

# Internal Conversion and Vibrational Energy Redistribution in Chlorophyll A

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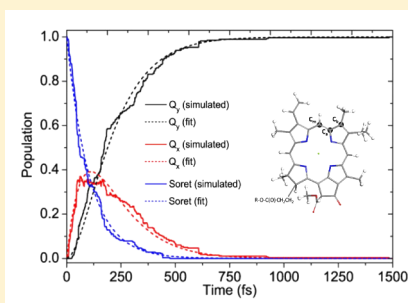
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## Supporting Information

**ABSTRACT:** We have computationally investigated the role of intramolecular vibrational modes in determining nonradiative relaxation pathways of photoexcited electronic states in isolated chlorophyll A (ChlA) molecules. To simulate the excited state relaxation from the initially excited *Soret* state to the lowest excited state  $Q_x$ , the approach of nonadiabatic excited state molecular dynamics has been adopted. The intramolecular vibrational energy relaxation and redistribution that accompany the electronic internal conversion process is followed by analyzing the excited state trajectories in terms of the ground state equilibrium normal modes. The time dependence of the normal mode velocities is determined by projecting instantaneous Cartesian velocities onto the normal mode vectors. Our analysis of the time evolution of the average mode energies uncovers that only a small subset of the medium-to-high frequency normal modes actively participate in the electronic relaxation processes. These active modes are characterized by the highest overlap with the nonadiabatic coupling vectors (NACRs) during the electronic transitions. Further statistical analysis of the nonadiabatic transitions reveals that the electronic and vibrational energy relaxation occurs via two distinct pathways with significantly different time scales on which the hopping from *Soret* to  $Q_x$  occurs thereby dictating the overall dynamics. Furthermore, the NACRs corresponding to each of the transitions have been consistently found to be predominantly similar to a set of normal modes that vary depending upon the transition and the identified categories. Each pathway exhibits a differential time scale of energy transfer and also a differential set of predominant active modes. Our present analysis can be considered as a general approach allowing identification of a reduced subset of specific vibrational coordinates associated with nonradiative relaxation pathways. Therefore, it represents an adequate prior strategy that can particularly facilitates mixed quantum-classical approaches.



## 1. INTRODUCTION

Given the supreme importance of natural photosynthesis in sustaining life on the Earth, unraveling its secrets has been a pivotal research area over many decades, especially in the light of depleting conventional energy sources. A sophisticated biological machinery enables harvesting of solar energy and its subsequent conversion to chemical energy inside the thylakoid membranes in the chloroplasts of green plants. Despite the great complexity of the whole process, it is surprisingly hinged upon very few pigment molecules known as chlorophylls. Chlorophylls not only act as primary photoactive pigments and enable energy transfer pathways to the so-called “reaction center” but also participate in charge separation that triggers photochemical conversion.<sup>1,6,7</sup> Dynamics of the photoexcited states in chlorophylls is thus central to a number of aspects of photosynthesis.<sup>8,9</sup>

Previously,<sup>10</sup> we have measured the time constants for high excited states to relax nonradiatively to the lowest excited state ( $Q_x$ ) in Chlorophyll A (ChlA) and B (ChlB) with ultrafast transient absorption spectroscopy and modeled it using the nonadiabatic excited state molecular dynamics (NA-ESMD) methodology.<sup>64</sup> The slight difference in the molecular structures of these two chlorophylls was found to lead to slightly slower dynamics of  $B \rightarrow Q_x$  internal conversion in ChlB in agreement with experimental spectroscopic data.<sup>10</sup> Shi et al. demonstrated that the time-constant for internal conversion increases with an increase in the solvent dielectric constant.<sup>11</sup> Dong et al. further estimated the vibrational relaxation time to be 1.5 ps for ChlA solvated in common solvents such as ethyl

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acetate and ethyl ether.<sup>12,13</sup> Even though the internal conversion in chlorophylls itself is surprisingly understudied, it has received significant attention in various other light-harvesting pigments such as bacteriochlorophylls,<sup>14–17</sup> carotenoids,<sup>18–22</sup> and porphyrins. Within a wider view, importance of understanding internal conversion is evident from a number of related studies in biological macromolecules such as nucleic acids<sup>23,24</sup> and many other organic/inorganic molecules or compounds.<sup>25,30,31</sup>

Nonradiative relaxation of excited electronic states is invariably accompanied by vibrational energy redistribution, and a number of studies point to the importance of “active” vibrational modes in assisting internal conversion processes.<sup>32–36</sup> Experimentally, such studies are generally based on variants of excited state IR spectroscopy,<sup>37–39</sup> ultrafast time-resolved IR spectroscopy,<sup>40–42</sup> and ultrafast transient IR spectroscopy.<sup>43</sup> These techniques have been shown to be powerful in the study of the excited states decay.<sup>47–57</sup> Much of the theoretical work on intramolecular energy transfer in organic molecules has involved classical and semiclassical Hamiltonians using harmonic approximations to identify the pathways and life times of energy transfer out of a given bond or mode.<sup>44–46</sup> Equilibrium normal modes (ENMs), typically calculated from second derivatives of the ground-state (GS) energy with respect to nuclear coordinates (Hessian), can be used to elucidate vibrational motions of polyatomic molecules.<sup>58–61</sup> Within this theoretical framework, one can then address the vibrational energy flow by following the time evolution of the kinetic energy of each ENM. Anharmonic effects and mode couplings have, however, been recognized to be important toward giving rise to vibrational energy redistribution. Previous work has extensively discussed these aspects in proteins.<sup>26–29</sup> In particular, for ChlA, for example, density functional theory calculations have been used to investigate the vibrational properties and to ascertain couplings between important infrared-active modes of the carbonyl groups.<sup>3,62</sup> In this work, following up on our earlier study, we focus on characterizing the role of vibrational energy redistribution in the nonradiative relaxation of high-energy excited states in ChlA. Our aim is to identify the nature of active normal modes related to the electronic couplings and intramolecular energy transfer throughout the nonadiabatic excited-state molecular dynamics. We show that only a small subset of vibrational modes of ChlA actively aid the internal conversion process.

