Exact simulation of pigment-protein complexes unveils vibronic renormalization of electronic parameters in ultrafast spectroscopy

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The primary steps of photosynthesis rely on the generation, transport, and trapping of excitons in pigment-protein complexes (PPCs). Generically, PPCs possess highly structured vibrational spectra, combining many discrete intra-pigment modes and a quasi-continuous of protein modes, with vibrational and electronic couplings of comparable strength. The intricacy of the resulting vibronic dynamics poses significant challenges in establishing a quantitative connection between spectroscopic data and underlying microscopic models. Here we show how to address this challenge using numerically exact simulation methods by considering two model systems, namely the water-soluble chlorophyll-binding protein of cauliflower and the special pair of bacterial reaction centers. We demonstrate that the inclusion of the full multi-mode vibronic dynamics in numerical calculations of linear spectra leads to systematic and quantitatively significant corrections to electronic parameter estimation. These multi-mode vibronic effects are shown to be relevant in the longstanding discussion regarding the origin of long-lived oscillations in multidimensional nonlinear spectra.
Light-harvesting (LH) antennas and photo-chemical reaction centers (RC) provide the elementary building blocks of the photosynthetic apparatus of plants, algae, and bacteria. Primarily these molecular aggregates consist of absorbing molecules (pigments) complexed with specific proteins to form a PPC. Despite its fundamental importance to biology, the dynamical characterization of these complexes to a degree that can reproduce all reported spectroscopic data in a single microscopic model remains an outstanding challenge.

Reduced models of excitonic dynamics subject to purely thermal fluctuations can achieve reasonable agreement with linear optical spectra. The quantitative explanation of all relevant aspects of multi-dimensional nonlinear spectroscopy though requires a more detailed model of the system-environment interaction that takes into account the full complexity of the environmental structure. Indeed, spectroscopic studies of PPCs at low temperatures reveal the presence of vibrational environments that consist of a broad spectrum of low-frequency modes with room temperature energy scales, and several tens of discrete high-frequency modes that originate mainly from intra-pigment dynamics. Nonlinear optical experiments on monomer pigments in solution at both 77 K and room temperature, as well as first-principles calculations further corroborate the underdamped nature of intra-pigment vibrational modes with picosecond lifetimes.

Recently, a range of vibronic models in which pigments are subject to the combined influence of a broad unstructured bosonic environment and a small number of vibrational modes with frequencies in the vicinity of excitonic transitions have been formulated. In this picture, vibrational lifetime borrowing can lead to long-lasting oscillatory dynamics of coherences between excitonic states, and observations of long-lasting oscillatory features in multi-dimensional spectroscopy have been attributed to this effect. Notwithstanding, the identification of a universally accepted origin of these long-lived oscillations remains a subject of active discussion.

An important obstacle that prevents the conclusive resolution of this debate is the fact that the interpretation of spectroscopic data and their underpinning dynamical features can be influenced significantly by the specific choice of electronic and vibrational parameters that enter the PPC models. We will demonstrate that by accounting for the full environmental spectral density, involving more than 50 intra-pigment modes per site in addition to a broad background, the presence of high-frequency long-lived vibrational modes can lead to quantitatively significant renormalization of the resulting excitons to vibrational degrees of freedom of the pigment’s environment. In the following, we will restrict our analysis to the transition between electronic ground and first excited states of the pigments, which suffices for the evaluation of the low-energy part of absorption spectra and is relevant for photosynthetic energy transfer.

Results

Electronic and vibronic couplings of PPCs. Absorption spectra of PPCs are determined by the electronic energy-level structure of pigments, their mutual electronic interactions and the coupling of the resulting excitons to vibrational degrees of freedom of the pigment’s environment. In the following, we will restrict our analysis to the Q transition between electronic ground and first excited states of the pigments, which suffices for the evaluation of the low-energy part of absorption spectra and is relevant for photosynthetic energy transfer.

For the dimeric WSCP and SP, the electronic Hamiltonian is then described by (see Supplementary Note 1)

\[ H_e = \sum_{i=1}^{2} \epsilon_i |\epsilon_i\rangle \langle \epsilon_i| + V(|\epsilon_1\rangle \langle \epsilon_2| + |\epsilon_2\rangle \langle \epsilon_1|). \]  

Here \( |\epsilon_i\rangle \) denotes the singly excited state of site \( i \) with on-site energy \( \epsilon_i \) that is in the visible (WSCP) or in the near infrared spectrum (SP). The on-site energies depend on their local environment and therefore suffer from static disorder inducing ensemble dephasing that will be included in our numerical treatment. The electronic coupling \( V \) leads to delocalized electronic eigenstates (excitons), \( H_e |E_k\rangle = E_k |E_k\rangle \), and an excitonic splitting \( \Delta = E_+ - E_- = \sqrt{4V^2 + (\epsilon_1 - \epsilon_2)^2} \). In WSCP, the mean site energies are identical, \( \langle \epsilon_1 \rangle = \langle \epsilon_2 \rangle \), due to the symmetry
of molecular structure, while in SP, the mean site energies are different as pigments are surrounded by nonidentical local protein environments. Another difference concerns the electronic coupling strength, which is stronger in SP due to electron exchange giving rise to short-range Dexter type contributions 56,57.

