, P.; Gaber, S.; Gaber, D. Raman spectroscopy, review.
Int. J. Tech. Res. 2016, 21, 24.
(2) Graves, P.; Gardiner, D. Practical Raman spectroscopy; Springer,
1989; Vol. 10.
(3) Ferraro, J. R. Introductory Raman spectroscopy; Elsevier, 2003.
(4) Kuhar, N.; Sil, S.; Verma, T.; Umaphathy, S. Challenges in
application of Raman spectroscopy to biology and materials. RSC adv
2018, 8, 25888–25908.
(5) Jones, R. R.; Hooper, D. C.; Zhang, L.; Wolverson, D.; Valey, V.
K. Raman techniques: fundamentals and frontiers. Nanoscale Res. Lett.
2019, 14, 1–34.
(6) Hess, C. New advances in using Raman spectroscopy for the
characterization of catalysts and catalytic reactions. Chem. Soc. Res.
2021, 50, 3519–3564.
(7) Das, R. S.; Agrawal, Y. Raman spectroscopy: Recent advances,
techniques and applications. Vib. Spectrosc. 2011, 57, 163–176.
(8) Balan, V.; Mihai, C.-T.; Cojocaru, F.-D.; Uritu, C.-M.; Dodi, G.;
Botezat, D.; Gardikiotis, I. Vibrational spectroscopy fingerprinting in
medicine: from molecular to clinical practice. Materials 2019, 12,
2884.
(9) Long, D. A.; Long, D. The Raman effect: a unified treatment of
the theory of Raman scattering by molecules; Wiley Chichester, 2002; Vol.
8.
(10) Schatz, G. C.; Ratner, M. A. Quantum mechanics in chemistry;
Courier Corporation, 2002.
(11) Tannor, D. J. Introduction to quantum mechanics: A time-
dependent perspective; University Science Books, 2007.
(12) Lee, S.-Y.; Heller, E. J. Time-dependent theory of Raman
scattering. J. Chem. Phys. 1979, 71, 4777–4788.
(13) Williams, S. O.; Imre, D. G. Raman spectroscopy: time-
dependent pictures. J. Phys. Chem. 1988, 92, 3363–3374.
(14) Kramers, H.; Heisenberg, W. Uber die Streuung von Strahlung
durch Atome. Z. Phys. 1925, 31, 681–709.
(15) Dirac, P. A. M.; Fowler, R. H. The quantum theory of
dispersion. Proc. R. Soc. London, Ser. A 1927, 114, 710–728.
(16) Tannor, D. J.; Heller, E. J. Polytomic Raman scattering for
general harmonic potentials. J. Chem. Phys. 1982, 77, 202–218.
(17) Albrecht, A. C. On the theory of Raman intensities. J. Chem.
Phys. 1961, 34, 1476–1484.
(18) Herzberg, G.; Teller, E. Schwingungsstruktur der Elektronen-
bergänge bei mehratomigen Molekülen. Z. Phys. Chem. 1933, 21,
410–446.
(19) Small, G. J. Herzberg–Teller vibronic coupling and the
Duschinsky effect. J. Chem. Phys. 1971, 54, 3300–3306.
(20) Kupka, H.; Cribb, P. Multidimensional Franck–Condon
integrals and Duschinsky mixing effects. J. Chem. Phys. 1986, 85,
1303–1315.
(21) Santoro, F.; Cappelli, C.; Barone, V. Effective Time-
Independent Calculations of Vibrational Resonance Raman Spectra
of Isolated and Solvated Molecules Including Duschinsky and
Herzberg–Teller Effects. J. Chem. Theory Comput. 2011, 7, 1824–
1839.
(22) Egidì, F.; Bolino, J.; Cappelli, C.; Barone, V. A robust and
effective time-independent route to the calculation of resonance
raman spectra of large molecules in condensed phases with the
inclusion of Duschinsky, Herzberg–Teller, anharmonic, and environ-
mental effects. J. Chem. Theory Comput. 2014, 10, 346–363.
(23) Neugebauer, J.; Reiher, M.; Kind, C.; Hess, B. A. Quantum
chemical calculation of vibrational spectra of large molecules—Raman
and IR spectra for buckminsterfullerene. J. Comput. Chem. 2002, 23,
895–910.
(24) Kane, K. A.; Jensen, L. Calculation of absolute resonance
Raman intensities: Vibronic theory vs short-time approximation. J.
Phys. Chem. C 2010, 114, 5540–5546.
(25) Heller, E. J.; Sundberg, R.; Tannor, D. Simple aspects of Raman
scattering. J. Phys. Chem. 1982, 86, 1822–1833.
(26) Jensen, L.; Zhao, L.; Autschbach, J.; Schatz, G. Theory and
method for calculating resonance Raman scattering from resonance
polarizability derivatives. J. Chem. Phys. 2005, 123, 174110.
(27) Jensen, L.; Schatz, G. C. Resonance Raman scattering of
rhodamine 6G as calculated using time-dependent density functional
theory. J. Phys. Chem. A 2006, 110, 5973–5977.
(28) Guthmuller, J.; Champagne, B. Time dependent density
functional theory investigation of the resonance Raman properties of
the julolidinemalononitrile push-pull chromophore in various
solvents. J. Chem. Phys. 2007, 127, 164507.
(29) Guthmuller, J.; Champagne, B. Resonance Raman scattering of
rhodamine 6G as calculated by time-dependent density functional
theory: vibronic and solvent effects. J. Phys. Chem. A 2008, 112,
3215–3223.
(30) Ma, H.; Liu, J.; Liang, W. Time-dependent approach to resonance Raman spectra including Duschinsky rotation and Herzberg–Teller effects: Formalism and its realistic applications. J. Chem. Theory Comput. 2012, 8, 4474–4482.