The remainder of the paper is organized as follows. The underlying theory for the NA-ESMD methodology is briefly described in the next section 2. Results are presented and discussed in detail in section 3 before drawing concluding remarks in the final section, section 4.

## 2. THEORETICAL AND COMPUTATIONAL DETAILS

**2.1. NA-ESMD Background.** The NA-ESMD framework<sup>63,64</sup> allows for simulation of photoinduced dynamics of large organic conjugated molecules involving multiple coupled electronic excited states, a formidable task carried out by combining the molecular dynamics with quantum transitions approach<sup>65,66</sup> with “on the fly” analytical calculations of excited-state energies,<sup>67–69</sup> gradients,<sup>70,71</sup> and nonadiabatic coupling<sup>63,72–74</sup> terms. For this purpose, the collective electron oscillator (CEO) method<sup>75–78</sup> is applied using the AM1 semiempirical Hamiltonian<sup>79,80</sup> at the configuration interaction singles level.<sup>81,82</sup> Previously, AM1 and similar semiempirical

models have been successfully applied to describe excited state properties of porphyrin-based molecules.<sup>80,83–91</sup> A detailed discussion on the implementation of, the advantages of, and various testing parameters of NA-ESMD can be found elsewhere.<sup>63,64,97</sup>

**2.2. Normal Mode Analysis.** ENMs are computed at the ground state state minimum  $\mathbf{R}_0$  from the mass-weighted Hessian matrix  $\mathbf{H}$  with elements

$$\mathbf{H}_{ij}(\mathbf{R}_0) = -\partial^2 E / \partial q_i \partial q_j \quad (1)$$

where  $E$  is the ground state potential energy and  $q_i = \sqrt{m_i}(R_i - R_{0,i})$  is the mass weighted Cartesian displacement of  $i$ th atom with mass  $m_i$  and Cartesian coordinates  $R_{0,i}(X_{0,i}, Y_{0,i}, Z_{0,i})$ .<sup>92–94,96,98–102</sup> Geometry optimization was performed on the chlorophyll molecule with the phytol tail replaced by a methyl group for computational ease, with the AM1 semiempirical Hamiltonian, using Gaussian 09.<sup>103</sup> Diagonalization of  $\mathbf{H}$  produces the set of orthonormal ENM with frequencies  $\nu_i$  related to the eigenvalues  $\lambda_i (i = 1, \dots, 3N - 6)$  of  $\mathbf{H}$  as  $\nu_i = (\sqrt{\lambda_i} / 2\pi)$ . ENMs are defined in a body-fixed reference frame with the origin at the center of mass of the molecule and axes corresponding to its principle axes of inertia. Upon diagonalization of  $\mathbf{H}$ , the set of ENM vectors  $\{Q_i\}$ ,  $i = 1, 3N - 6$  are obtained from the corresponding eigenvector matrix  $\mathbf{L}$ , which allows to express their amplitudes throughout the NA-ESMD as a linear combination of the set  $\{q_i\}_{i=1,3N}$  as

$$Q_i(t) = \sum_{j=1}^{3N} l_{ji} q_j(t) \quad i = 1, \dots, 3N - 6 \quad (2)$$

where  $l_{ji}$  are elements of the eigenvector matrix  $\mathbf{L}$ . By differentiating this expression with respect to time, we obtain the relationship between the momenta

$$\dot{Q}_i(t) = \sum_{j=1}^{3N} l_{ji} \dot{q}_j(t) \quad i = 1, \dots, 3N - 6 \quad (3)$$

In this way, the vibrational kinetic energy transforms in terms of the ENM momenta as

