Nonadiabatic Sunlight-harvesting

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Experimental and theoretical evidence point out to the crucial role of specific resonant intramolecular vibrational modes in the interpretation of long-lived coherences observed in two-dimensional spectra of some natural and synthetic light harvesting complexes. For the natural situation of illumination by incoherent (sun)light, the relevance of these vibrations is analyzed here for light-harvesting vibronic prototype dimers. The detailed analysis of the density matrix dynamics reveals that the inclusion of the intramolecular vibrational modes reinforce up to one order of magnitude the exciton coherence and may increase the populations of lowest energy single exciton states, as well as populations and coherences in the site basis. In sharp contrast to the case of initial-state preparation by coherent (laser)light-sources, the initial thermal state of the local vibrational modes, as well as that of the anti-correlated mode, evolves devoid of non-classical correlations as confirmed by the absence of negative values of its phase-space quasi–probability distribution at all times. Therefore, not only the long-lived coherences observed in two-dimensional spectra are induced by the coherent character of pulsed laser sources, but it is unambiguously shown here that the non-classical character generally assigned to the anti-correlated vibrational mode also comes as the result of the preparation of the initial state by coherent pulsed laser sources.

In the last three decades, new experimental, theoretical and computational techniques have been developed to resolve the interplay between the multiple electronic and vibrational degrees of freedom, together with the variety of energy scales involved in molecular-aggregates energy-transfer-processes. [1–7]. In doing so, two-dimensional-electronic-spectroscopy experiments have revealed long-lived oscillations in two-dimensional spectra of several photosynthetic light-harvesting complexes [8–10] that pointed out to the potential existence of non-trivial quantum effects related to the interplay of the electronic and vibrational degrees of freedom [1, 2, 11–13]. In particular, the coupling between the electronic degrees of freedom and intramolecular vibrations in quasi-resonance to excitonic transitions (vibronic coupling) has been proposed as a consistent physical design principle that could explain the origin of long-lived oscillations observed in two-dimensional spectra, and possibly related to the high efficiency of the energy transfer process [14–22]. Unexpectedly, during the course of potentially being supporting long-lived oscillations, the state of the intramolecular vibrations evolve from a thermal state with non-quantum correlations into a state provided with genuinely quantum correlations even at room temperature [23].

Two-dimensional-electronic-spectroscopy is a laser-pulsed non-linear technique [24] and therefore, the extend to which their results are representative for natural conditions with continuous incoherent light sources has been intensively addressed in the literature [7, 18, 25–34]. It is by now clear that the dynamics induced by suddenly-turned-on incoherent sunlight are qualitatively different from coherent laser sources and also different from a bare white-noise-source provided that incoherent light has a super-Ohmic character and does not induce pure dephasing dynamics [31, 35]. The relevance of intramolecular vibrational modes, their impact on the energy transfer, as well as their non-trivial quantum character are explored here for the natural scenario of illumination by sunlight.

To account for that natural scenario in which chromophores harvest incoherent sunlight in the presence of localized vibrational modes, consider vibronic dimers formed by chromophores treated within the two-level approximation and a quantized intramolecular vibrational mode in interaction with each monomer. The vibronic dimers are initially set up in their electronic ground and thermal vibrational state. The excitation ignited by sunlight is then dissipated by a thermal phonon bath that accounts for the effects of the protein and solvent environments. The subsequent dynamics of the vibronic dimer are compared, in the site and exciton bases, with the dynamics of its corresponding electronic dimer with no specific intramolecular vibrational modes. As concrete examples, the two phycoerythrobilin chromophores from the protein-antenna phycoerythrin 545 and the two dihydrobiliverdin chromophores from the protein-antenna phycoerythrin 545 of marine cryptophyte algae are considered below.

Under sunlight illumination, it is found that the introduction of intramolecular vibrational modes may increase the population amplitudes in the vibronic dimers compared to the electronic ones. The amplitude of the single
exciton coherence increases up to one order of magnitude with the inclusion of the intramolecular vibrational modes, but coherence between site states is of the same order for both electronic and vibronic dimers. To account for the assistance of intramolecular vibrational modes to long-decoherence times, the decoherence rate of vibronic single exciton state superpositions is systematically analyzed in a broad regime of the dimer parameter space. It is shown that the decoherence rate displays a non-trivial behavior and that neither the nonadiabatic regime nor the values of electronic and vibronic couplings of the dynamics analyzed lead to the lowest value in the decoherence rate.