(31) Thomas, M.; Latorre, F.; Marquetand, P. Resonance Raman spectra of ortho-nitrophenol calculated by real-time time-dependent density functional theory. J. Chem. Phys. 2013, 138, 044101.

(32) Mattiat, J.; Lubr, S. Efficient calculation of (resonance) Raman spectra and excitation profiles with real-time propagation. J. Chem. Phys. 2018, 149, 174108.

(33) Guthmuller, J. Comparison of simplified sum-over-state expressions to calculate resonance Raman intensities including Franck-Condon and Herzberg-Teller effects. J. Chem. Phys. 2016, 144, 064106.

(34) Guthmuller, J. Calculation of Vibrational Resonance Raman Spectra of Molecules Using Quantum Chemistry Methods. Molecular Spectroscopy: A Quantum Chemistry Approach 2019, 2, 497–536.

(35) Guthmuller, J. Sum-over-state expressions including second-order Herzberg–Teller effects for the calculation of absorption and resonance Raman intensities. J. Chem. Phys. 2021, 155, 084107.

(36) Baiardi, A.; Bloino, J.; Barone, V. A general time-dependent route to Resonance-Raman spectroscopy including Franck-Condon, Herzberg-Teller and Duschinsky effects. J. Chem. Phys. 2014, 141, 114108.

(37) Baiardi, A.; Bloino, J.; Barone, V. Time-dependent formulation of resonance Raman optical activity spectroscopy. J. Chem. Theory Comput. 2018, 14, 6370–6390.

(38) Barone, V.; Biczysko, M.; Bloino, J. Fully anharmonic IR and Raman spectra of medium-size molecular systems: accuracy and interpretation. Phys. Chem. Chem. Phys. 2014, 16, 1759–1787.

(39) Bloino, J.; Biczysko, M.; Barone, V. Anharmonic effects on vibrational spectra intensities: infrared, Raman, vibrational circular dichroism, and Raman optical activity. J. Phys. Chem. A 2015, 119, 11862–11874.

(40) Baiardi, A.; Bloino, J.; Barone, V. Accurate simulation of resonance-Raman spectra of flexible molecules: An internal coordinates approach. J. Chem. Theory Comput. 2015, 11, 3267–3280.