$$K(t) = \frac{1}{2} \sum_{i=1}^{3N-6} (\dot{Q}_i(t))^2 \quad (4)$$

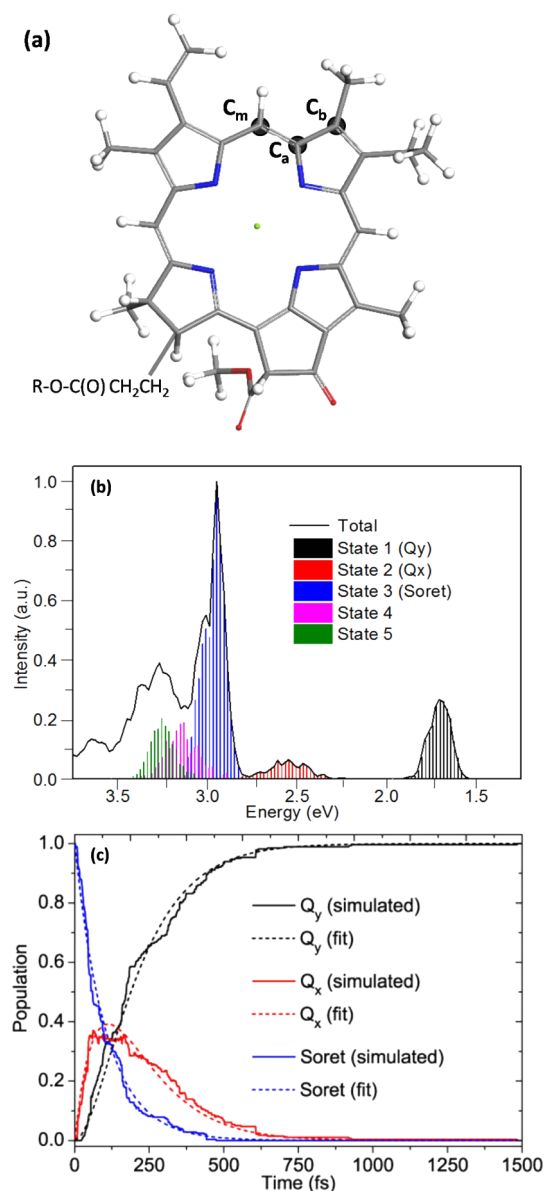
The total vibrational energy associated with a given mode is calculated using the virial theorem,  $E_i(t) = 2K_i(t)$ .

**2.3. Molecular Dynamics Simulations.** The ground state molecular dynamics simulation was then performed at 300 K with a time step  $\Delta t = 0.5$  fs. The system was heated to allow thermal equilibration at a final temperature of 300 K during the first 10 ps. The Langevin equation at constant temperature was utilized with a friction coefficient of  $\gamma = 2.0$  ps. This value has been chosen to allow an efficient temperature coupling<sup>95</sup> to obtain the adequate ground state conformational sampling. In the 8 ns long production run following equilibration, we sample atomic coordinates and momenta at every 20 ps generating a set of 400 initial positions and momenta for the subsequent NA-ESMD simulations. Details of the assignment of the initial excited state for each NA-ESMD trajectories according to a Franck–Condon window centered at 2.94 eV (lowest state in the Soret band) can be found elsewhere.<sup>10</sup> As opposed to the initial ground state simulation, NA-ESMD simulations have been carried out at constant energy during 1.5 ps using a

classical time-step of  $\Delta t = 0.1$  fs and  $N_q = 4$  quantum time steps per classical step to simultaneously propagate the electronic coefficients for the 10 lowest excited states. The existence of trivial unavoided crossings has been examined by tracking the identities of the states.<sup>104</sup>

### 3. RESULTS AND DISCUSSIONS

We have investigated the dynamics of the internal conversion process of ChlA (Figure 1a) that takes place after its initial photoexcitation in the red edge ( $S_3$ ) of the *Soret* (B) band (see Figure 1b). Subsequent to laser excitation, the molecular system experiences a fast  $B \rightarrow Q_x \rightarrow Q_y$  electronic energy

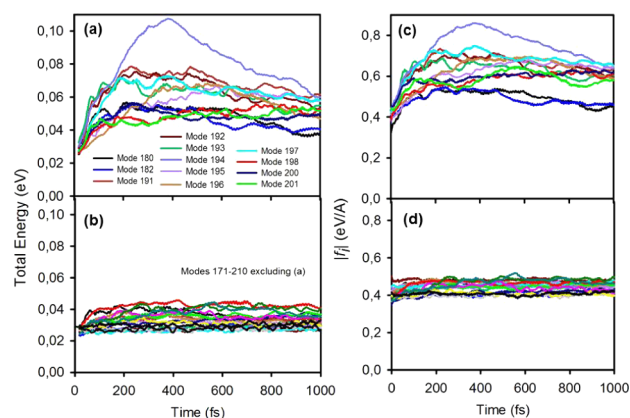


**Figure 1.** (a) Molecular structure of chlorophyll A (carbon atoms in gray, oxygen in red, nitrogen in blue, magnesium in green and hydrogen in light gray), where R denotes the phytol tail. (b) Simulated absorption spectrum of ChlA where contributions from the 5 lowest energy excited states have been included; and (c) Populations of excited states following initial excitation centered at the lowest state in the *Soret* band. Results are fitted (dashed curves) with a sequential kinetic rate constant model.

relaxation and redistribution that involves the selective participation of specific atomic groups and complex global migration of the electronic wave function within the total carbon macrocycle of the porphyrin ring structure.<sup>10</sup> Figure 1c represents the average populations of various electronic states as a function of time. Here, the population of the *Soret* state is taken to be a summation of the populations on  $S_3$  and  $S_4$  as a small percentage (<4%) of configurations are initially excited at  $S_4$ . The lowest excited state is completely populated within 1.5 ps. By fitting the simulations data with a sequential kinetic model we can extract the time constant as  $\tau_1 = 105$  fs for the *Soret*  $\rightarrow Q_x$  process and  $\tau_2 = 124$  fs for the  $Q_x \rightarrow Q_y$  relaxation. The time constants obtained through the constant energy simulations are close to those obtained by us in the previous study using constant temperature simulations.