The fact that the dynamics induced by the *suddenly-turning-on* of incoherent radiation are effectively coherent, in the vibronic single exciton basis, may equivocally lead to the conclusion that the temporal coherence of the light source plays a minor role on the energy-transfer process. By focusing on the non-trivial quantum characteristics of the states of intramolecular vibrational degrees of freedom, it is ambiguously shown that under sunlight illumination conditions, the state of the two vibrational modes and that of the anticorrelated vibrational mode evolve devoid of non-classical correlations. This is in sharp contrast to the case of illumination by coherent light sources [23] and can be considered as a genuine and experimentally verifiable difference between natural and in-lab conditions, independent of the suddenly-turning-on condition.

I. LIGHT-HARVESTING ANTENNA SYSTEMS UNDER SUNLIGHT

Consider a molecular aggregate immersed within a protein that is excited by incoherent sunlight radiation. The global system-bath Hamiltonian reads $\hat{H} = \hat{H}_S + \hat{H}_{SB} + \hat{H}_B$. The light-harvesting system corresponds to a set of $N$ chromophores (sites) whose electronic degrees of freedom are coupled to quantized intramolecular vibrational modes and described by the Hamiltonian

$$\hat{H}_S = \sum_{i=1}^{N} (E_{g_i} \hat{1}_i + \epsilon_i \hat{a}_i^+ \hat{a}_i^-) + \sum_{i \neq j}^{N} V_{ij} \hat{a}_i^+ \hat{a}_j^- + \sum_{i=1}^{N} \hbar g_i \hat{a}_i^+ \hat{a}_i^- (\hat{b}_i^+ + \hat{b}_i^-) + \sum_{i=1}^{N} \hbar \kappa \hat{b}_i^+ \hat{b}_i,$$

being $E_{g_i}$ the ground state energy, $\epsilon_i$ the electronic energy of the $i$th site, $\hat{a}_i^+$ ($\hat{a}_i^-$) creates (annihilates) an electronic excitation in the $i$th site, $V_{ij}$ is the electronic coupling between the $i$th and the $j$th site. Here, $\hat{b}_i^+$ ($\hat{b}_i^-$) is the creation (annihilation) operator of the $i$th intramolecular vibrational mode of frequency $\kappa_i$, and $g_i = \sqrt{\kappa_i \kappa_i}$ represents the coupling between the $i$th excited electronic state and the $i$th intramolecular vibrational mode, and $S_i$ is the Huang-Rhys factor.

The action of the operators $\hat{a}_i^+$ and $\hat{b}_i^\dagger$ allows for defining, in an adiabatic basis, the vibronic states (i.e., electronic-vibrational states) in the electronic ground, singly excited, and doubly excited states through the relations $|g_i \nu_i\rangle = \left(\hat{b}_i^\dagger\right)^{\nu_i}/\sqrt{\nu_i!} |g_i, 0\rangle$, $|\epsilon_i \nu_i\rangle = \hat{\sigma}_i^+ \left(\hat{b}_i^\dagger\right)^{\nu_i}/\sqrt{\nu_i!} |g_i, 0\rangle$, $|f_{ij} \nu_i\rangle = \hat{\sigma}_i^+ \hat{\sigma}_j^+ \left(\hat{b}_i^\dagger\right)^{\nu_i}/\sqrt{\nu_i!} |g_i, 0\rangle$, respectively. $\nu_i$ stands for the vibrational quantum number of the $i$th site and $|g_i, 0\rangle$ for the electronic-vibrational ground state. The eigenstates of the Hamiltonian $\hat{H}_S$ correspond to the vibronic exciton states $\{|\psi_n\rangle\}$ defined by $\hat{H}_S |\psi_n\rangle = \xi_n |\psi_n\rangle$. The vibronic single exciton states are defined by $|\psi_n\rangle = \sum_{\nu_i} C_{n,\nu_i}^{|\psi_n\rangle, \nu_i}$, the localization of the $n$th vibronic single exciton state on the $k$th site is given by $l_k(\psi_n) = \langle \psi_n | \psi_n \rangle = \sum_{\nu_i} C_{n,\nu_i}^{|\psi_n\rangle, \nu_i}^2$, being $|\psi_n\rangle = \sum_{\nu_i} C_{n,\nu_i}^{|\psi_n\rangle, \nu_i} |\psi_{\nu_i}\rangle$ a vibronic single exciton state completely localized on the $k$th site. The intersite mixing ratio for a superposition between the $n$th and the $m$th vibronic single exciton states of a vibronic dimer is defined by [20] $\zeta_{nm} = \xi_n l_k(\psi_n) l_k(\psi_m) + \xi_m l_k(\psi_n) l_k(\psi_m)$. This ratio characterizes the type of coherence in the vibronic single exciton basis: $\zeta_{nm} = 1$ for a pure electronic coherence (superposition between vibronic single exciton states localized each one on different sites), and $\zeta_{nm} = 0$ for a pure vibronic coherence (superposition between vibronic single exciton states localized on the same site).