(41) Holtum, T.; Bloino, J.; Pappas, C.; Kumar, V.; Barone, V.; Schlücker, S. Ultraviolet resonance Raman spectroscopy of anthracene: Experiment and theory. J. Raman Spectrosc. 2021, 52, 2292–2300.

(42) Avila Ferrer, F. J.; Barone, V.; Cappelli, C.; Santoro, F. Duschinsky, Herzig-Teller, and Multiple Electronic Resonance Interferential Effects in Resonance Raman Spectra and Excitation Profiles. The Case of Pyrene. J. Chem. Theory Comput. 2013, 9, 3597–3611.

(43) Coccia, E.; Troiani, F.; Corni, S. Probing quantum coherence in ultrafast molecular processes: An ab initio approach to open quantum systems. J. Chem. Phys. 2018, 148, 204112.

(44) Coccia, E.; Corni, S. Role of coherence in the plasmonic control of molecular absorption. J. Chem. Phys. 2019, 151, 044703.

(45) Dall’Osto, G.; Coccia, E.; Guido, C. A.; Corni, S. Investigating ultrafast two-pulse experiments on single DNQDI fluorophores: a stochastic quantum approach. Phys. Chem. Chem. Phys. 2020, 22, 16734–16746.

(46) Domcke, W.; Stock, G. Theory of ultrafast nonadiabatic excited-state processes and their spectroscopic detection in real time. Adv. Chem. Phys. 1997, 100, 15.

(47) Santoro, F.; Petrongolo, C.; Lami, A. Time- and frequency-resolved spontaneous emission: Theory and application to the NO_2AXA’/A2A’ conical intersection. J. Chem. Phys. 2000, 113, 4073–4082.

(48) Gelin, M. F.; Pislakow, A. V.; Domcke, W. Time- and frequency-gated spontaneous emission as a tool for studying vibrational dynamics in the excited state. Phys. Rev. A 2002, 65, 062507.

(49) Sun, Z.; Lu, J.; Zhang, D. H.; Lee, S. Y. Quantum theory of (femtosecond) time-resolved stimulated Raman scattering. J. Chem. Phys. 2008, 128, 144114.

(50) Gelin, M. F.; Domcke, W.; Rao, B. J. Femtosecond stimulated Raman spectroscopy as a tool to detect molecular vibrations in ground and excited electronic states. J. Chem. Phys. 2016, 144, 184307.

(51) Langer, J.; Jimenez de Aberasturi, D.; Aizpurua, J.; Alvarez-Puebla, R. A.; Auguè, B.; Baumberg, J. J.; Bazan, G. C.; Bell, S. E. J.; Boisen, A.; Brolo, A. G.; et al. Present and Future of Surface-Enhanced Raman Scattering. ACS Nano 2020, 14, 28–117.

(52) Le, F.; Brandl, D.; Urzhumov, Y.; Wang, H.; Kundu, J.; Halas, N.; Aizpurua, J.; Nordlander, P. Metallic Nanoparticle Arrays: A Common Substrate for Both Surface-Enhanced Raman Scattering and Surface-Enhanced Infrared Absorption. ACS Nano 2008, 2, 707–718.

(53) Ding, S.-Y.; Yi, J.; Li, J.-F.; Ren, B.; Wu, D.-Y.; Panneerselvam, R.; Tian, Z.-Q. Nanostructure-based plasmon-enhanced Raman spectroscopy for surface analysis of materials. Nat. Rev. Mater. 2016, 1, 1–16.

(54) Corni, S.; Tomasi, J. Surface enhanced Raman scattering from a single molecule adsorbed on a metal particle aggregate: A theoretical study. J. Chem. Phys. 2002, 116, 1156–1164.

(55) Pipolo, S.; Corni, S. Real-Time Description of the Electronic Dynamics for a Molecule Close to a Plasmonic Nanoparticle. J. Phys. Chem. C 2016, 120, 28774–28781.

(56) Dall’Osto, G.; Gil, G.; Pipolo, S.; Corni, S. Real-time dynamics of plasmonic resonances in nanoparticles described by a boundary element method with generic dielectric function. J. Chem. Phys. 2020, 153, 184114.