The intramolecular vibrational energy flow in ChlA, accompanied by the electronic energy transfer, is monitored by evaluating the time evolution of the normal mode velocities obtained by projection of the body-fixed mass weighted Cartesian velocities onto the normal modes vectors (see Section Normal Mode Analysis).

Parts a and b of Figure 2 show the vibrational energy  $E_i$  along different intermediate-range and high-frequency ENMs (twice



**Figure 2.** Time evolution of the total energy of trajectories along (a) normal modes 180, 182, and 191–201 and (b) all the normal modes from 171 to 210 excluding those plotted in part a; time evolution of the norm of the normal mode forces averaged over the trajectories for (c) normal modes 180, 182, 191–201; (d) all the normal modes from 171 to 210 excluding those plotted in part c.

the kinetic energy considering the equipartition of energy) averaged over the ensemble of NA-ESMD trajectories. It is observed first that the intramolecular vibrational energy redistribution is not statistical, since the energy deposited in each ENM varies significantly. It is important to stress that, in the present work, NA-ESMD simulations have been performed at constant energy. Therefore, the lack of dominant vibronic relaxation paths during the internal conversion process should lead to transient and final accumulations of slightly different amounts of the initial excess of energy spread among all the 240 vibrational normal modes of the molecule. Except for the 191th to 201th, and also 180th and 182th ENMs, the energies of all other ENMs deviate only slightly from their equilibrium values during the internal conversion process. We have defined a normal mode as active during the internal conversion process if, at any time during the NA-ESMD simulations, its average energy exceeds twice the initial  $kT = 0.026$  eV equilibrium value, thus reflecting a strong coupling to the electronic degrees

of freedom. These reduced number of active modes experience a fast increase in their energy reaching their maxima at about 400 fs. The maximum combined energy stored in all of these active modes is 0.874 eV (7049  $\text{cm}^{-1}$ , with the highest contribution coming from mode 194 at 0.108 eV (871  $\text{cm}^{-1}$ ). After that, they experience a slow energy relaxation that is extended for more than 1 ps. The rest of the modes seems to act as a bath of coupled harmonic oscillators to which the excess of energy, transiently accumulated in these active modes, is finally transferred.

A careful analysis along each of the 240 normal modes reveals that only a small subset of them indexed predominantly from modes number 180 (1465  $\text{cm}^{-1}$ ), 182 (1491  $\text{cm}^{-1}$ ), and 191–201 (1665–1908  $\text{cm}^{-1}$ ) are strongly excited during the excited state relaxation process as shown in Figure 2a. The energy plots in Figure 2b along any of the remaining normal modes in a similar frequency bracket suggest that they are not involved in the excited state dynamics. We have thus identified a few vibrational modes that are intricately associated with *Soret* to  $Q_y$  internal conversion in ChlA. These modes can be associated with the 1597–1620  $\text{cm}^{-1}$  band observed in the IR spectra of most chlorophyll systems.<sup>2,3</sup> This assignment is based on previous work performed by Wang and co-workers.<sup>3</sup> Despite the frequency difference between the experimental IR band and the calculated one, the assignment can be performed based on the common relative nuclear displacements between vibrations associated with this experimental band and relative nuclear displacements involved in the active normal modes. On the basis of the normal mode assignments performed by Boldt et al.,<sup>5</sup> this band is predominantly associated with  $C_aC_m$ ,  $C_aC_b$ , and  $C_bC_b$  stretching vibrations (see Figure 1a).<sup>4,5</sup>

Parts c and d of Figure 2 show the time-dependence of the norm of different ENM forces averaged over the trajectories. Thereof we can point out the role that gradients/forces on the excited states play during this intramolecular vibrational energy redistribution. Upon vertical excitation to the Soret band, an ultrafast transfer of electronic to vibrational energy is observed as an increase of the forces in the direction of the modes previously identified as active normal modes. The relative low initial values of these forces increase in time until they reach their maxima at about 400 fs. This behavior indicates that the redistribution of the energy does not take place as the result of the initial ultrafast adiabatic relaxation that occurs right after photoexcitation, but rather as a consequence of nonadiabatic dynamics.

In order to further elucidate the role played by these vibrational normal modes in the internal conversion process, we investigate the correspondence between such active modes and the nonadiabatic coupling vectors (NACRs) defined as

$$d_{ij} = \langle \psi_i(r, R) | \nabla_R \psi_j(r, R) \rangle \quad (5)$$

where  $r$  and  $R$  are the electronic and nuclear vector coordinates, respectively, and  $\psi_i(r, R)$  is the  $i$ th CI electronic state.