Due to the exponential scaling of the dimension of the full Hilbert space, for all simulations below, only the first four states (ground state and three excited levels) of each intramolecular vibrational mode are considered. Convergence of the density matrix time evolution was verified with the case of four excited levels. For the vibronic dimers considered here (two monomers and two intramolecular vibrations), the vibronic exciton manifold has then dimension 64: 16 vibronic ground exciton states $\{|\psi_1\rangle, \ldots, |\psi_{16}\rangle\}$, 32 vibronic single exciton states $\{|\psi_{17}\rangle, \ldots, |\psi_{48}\rangle\}$ and 16 vibronic double exciton states $\{|\psi_{49}\rangle, \ldots, |\psi_{64}\rangle\}$. The comparison with the case of an electronic dimer with no specific intramolecular vibrational modes, in the site $\{|\epsilon_1\rangle, |\epsilon_2\rangle\}$ and in the exciton basis $\{|\epsilon_1\rangle, |\epsilon_2\rangle\}$, follows after tracing over the intramolecular vibrational degrees of freedom in the density matrix of the vibronic dimer. For electronic dimers, the Frenkel Hamiltonian corresponds to the first two terms of the Hamiltonian described in Eq. (1). Thus, the two monomers have a site representation described by the states $\{|\epsilon_1\rangle, |\epsilon_2\rangle\}$ and by two single exciton states $\{|\epsilon\rangle, |\epsilon'\rangle\}$.

Specifically, the two phycoerythrobilin (PEB$_{50/61}$C and PEB$_{50/61}$D) chromophores from the protein-antenna phycoerythrin 545 (PE545), and the two dihydrobiliverdin (DBV$_{50/61}$C and DBV$_{50/61}$D) chromophores from the protein-antenna phycocyanin 645 (PC645) of marine cryptophyte algae are considered below. The PEB dimer has a large energy gap between excited electronic
states $\Delta \varepsilon = 1042 \text{ cm}^{-1}$, and due to the large spatial separation between chromophores, the electronic coupling is small ($V = 92 \text{ cm}^{-1}$); in consequence, each excitonic state is highly localized over a specific chromophore. The exciton energy splitting is $\Delta \varepsilon = 1058 \text{ cm}^{-1}$. The DBV dimer has a small energy gap between excited electronic states $\Delta \varepsilon = 73 \text{ cm}^{-1}$, and moderate electronic coupling ($V = 319.4 \text{ cm}^{-1}$) that results in the formation of delocalized exciton states with an exciton energy splitting $\Delta \varepsilon = 1042 \text{ cm}^{-1}$. Figure 1 depicts the functional dependence of $\langle \psi_a | \hat{H}_S | \psi_a \rangle$ on the ratio between the exciton energy splitting and the intramolecular vibrational frequency $\omega$, and on the electronic coupling $V$ for the first ten vibrionic single exciton states ($| \psi_1 \rangle, \ldots, | \psi_{26} \rangle$) for the PEB (top panels) and DBV (bottom panels) dimers. The vertical dashed lines indicate the conditions considered in the simulations below, and corresponds to a nonadiabatic framework [16, 22] that has been related to an enhancement of energy transfer process and the appearance of non-trivial quantum correlations driven by strong vibronic interactions [19, 23, 36].