(57) Baer, M. Beyond Born-Oppenheimer: electronic nonadiabatic coupling terms and conical intersections; John Wiley & Sons, 2006.

(58) Worth, G. A.; Cederbaum, L. S. Beyond Born-Oppenheimer: molecular dynamics through a conical intersection. Annu. Rev. Phys. Chem. 2004, 55, 127–58.

(59) Yonehara, T.; Hansukai, K.; Takatsuka, K. Fundamental approaches to nonadiabaticity: Toward a chemical theory beyond the Born–Oppenheimer paradigm. Chem. Rev. 2012, 112, 499–542.

(60) Barone, V. Accurate vibrational spectra of large molecules by density functional computations beyond the harmonic approximation: the case of azabenzenes. J. Phys. Chem. A 2004, 108, 4146–4150.

(61) Carbonnier, P.; Lucca, T.; Pouchan, C.; Rega, N.; Barone, V. Vibrational computations beyond the harmonic approximation: Performances of the B3LYP density functional for semirigid molecules. J. Comput. Chem. 2005, 26, 384–388.

(62) Santoro, F.; Lami, A.; Improta, R.; Barone, V. Effective method to compute vibrationally resolved optical spectra of large molecules at finite temperature in the gas phase and in solution. J. Chem. Phys. 2007, 126, 184102.

(63) Pipolo, S.; Corni, S.; Cammi, R. Equation of motion for the solvent polarization apparent charges in the polarizable continuum model: Application to time-dependent CI. J. Chem. Phys. 2017, 146, 064116.

(64) Van Kampen, N. G. Stochastic Processes in Physics and Chemistry; North Holland, 2007.

(65) Tremblay, J. C.; Klamroth, T.; Saalfrank, P. Time dependent configuration interaction calculations of laser driven dynamics in presence of dissipation. J. Chem. Phys. 2008, 129, 84302.

(66) Biele, R.; D’Agosta, R. A stochastic approach to open quantum systems. J. Condens. Matter Phys. 2012, 24, 273201.

(67) Fritzsch, M. J.; Trucks, G. W.; Schlegel, H. B.; Scuseria, G. E.; Robb, M. A.; Cheeseman, J. R.; Scalmani, G.; Barone, V.; Petersson, G. A.; Nakatsuji, H.; et al., Gaussian 16, rev B.01; Gaussian Inc: Wallingford CT, 2016.

(68) Santoro, F.; Cerezo, J. FCclasses3, a code for vibronic calculations. http://www.icom.cn.it/en/fcclasses, 2022 (accessed 27-July-2022).

(69) Coccia, E.; Dall’Osto, G.; Fregoni, J.; Gil, G.; Marsili, M.; Pipolo, S.; Romanelli, M.; Rosa, M.; Corni, S. WaveT-TDPolas. https://github.com/stefano-corni/WaveT_TDPlas, 2021 (accessed 10-January-2022).

(70) Marcelli, A.; Foggii, P.; Moroni, L.; Gellini, C.; Salvi, P. R. Excited-state absorption and ultrafast relaxation dynamics of...
porphyrin, diprotonated porphyrin, and tetraoxaporphyrin dication. J. Phys. Chem. A 2008, 112, 1864–1872.

(71) Taniguchi, M.; Mass, O.; Boyle, P. D.; Tang, Q.; Diers, J. R.; Bocian, D. F.; Holten, D.; Lindsey, J. S. Structural studies of sparsely substituted synthetic chlorins and phorhines establish benchmarks for changes in the ligand core and framework of chlorophyll macrocycles. J. Mol. Struct. 2010, 979, 27–45.

(72) Halls, M. D.; Velkovski, J.; Schlegel, H. B. Harmonic frequency scaling factors for Hartree-Fock, S-VWN, B-LYP, B3-LYP, B3-PW91 and MP2 with the Sadlej pVTZ electric property basis set. Theor. Chem. Acc. 2001, 105, 413–421.

(73) Irikura, K. K.; Johnson, R. D.; Kacker, R. N. Uncertainties in scaling factors for ab initio vibrational frequencies. J. Phys. Chem. A 2005, 109, 8430–8437.