The direction of NACR can be interpreted as the nonadiabatic contribution to the direction of the main driving force on the nuclei during electronic transitions.<sup>33</sup> Figure 3 depict the projection of the forces on the direction of NACR throughout the NAESMD simulations. After photoexcitation, the molecular system enters in regions of the configurational phase space with strong nonadiabatic couplings. The non-adiabatic contributions to the nuclear forces rapidly increase in time. At earlier times, the direction of  $\text{NACR}(\text{Soret}-Q_x)$

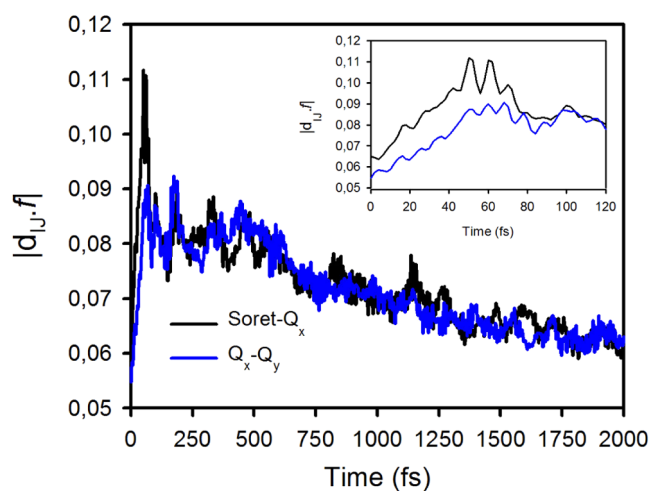
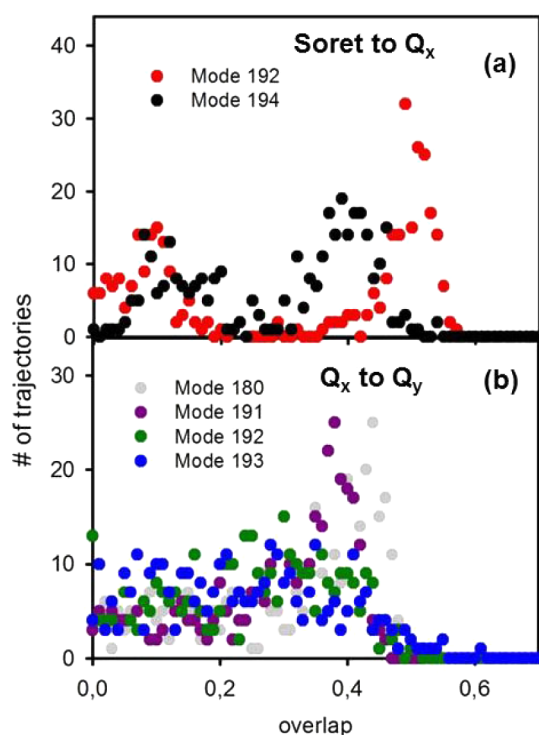


Figure 3. Time evolution of the projection of the forces on the direction of NACR averaged over the trajectories.

represents a higher contribution than the  $\text{NACR}(Q_x-Q_y)$  one. After the first 100 fs of the simulations, the nonadiabatic contributions of either  $\text{NACR}(\text{Soret}-Q_x)$  and  $\text{NACR}(Q_x-Q_y)$  persist at relatively higher values during the next 400 fs. After that, both contributions decay while the system adiabatically relaxes on the  $Q_y$  state. Therefore, the nonadiabatic contribution of the nuclear forces seems to dictate the fast intramolecular vibrational energy redistribution that occurs during the internal conversion process. During nonadiabatic dynamics in the vicinity of level crossings, both the Pechukas forces and redistribution of electronic energy excess upon quantum transitions are generally raising classical momenta of nuclei in the direction of NACR vectors. This ensures an effective flow of electronic energy into specific vibrational modes aligned with NACR vectors.

It is interesting at this point to associate each step of the *Soret* to  $Q_y$  internal conversion with the participation of the different active normal modes in the corresponding concomitant intramolecular vibrational energy pathways. To this end, from the ensemble of excited state trajectories, we first analyze two sets of NACRs - one corresponding to the *Soret* to  $Q_x$  transition and the other corresponding to the  $Q_x$  to  $Q_y$  relaxation. Each NACR vector, expressed in body-fixed reference frame, is projected on the basis of normal modes  $\{Q_j\}_{j=1,3N-6}$ . In Figure 4, we show the distributions of the overlap between NACR vectors and various selected normal modes. Figure 4a indicates a considerable overlap (>20%) between a substantial subset of *Soret* to  $Q_x$  NACRs and only a few normal modes indexed 192 and 194 while the overlap with the other modes is relatively small. On the other hand, for  $Q_x$  to  $Q_y$  transition, the overlaps between the NACRs and only the normal modes 180, 191, 192, and 193 are significant as shown in Figure 4b. This implies that the nuclear motions that modulate the electronic couplings, and therefore contribute to the ultrafast electronic energy transfer, involve directions given by only a few normal modes of intermediate-to-high frequencies. Besides, these modes that overlap the most with the NACR vectors, are in agreement with our previously identified active modes due to the significant accumulation of the excess vibrational energy during the relaxation process.

It is worth to mention that, according to the surface hopping prescription,<sup>65,105,106</sup> adjustments to the nuclear velocities are required in order to conserve total energy following hops



**Figure 4.** Distribution of the values of overlaps between specific normal modes and the set of NACRs corresponding to (a) *Soret* to  $Q_x$  and (b)  $Q_x$  to  $Q_y$  transitions.

between electronic states. The direction in which the velocity  $\dot{R}_i$  of each of the  $i$ th atom in the molecular system is commonly rescaled, corresponds to the direction of the NACR vector. Via an important test of reliability, we have ensured that changing the rescaling prescription does not alter the results of the dynamics (see [Supporting Information](#)).