The environment of the protein complex in Eq. (1) can be treated as a local phonon bath, whereas sunlight is formally described as blackbody radiation at 5600 K. Due to the highly mixed character of electronic-vibrational coherences, it is then necessary also to consider the coupling of the vibrational modes to the environment of the protein complex. The system-bath Hamiltonian is given by

$$\hat{H}_{SB} = \sum_{i,l}^{N, \infty} \hbar g_{il}^{(e)} \hat{\sigma}_i^+ \hat{\sigma}_l + \sum_{j}^{N} \mu_j \cdot \hat{\mathbf{E}}(t) + \sum_{i,m}^{N, \infty} \hbar g_{im}^{(v)} (\hat{b}_i^+ + \hat{b}_i) (\hat{b}_m^+ + \hat{b}_m).$$

Here, $g_{il}^{(e)}$ ($g_{im}^{(v)}$) represent the coupling between the $i^{th}$ site ($i^{th}$ intramolecular vibrational mode) and the $l^{th}$ ($m^{th}$) phonon mode. $\hat{b}_i^+$ ($\hat{b}_i$) is the creation (annihilation) operator of a $l^{th}$ phonon mode of frequency $\omega_{il}^{(i)}$ which interacts with the $i^{th}$ site. $\hat{b}_m^+$ ($\hat{b}_m$) is the creation (annihilation) operator of a $m^{th}$ phonon mode of frequency $\omega_{im}^{(i)}$ which interacts with the $i^{th}$ intramolecular vibrational mode. $\hat{\mathbf{E}}(t)$ is the dipole operator for the $i^{th}$ site and the electric field of the radiation [37] is given by $\hat{\mathbf{E}}(t) = \mathbf{E}^{(+)}(t) + \mathbf{E}^{(-)}(t)$, with $\mathbf{E}^{(+)}(t) = i \sum_{k,s} \left( \frac{\hbar \omega}{2 e \nu V} \right)^{1/2} \hat{a}_{k,s} \mathcal{E}(k,s) e^{-i \omega t}$ and $\mathbf{E}^{(-)}(t) = \left[ \mathbf{E}^{(+)}(t) \right]^{\dagger}$, and $\hat{a}_{k,s}^\dagger (\hat{a}_{k,s})$ being the creation (annihilation) operator for the $k^{th}$ radiation field mode in the $s^{th}$ polarization state.

### A. System dynamics

The effect of the incoherent radiation environment (blackbody bath) and the vibrational environment (phonon bath) is encoded in the spectral densities $J_{il}^{(e)}(\omega)$ and $J_{im}^{(v)}(\omega)$, respectively. The blackbody radiation bath of frequency $\omega$...
is characterized by the super-Ohmic spectral density [31]

$$\omega^2 j_{\text{phon}}^{(\nu)}(\omega) = \frac{2\hbar \omega^3}{3(4e0\pi^2 c^3)}.$$  \hspace{0.5cm} (4)

This spectral density generates long-lasting coherent dynamics (see Fig. 2, \(\Lambda^{(e,v)} = 0\) case) provided by the lack of pure dephasing dynamics and the strong dependence of the decoherence rate on the system level spacing [31] (see Fig. 3 A and B). The spectral density of the phonon-baths reads

$$\omega^2 j_{\text{j}}^{(\nu)}(\omega) = \frac{2\Omega_j^{(e,v)} \Lambda_j^{(e,v)} \omega}{\hbar(\omega^2 + \Omega_j^{(e,v)} \omega^2)},$$  \hspace{0.5cm} (5)

where \(\Omega_j^{(e,v)}\) represents the cutoff frequency and \(\Lambda_j^{(e,v)}\) the reorganization energy of the phonon baths coupled to the electronic (e) and intramolecular vibrational (v) degrees of freedom. The dynamics of light-harvesting systems with spectral densities of the form (5) are commonly solve with the hierarchical equations of motion (HEOM) method [38–40]. However, this method cannot be applied to super-ohmic spectral densities. To circumvent this, the HEOM method has been used to treat the phonon bath whereas the non-unitary effect of the incoherent light has been accounted for by a Lindblad dissipator [41, 42]. However, this hybrid approach does not properly get into account the influence of the super-Ohmic spectral density in the density matrix dynamics since it does not consider the dependence of the decoherence rates on the system level spacing. To adequately describe the correlations induced by the super-Ohmic spectral density of the blackbody radiation together with the phonon bath effects, the dynamics are solved in the vibronic exciton basis \(\{\psi_n\}\), i.e., in the eigenstates of the Hamiltonian in Eq. (1), by using the standard Redfield master equation (second-order and non-secular) [31, 43]. Recent works on energy transfer dynamics of vibronic dimers excited with coherent light have considered the Redfield approach and have shown similar results to the HEOM method under parameters used in experimental conditions [44–46]. Further details about both methods and comparison between them are presented in SI, Sec. 1 and 2.

II. DYNAMICS IN THE PRESENCE OF BLACKBODY RADIATION AND PHONON BATHS

Energy transfer starts with the rapid incoherent excitation of the electronic sites in their electronic ground states with transition dipole moments of 11.87 D (PEB50/61 D), 12.17 D (PEB50/61 c), 13.1 D (DBV50/61 D) and 13.2 D (DBV50/61 c). Initially, each chromophore is in its ground electronic state and the intramolecular vibrational modes are in thermal equilibrium at \(T = 300\) K. After the dynamics begin, the vibronic dimer remains coupled to the incident blackbody radiation [31, 35]; this is in sharp contrast to the pulsed laser excitation conditions [7]. The frequency of the two intramolecular vibrational modes is in full resonance with the exciton splitting, i.e., \(\omega_1 = \omega_2 = \Delta\varepsilon\). The vibronic coupling strength to each monomer is the same, i.e., \(g_1 = g_2 = g\), specifically, \(g = 267.1\) cm\(^{-1}\) for the PEB dimer and \(g = 250\) cm\(^{-1}\) for the DBV dimer. For the simulations below, same spectral densities on each monomer are taken, with \(\Omega_j^{(e)} = 100\) cm\(^{-1}\) and for various values of the reorganization energy \(\Lambda_j^{(e)}\). Besides, same spectral densities on each intramolecular vibrational mode are taken, with \(\Omega_j^{(v)} = 50\) cm\(^{-1}\) and reorganization energy \(\Lambda_j^{(v)} = 10\) cm\(^{-1}\) provided that \(\Lambda_j^{(e)} \neq 0\).

A. Vibronic single exciton basis

Consider first the dynamics of the vibronic dimer in interaction with blackbody radiation only, i.e., set to zero the non-unitary effects related to the phonon baths and assume that the system and the blackbody radiation are initially decoupled, i.e., \(\hat{\rho}(t_0) = \hat{\rho}_S(t_0) \otimes \hat{\rho}_{BB}(t_0)\). The transition dipole moments are considered parallel to the incident electric field and constant in time so that the effect of different orientations of the transition dipole moments is neglected.
The linear increase of the populations is expected in panels), Λ Green points represent the values adopted for the simulations of the density matrix dynamics in the vibronic single exciton,herent superpositions arising between them in the PEB ∼ences (coherences depicted in Fig. 3. Decoherence rate γ_{18,20} [s^{-1}] (color map) for the PEB dimer with the reorganization energies Λ^{(e,v)} = 0 cm^{-1} (top panels), Λ^{(e)} = 10 cm^{-1}, Λ^{(v)} = 10 cm^{-1} (middle panels) and Λ^{(e)} = 100 cm^{-1}, Λ^{(v)} = 10 cm^{-1} (bottom panels), as a function of the ratios g/Δε, π/Δε, V/Δε and the transition dipole moment amplitude D, where Δε represents the site energy difference. Green points represent the values adopted for the simulations of the density matrix dynamics in the vibronic single exciton, exciton and sites bases, discussed throughout the paper. Baths parameters are T_{vb}^{(e,v)} = 300 K, T_{bb} = 5600 K.

Figure 2 (Λ^{(e,v)} = 0 case) shows the populations (> 10^{-6}) of the vibronic single exciton states and the coherent superpositions arising between them in the PEB dimer after suddenly-turning-on the incoherent radiation. The linear increase of the populations is expected in low-intensity incoherent radiation [31]. In chromophores isolated from the vibrational phonon environment, suddenly turned-on incoherent-light-induced-dynamics are effectively coherent and last for hundreds of picoseconds [31]. Nevertheless, the amplitude of the vibronic coherences (∼ 10^{-8}) is approximately two orders of magnitude smaller than the populations; hence, they turn out to quickly become irrelevant for the dynamics of populations [28]. Most of these vibronic coherences display a highly mixed electronic-vibrational character, quantified through the intersite mixing ratio. Specifically, for the coherences depicted in Fig. 2, ζ_{18,19} = 0.52, ζ_{18,20} = 0.50 and ζ_{19,20} = 0.48. Therefore, vibronic coherence dynam-ics are influenced by the decoherence and dissipation of the electronic as well as intramolecular degrees of freedom. This is the reason for introducing a thermal bath of each intramolecular vibrational mode in Eq. (3).