For a more quantitative characterization of the electronic and vibrational energy pathways, representative NACR vectors associated with each  $I \rightarrow J$  transition can be defined. We construct matrices  $A_{IJ}$  of dimension  $3N \times K$ , with  $K$  being the number of trajectories with an effective  $I \rightarrow J$  hop. Matrices  $A_{IJ}$  are built with columns representing the NACR vector at the moment of effective  $I \rightarrow J$  transition in each of the  $K$  NA-ESMD trajectories. As these are not square matrices, in order to capture the representative NACR vectors of the whole set, Singular Value Decomposition (SVD) of  $A_{IJ}$  can be performed. Essentially it means that  $A_{IJ}$  is written as the product of a  $3N \times K$  column-orthogonal matrix  $U$ , a  $K \times K$  diagonal matrix  $W$  with positive or zero elements (the singular values), and the transpose of a  $K \times K$  orthogonal matrix  $V$ :

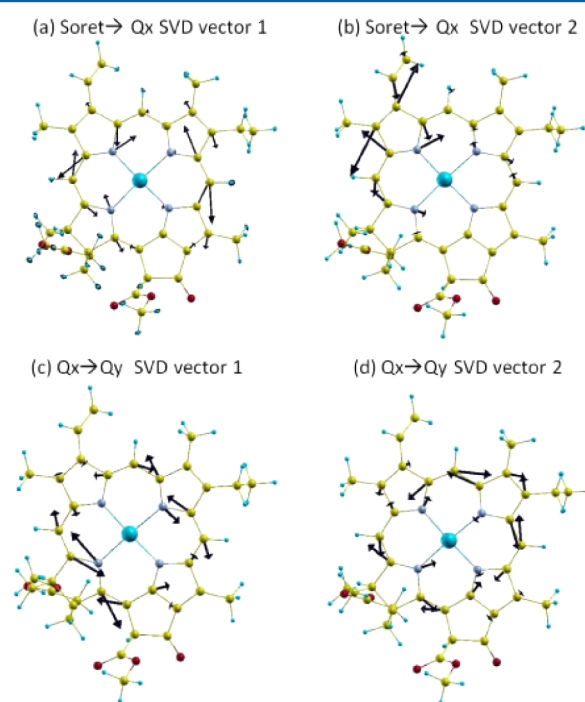
$$[A_{IJ}] = [U_{IJ}] \cdot [W_{IJ}] \cdot [V_{IJ}^T] \quad (6)$$

Thus, the  $a_{IJ}^{ij}$  elements of the matrix  $A_{IJ}$  can be expressed as the sum of products of columns of  $U_{IJ}$  and rows of  $V_{IJ}^T$  with the “weighting factors” being the singular values  $w_{IJ}^j$

$$a_{IJ}^{ij} = \sum_{l=1}^k w_{IJ}^{jl} \cdot u_{IJ}^{il} \cdot v_{IJ}^{jl} \quad (7)$$

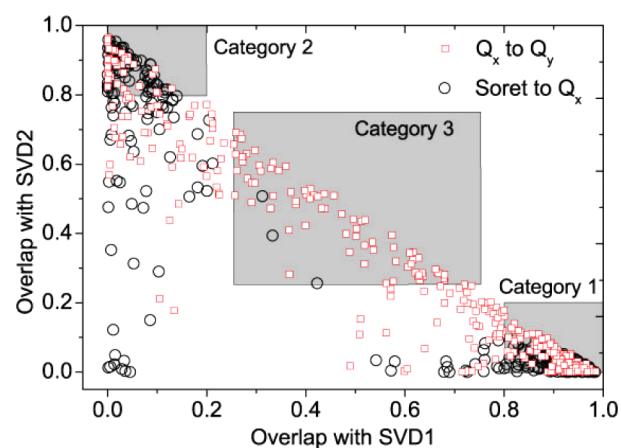
Subsequently, in this work, the  $u_{IJ}^l$  vectors, in the decreasing order of the values of  $w_{IJ}^l$ , are referred to as  $l$ th SVD vectors. The eigenvalues  $w_{IJ}^l$  indicate the correspondence between the entire data set and the individual SVD vectors. Here we find

that the first two SVD vectors, shown in [Figure 5](#), with the largest eigenvalues can sufficiently well describe more than 80%



**Figure 5.** Visualization of the first two SVD vectors for the NACRs corresponding to (a, b) *Soret* to  $Q_x$  and (c, d) to  $Q_x$  to  $Q_y$  internal conversion.

of the NACRs from the ensemble. An intuitive understanding of the SVD vectors can be obtained by plotting the overlap between an NACR and the first SVD vector 1 (SVD1) versus the overlap between the same NACR and SVD2. [Figure 6](#)

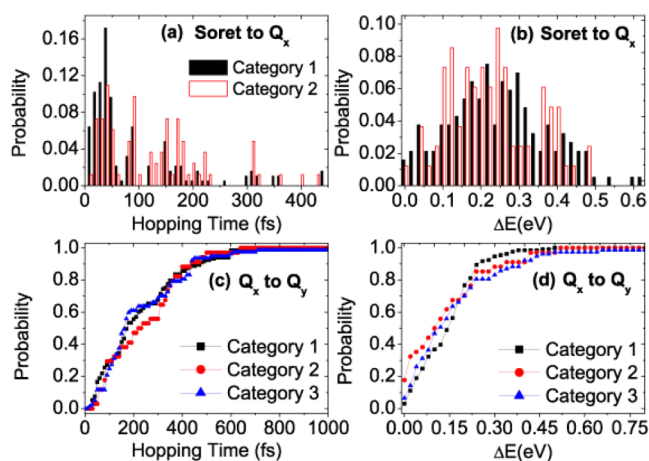


**Figure 6.** Overlap of each of the NACRs from the ensemble with the first SVD vector versus the overlap with the second SVD vector for *Soret* to  $Q_x$  (circles) and for  $Q_x$  to  $Q_y$  (squares) transitions.