To incorporate the effect of the phonon bath (Λ^{(e,v)} ≠ 0), assume that it is initially decoupled from the dimers \( \hat{\rho}(t_0) = \hat{\rho}_S(t_0) \otimes \hat{\rho}_{PB}(t_0) \otimes \hat{\rho}_{PB}(t_0) \). Figure 2 depicts the populations of vibronic single exciton states for different values of the reorganization energy Λ^{(e)} = 10, 30, 100 cm^{-1}. Specifically, the population of the lowest energy vibronic single exciton state \( |\psi_{17} \rangle \) of the PEB dimer increases for increasing values of the reorganization energy. In the case of the DBV dimer, there is an increase of two orders of magnitude in the population of the state \( |\psi_{17} \rangle \), see SI, Fig. S2. This is a consequence of the intricate interplay between bath-enhanced rates and non-adiabatic dynamics of vibronic single exciton states with small energy gaps. Figure 2 also depicts the dynamics of...
superpositions between vibronic single exciton states (vibronic coherences). The vibronic coherences, originated by the turning-on of the incoherent radiation, decay due to the interaction with the phonon bath. Their influence on the population of the vibronic single exciton states is negligible, owing to the amplitude of the vibronic coherences is approximately one (PEB dimer) and two (DBV dimer) orders of magnitude smaller than the populations of vibronic single exciton states.

To explore the role of the vibronic coupling and the incoherent light excitation process on the lifetime of quantum superpositions between vibronic single exciton states, the decoherence rate of the highest amplitude vibronic coherence $\mathcal{R}_B(t)$ in the PEB dimer is analyzed (see Fig. 2). Figure 3 depicts the functional dependence of the decoherence rate $\gamma_{18,20}$ on the ratio between the vibronic coupling and the site energy difference $g/\Delta e$, the ratio between the intramolecular vibrational frequency and the site energy difference $\omega/\Delta e$, the ratio between the electronic coupling and the site energy difference $V/\Delta e$, and the dipole moment amplitude $D$. The green points in Fig. 3 depict the specific values for the PEB dimer, $g/\Delta e = 0.26$, $\omega/\Delta e = 1.02$, $V_1/\Delta e = 0.09$ and $D = 1$ (For the DBV dimer case, see SI, Fig. S3).

In absence of thermal baths for the intra-molecular vibrational modes (not shown), the decoherence rate $\gamma_{18,20}$ decreases for increasing values of the vibronic coupling $g$, however, in the more realistic scenario depicted in Fig. 3, increasing the vibronic coupling may lead to regions of parameter space with higher decoherence rates. Thus, the decoherence rate $\gamma_{18,20}$ displays a non-trivial behavior under the variation of the physical quantities defined above and neither the vibronic resonance condition $\omega_1 = \omega_2 = \Delta e$ (i.e., nonadiabatic regime) nor the values of electronic and vibronic couplings of the dynamics discussed (see green points in Fig. 3) lead to the lowest value in the decoherence rate. Thus, the longest decoherence time for vibronic single exciton state superpositions is not reached under the physical conditions considered usually in two-dimensional-electronic-spectroscopy studies [22, 47].

For natural light-matter coupling strengths (Figs. 3 A, B, D, E, G and H): (i) increasing the reorganization energy increases the decoherence rate $\gamma_{18,20}$ for $\Lambda(e) = 0, 10, 100$ [cm$^{-1}$] as $\gamma_{18,20} \sim 10^4, 10^{13}, 10^{15}$ [s$^{-1}$], respectively. (ii) For $\Lambda(e) = 0$, the decoherence rate $\gamma_{18,20}$ is at least five orders of magnitude smaller than for cases with $\Lambda(e) \neq 0$. This follows from the low intensity of sunlight and the energy-gap dependence of blackbody radiation decoherence rates $\gamma_B^{\Lambda(e)} \sim (\mu_{e,e}'^2 \omega_{e,e}'^2 / 3h \pi \epsilon_0 \epsilon_r^3) \cosh(\hbar \omega_{e,e}' / 2 k_B T_B^{\Lambda(e)})$. Thus, for small energy gaps $\hbar \omega_{e,e}'$, the decoherence rate $\gamma_B^{\Lambda(e)}$ may be considerably smaller than the case of an Ohmic thermal phonon bath $\gamma_B^{\Lambda(e)} \sim 4 k_B T^{\Lambda(e)} / h^2 \Omega$, which is energy-gap independent [31]. For the values of the reorganization energy considered in Fig. 3, the increase of the transition dipole moment amplitude leads to higher values in the decoherence rate $\gamma_{18,20}$ (see Fig. 3 C, F and I). Increasing of the vibronic coupling $g$ in Fig. 3 for the values of the reorganization energy considered does not lead to decrease of the decoherence rate.

**B. Reduced exciton and site bases**

The role of the high frequency intramolecular vibrational modes is explored by comparing the reduced electronic dynamics (tracing over the intramolecular vibrations) of the vibronic dimers considered above with their corresponding electronic dimer dynamics (no specific intramolecular vibrational modes). Figures 4 and 5 show the populations and coherences in the exciton (top panels) and site bases (bottom panels), with (vibronic dimer and without (electronic dimer) intramolecular vibrational modes for the PEB and DBV dimers, respectively.

In the exciton basis, and in the absence of the phonon baths ($\Lambda(e,v) = 0$ [cm$^{-1}$]), the population of the lowest energy exciton state of the vibronic dimer is higher than that of its corresponding electronic dimer case: one and half times for the PEB dimer (see Fig. 4 A), and two orders of magnitude for the DBV dimer (see Fig. 5 A). The amplitude of the coherence between single exciton states in the vibronic dimer model (see Fig. 4 C) is one order of magnitude higher than in the electronic dimer case (see Fig. 4 D) for the PEB dimer, and two orders of magnitude in the case of DBV dimer (see Fig. 5 C and D). The increase in population and coherence can be understood as the result of the smaller energy gaps [1, 12] induced by intramolecular vibrations, i.e., a consequence of the non-adiabatic character of the dynamics (see Fig. 1). Hence, for $\Lambda(e,v) = 0$, the population of the lowest energy exciton state and the coherence between single exciton states increase with the inclusion of intramolecular vibrational modes.

In the presence of the phonon bath ($\Lambda(e,v) \neq 0$), the behavior of populations are similar to $\Lambda(e,v) = 0$. The amplitude of the coherences increases slightly in the PEB dimer (see Fig. 4 C and D) and by up to one order of magnitude in the DBV dimer (see Fig. 5 C and D). The population of the lowest energy exciton state of the PEB dimer have higher amplitudes in the vibronic dimer than in the electronic dimer; however, as the value of the reorganization increases, this population difference decreases. Moreover, the population rate of the reduced vibronic dimer barely changes by increasing the reorganization energy, thus seeing to be robust against the fluctuations of the phonon environment (see Figs. 4 A and 5 A). For small values of the reorganization energy ($\Lambda(e) \sim 10$cm$^{-1}$), a similar population trend is found for the DBV dimer; however, for moderate ($\Lambda(e) \sim 30$cm$^{-1}$) and large values ($\Lambda(e) \sim 100$cm$^{-1}$) of the reorganization energy, the population of the lowest energy exciton state has slightly higher amplitudes in the case of the electronic dimer than the vibronic dimer (see Fig. 5 A). This clearly shows the highly non-trivial interplay between bath-enhanced population rates and
non-adiabatic dynamics for highly localized (PEB) and highly delocalized (DBV) excitons.