shows that for the *Soret* to  $Q_x$  transition the complete set of NACRs can be distinctly categorized in two subsets—one (containing nearly 70% of the total NA-ESMD trajectories) which shows close resemblance with the SVD1 and the other with a very high overlap with the SVD2. This indicates that the *Soret* to  $Q_x$  transition may take place along two different pathways characterized by two different NACRs and sub-

sequently, the excited state dynamics along the two paths may exhibit important differences. Therefore, SVD1 and SVD2 vectors can be considered as an average representation of two distinguished original ensembles of NACR vectors, each of them identified with a particular *Soret* to  $Q_x$  transition. On the other hand, for the  $Q_x$  to  $Q_y$  transition, while the whole data set can still be represented by the first two SVD vectors, a distinct categorization does not emerge. The NACRs for this process are uniform combinations of SVD1 and SVD2. The subspace (plane) formed by both SVD vectors contains the overall ensemble of NACR vectors associated with the *Soret* to  $Q_x$  transition. According to the structural diagram of Chla that can be found elsewhere,<sup>4</sup> these vectors contain substantial contributions from  $C_a C_m$ ,  $C_a C_b$ , and  $C_b C_b$  stretching vibrations<sup>4,5</sup> that are associated with the IR spectra band observed at 1597–1620  $\text{cm}^{-1}$ .<sup>2,4</sup>

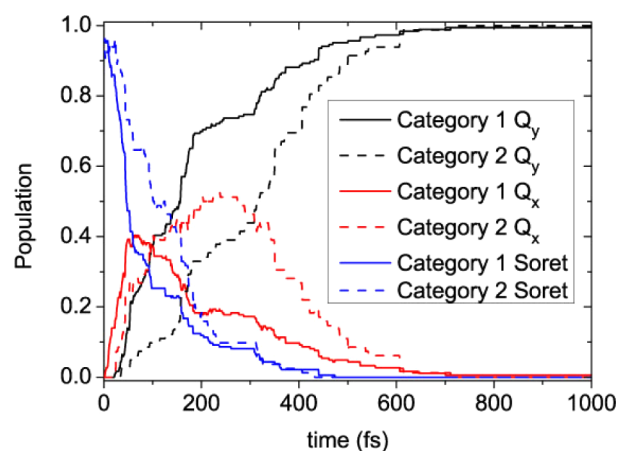
In order to ascertain whether the two distinct NACRs lead to two different relaxation pathways, we segregated the excited state dynamics trajectories based on the following criterion. Category 1 contains trajectories for which the NACRs exhibit more than 80% overlap with SVD1 for a given transition whereas Category 2 contains trajectories for which the NACRs show greater than 80% overlaps with SVD2. A third category relevant only to the  $Q_x$  to  $Q_y$  transition accounts for those trajectories for which the NACRs are a mixture of the first two SVD vectors. The shaded regions in Figure 6 also provide a visual aid to this categorizations. Figures 7a,b show the



**Figure 7.** Probability distribution of the (a) hopping times and (b)  $\Delta E$  at the time of hopping during *Soret* to  $Q_x$  transition for different categories based on SVD analysis described in text. For  $Q_x$  to  $Q_y$  transition, the corresponding plots for (c) hopping times and (d)  $\Delta E$  as cumulative distributions.

probability distribution of the hopping time and the difference in the energies at the time of hopping ( $\Delta E$ ), respectively, for the two categories of the *Soret* to  $Q_x$  internal conversion. The distribution of  $\Delta E$  can be seen to be relatively similar for both the categories, with the average value equal to 0.2554 eV for category 1 and 0.2343 eV for category 2. In order to clarify whether degeneracy (with  $\Delta E = 0$ ) can be reached by any of these two pathways, we adopt an arbitrary limit of  $\Delta E < 0.1$  eV to separate cases of strict degeneracy or their immediate regions and cases with larger  $\Delta E$ . With this separation in mind, we can say that less than only 16% of trajectories corresponding to each relaxation pathway pass through the degeneracy. The distributions of the hopping times are, however, significantly

different. This distinction becomes readily apparent in the average hopping time, which is much greater for category 2 trajectories (131.87 fs) than that for category 1 (90.93 fs). Both of these properties are however much more homogeneous for the  $Q_x$  to  $Q_y$  relaxation across all 3 categories as evident in the cumulative probability plots in Figure 7c,d. We have also subjected all of these distributions to the Kolmogorov–Smirnov test, which allows for an objective assessment of the similarity between any two distributions and have found that it shows that only the distributions in Figure 7a are distinct. We thus find that the *Soret* to  $Q_x$  transition occurs along two different pathways characterized by distinct NACRs. However, once the system reaches state  $Q_y$ , the subsequent relaxation process has uniform characteristics. Furthermore, we have calculated the population evolution on the different trajectory subsets based on the two categories obtained via SVD analysis of only the *Soret* to  $Q_x$  transition. The resulting plots shown in Figure 8 clearly point to the entirely different time scales of



**Figure 8.** Excited state dynamics for *Soret* to  $Q_x$  internal conversion for category 1 trajectories (solid curves) and category 2 trajectories (dashed curves).

both *Soret* to  $Q_x$  and the  $Q_x$  to  $Q_y$  internal conversion processes in the two categories. The time constant extracted via fitting simulated data with a sequential kinetic model is found to be 89.0 fs for category 1 trajectories and 133.3 fs for category 2 trajectories for the *Soret* to  $Q_x$  transition. For the  $Q_x$  to  $Q_y$  transition, the time constants are 94.4 and 187.3 fs for category 1 and 2, respectively.

Finally, we analyze the relationship between the categorization of the NA-ESMD trajectories for the two-stepped internal conversion process and the subset of the active normal modes identified earlier. We thus calculate the overlaps between the selected normal modes and the NACRs corresponding to each of the two transitions and average them within different ensembles determined by the aforementioned SVD based categorization. Table 1 lists the obtained results for all those active modes which appear in more than 90% of the trajectories within each category. For easier comparison, the most important modes, which show an overlap close to or higher than 0.25, are highlighted in bold. For the *Soret* to  $Q_x$  transition we can observe that the active modes are distinctly different for the two categories. Considering that the categorization is based on the SVD vectors, this implies that the *Soret* to  $Q_x$  transition is associated with two distinct NACRs that resemble different sets of normal modes. Furthermore, as

**Table 1. Average Values of Overlaps between Various Normal Modes and the NACRs Corresponding to Soret to  $Q_x$  and  $Q_x$  to  $Q_y$  Transitions<sup>a</sup>**

mode	Soret to $Q_x$		$Q_x$ to $Q_y$		
	cat 1	cat 2	cat 1	cat 2	cat 3
180	<b>0.269</b>	–	<b>0.436</b>	–	<b>0.289</b>
188	–	<b>0.317</b>	0.118	–	–
189	0.175	–	<b>0.299</b>	–	0.210
191	<b>0.300</b>	0.215	<b>0.391</b>	–	<b>0.272</b>
192	<b>0.502</b>	–	<b>0.334</b>	0.204	<b>0.253</b>
193	–	<b>0.238</b>	–	<b>0.335</b>	<b>0.301</b>
194	<b>0.403</b>	–	–	<b>0.284</b>	0.213
195	–	<b>0.281</b>	<b>0.243</b>	–	–
198	–	–	0.172	0.162	–
199	0.147	–	0.172	<b>0.251</b>	–
200	0.188	<b>0.272</b>	–	<b>0.424</b>	<b>0.276</b>
201	0.142	<b>0.428</b>	–	<b>0.311</b>	<b>0.241</b>

<sup>a</sup>Overlap values are shown for only those active modes which appear in >90% of the trajectories under each category. The values in bold emphasize the most important modes showing an overlap nearly equal to or greater than 0.25.

Figure 4a shows, the normal modes with prominent overlaps with the Soret to  $Q_x$  NACRs are indexed 192 and 194, which is consistent with the fact that nearly 70% of the trajectories belong to category 1. Modes 191, 192, and 194 are indeed also those, along which the strongest modulations of total energies are observed in Figure 2a. For the  $Q_x$  to  $Q_y$  transition as well one can note the distinctness of participating normal modes between category 1 and 2. However, the presence of the third category lends a significantly more mixed character that is consistent with the results presented in Figure 4b.

#### 4. CONCLUSIONS

We have identified the normal modes that participate actively in the intramolecular electronic energy relaxation and redistribution during the internal conversion process of the pigment chlorophyll A after its photoexcitation to the Soret band. The intramolecular vibrational energy flow, concomitant with the electronic energy transfer, is monitored by evaluating the temporal variance of the energy in each individual normal mode. Our analysis indicates that this process is not statistical, since the energy deposited on each mode varies significantly. Only a small subset of intermediate-to-high-frequency normal modes, with the highest overlap with the nonadiabatic coupling vectors at the instant of effective hops, experience a substantial increase in their vibrational energy during the process. Therefore, velocity adjustments, performed according to the surface hopping prescription, in the direction of the non-adiabatic coupling vectors can be well validated by the confirmation that this direction is the main vibrationally active direction during the internal conversion process. Two distinct pathways for nonradiative relaxation have been discovered, involving two different sets of active normal modes and significantly different time scales for the  $S_3 \rightarrow S_2$  transition. We consider that this present analysis can be used as a general approach allowing identification of a reduced subset of specific vibrational coordinates associated with nonradiative relaxation pathways. It is our hope that results reported here shall inspire further developments of hybrid quantum-classical molecular dynamics simulations in extended conjugated molecules that deal explicitly with the quantum mechanical treatment of a

small subset of normal modes and the conventional classical treatment of the remaining degrees of freedom.

#### ■ ASSOCIATED CONTENT

##### Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.jpbc.5b09548.

Description of a modified method for the rescaling of nuclear velocities as well as the results from a set of simulations with the modified algorithm (PDF)

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##### Notes

The authors declare no competing financial interest.

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