In the basis, and for all values of the reorganization energy considered, the population of the chromophores PEB_{50/61C} and DBV_{50/61C} is higher than that of PEB_{50/61D} and DBV_{50/61D}, respectively, in the case of the vibronic dimer than in the electronic dimer case (see Fig. 4 E and F, and Fig. 5 E and F). Since population is pumped from the ground state and the transition dipole moments favor excitation to singly excited exciton states with low energy, the population difference occurs due the localization l_{\psi} of the vibronic single exciton states on those chromophores; specifically, l_{\psi}(\psi_{17}) = 0.007, l_{\psi}(\psi_{18}) = 0.007, l_{\psi}(\psi_{17}) = 0.993, l_{\psi}(\psi_{18}) = 0.524, l_{\psi}(\psi_{18}) = 0.476, l_{\psi}(\psi_{19}) = 0.007, l_{\psi}(\psi_{19}) = 0.993, and l_{\psi}(\psi_{20}) = 0.475, l_{\psi}(\psi_{20}) =
depicts the occupation number distribution of a coherent exciton-vibration interactions, whereas PEB and DBV dimers develop a genuinely non-classical character due to coherent exciton-vibration interactions. The quantized vibrational modes do not display non-classical correlations quantified by the Mandel parameter. The Mandel parameter identifies the non-classical character of bosonic states through the comparison of occupation number distribution for a given bosonic state with the occupation number distribution of a coherent state. The Mandel parameter is given by $Q = \frac{\langle \hat{n}^2 \rangle - \langle \hat{n} \rangle^2}{\langle \hat{n} \rangle} - 1$, where $\hat{n}$ is the occupation number operator of the bosonic state. For the case of a coherent state the occupation number distribution corresponds to a Poisson distribution ($Q = 0$). Thus, for any occupation number distribution narrower than a Poisson distribution, i.e., with $Q < 0$, the associated bosonic state has a quantum character with no classical analog.

For different values of the reorganization energy and under sunlight illumination conditions, Fig. 6 depicts the Mandel parameter for the two vibrational modes of frequencies $\omega_1$ and $\omega_2$ considered in the vibronic model of the PEB and DBV dimers. The reduced dynamics of the anticoherent vibrational mode of frequency $\omega_{ac}$, previously analyzed in the seminal work by Jonas et al. [16] and responsible for the non-adiabatic character of the dynamics (see Fig. 1), is also considered. For every case considered, the Mandel parameter adopted positive values, which indicates that during the dynamics, the state of the intramolecular vibrations modes has a classical character. Even, in the case of absence of phonon bath ($\Lambda^{(e,v)} = 0$), the value of the Mandel parameter remains positive.

Under sunlight illumination conditions, dimers are initially in their the electronic ground state while intramolecular vibrations, that are decoupled from the ground state, are initially at thermal equilibrium; therefore, the initial condition is devoid of quantum superpositions. Under pulsed-laser-excitation conditions, vibrations are also assumed to be at thermal equilibrium; however, in sharp contrast to natural conditions, the dimer is assumed to have been prepared in, e.g., an exciton state. Therefore, under pulsed-laser-excitation conditions, the dimer is initially prepared in a coherent superposition of vibronic exciton states provided that the chromophore-chromophore dipole interaction is finite.

For vanishing chromophore-chromophore dipole interaction...
action, the electronic and vibrational contributions to the vibronic Hamiltonian in Eq. (1) commute; thus indicating that a product state of electronic and vibrational single eigenstates will also be an eigenstate— not a coherent superposition of eigenstates of the vibronic Hamiltonian. The fact that for this product state the Mandel parameter adopts only positive values led to conclude [23] that the transient formation of vibronic exciton states establishes non-classical correlations in the vibrational modes. However, the chromophore-chromophore dipole interaction is finite under sunlight illumination but not non-classical correlations are established provided the lack quantum correlations in the initial state. Therefore, non-classical correlations does not emerge due to the transient formation of vibronic exciton states, but as a consequence of the initial quantum correlations established in the light-harvesting system by the pulsed-laser-preparation of the initial state.

### IV. SUMMARY

The role of intramolecular vibrations resonant with excitonic transitions in light-harvesting systems was analyzed under natural sunlight illumination. The comprehensive analysis shows that the populations of single exciton and site states of vibronic dimers are not significantly affected as compared to their corresponding electronic dimers. Therefore, there is no direct evidence of an enhancement in the energy transport mediated by the inclusion of resonant intramolecular vibrational degrees of freedom under natural conditions. Recently, similar conclusions were elucidated about the impact of the vibronic coupling in the electronic and vibrational coherences observed in two-dimensional-electronic-spectroscopies [47]. However, the inclusion of the intramolecular vibrational modes reinforces the exciton coherence by up to one order of magnitude, as was shown for the DBV dimer.

Under incoherent light excitation conditions, it was further shown that intramolecular vibrational modes do not develop non-classical correlations provided that the initial state is devoid of quantum coherence. Therefore, the generation of non-classical correlations via the transient formation of vibronic exciton states lacks of theoretical support and should be replaced in favor of the natural dynamics of the initial quantum correlations established in the light-harvesting system by the pulsed-laser-preparation of the initial state.

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