The exciton dynamics of PPCs is driven by vibrational modes that influence fluctuations in the transition energies $\epsilon_i$ of pigments. The full electronic-vibrational interaction, induced by $N$ vibrational modes per site, is described by the Hamiltonian

$$H = H_v + H_h + H_{ve}$$

where

$$H_v = \sum_{i=1}^{N} \sum_{k=1}^{\omega_k} \omega_k b_{i,k}^+ b_{i,k},$$

$$H_{ve} = \sum_{i=1}^{N} \left( \epsilon_i - \sum_{k=1}^{\omega_k} \omega_k \sqrt{\gamma_k} (b_{i,k} + b_{i,k}^+) \right).$$

Here the annihilation (creation) operator $b_{i,k}$ ($b_{i,k}^+$) describes a local vibrational mode of frequency $\omega_k$ coupled to site $i$ with a strength quantified by the Huang-Rhys (HR) factor $\gamma_k$. For an environment initially in a thermal state, the ensuing dynamics is fully determined by the environmental spectral density $J(\omega) = \sum_k \omega_k b_{i,k}^+ b_{i,k}$ whose structure needs to be determined experimentally or theoretically.

**Structure of the environmental spectral density.** Generally, in PPCs the spectral density $J(\omega)$ consists of a broad background and multiple sharp peaks distributed across a broad range of frequencies. These can be determined by fluorescence line-narrowing (FLN) and hole burning experiments which reveal that the environmental spectral densities of WSCP and SP consist of low-frequency broad features originating from protein motions, and 55 intra-pigment modes resulting in multiple narrow peaks in the high-frequency part of the spectrum. The contribution of the protein modes of WSCP may be described by log-normal distribution functions of the form

$$J_p(\omega) = \frac{1}{\sqrt{2\pi} \sigma_p} \exp\left(-\ln(\omega/\Omega_p)^2/2\sigma_p^2\right),$$

which provides a satisfactory description of the low-energy part of experimentally measured FLN spectra of WSCP 58. Alternatively, the protein motions of WSCP have been modeled by the following functional form:

$$J_p(\omega) = \sum_{i=1}^{3} \sum_{j=1}^{2} \omega_i^2 e^{-\omega_i^2/(\omega_j)^2}$$

that has been extracted from FLN spectra of B777 photosynthetic complexes 59 and considered in the simulations of WSCP 58. Every underdamped intra-pigment mode contributes a Lorentzian of width $\gamma_k \sim 1 \text{ ps}^{-1}$, resulting in $\Delta = J_h(\omega) + J_p(\omega)$ where

$$\Delta = \sum_{i=1}^{N} \left( \epsilon_i - \sum_{k=1}^{\omega_k} \omega_k \sqrt{\gamma_k} (b_{i,k} + b_{i,k}^+) \right),$$

and the reorganization energy of the high-frequency modes is given by

$$\lambda_h = \sum_{k=1}^{\omega_k} \omega_k \sqrt{\gamma_k}.$$
an electronic coupling strength estimate of $V \approx 70 \text{ cm}^{-1}$, as shown in red in Fig. 2b. Such an electronic 
 coupling results in an excitonic splitting $\Delta \approx 2V \approx 140 \text{ cm}^{-1}$ which is consistent with 
 the experimentally observed energy-gap between two absorption 
 peaks at 656 and 662 nm, respectively. Since all the high 
 frequency intra-pigment modes are neglected in the coarse-grained 
 spectral density and the energy-gap between absorption peaks 
 is smaller than the vibrational frequencies of the intra-pigment 
 modes ($\Delta < \omega_k$), the estimated value could be interpreted as the 
 effective coupling $V_{00}$ between $|e_1, 0\rangle$ and $|e_2, 0\rangle$ where $|0\rangle$ denotes the common 
 vibrational ground state of the intra-pigment modes in the electronic excited state manifold. 
 As shown in Fig. 3a, the transition dipole strength between $|g, 0\rangle$ and $|e_1, 0\rangle$ 
 (0-0 transition) of a monomer is reduced by a factor of 
 $\exp(-\sum k \omega_k / 2)$, as the total transition dipole strength of the monomer is redistributed to 
 0-1 transitions between $|g, 0\rangle$ and $|e_1, 1\rangle$ where only the $k$-th mode is singly excited (see 
 Supplementary Note 10). As a result, the effective coupling between 0-0 transitions, shown in Fig. 3b, 
 is reduced to $V_{00} = V \exp(-\sum k \omega_k)$ depending on the HR factors $s_k$ of the intra-pigment modes. 
 This implies that $V_{00} \approx 70 \text{ cm}^{-1}$ corresponds to a bare electronic coupling 
 $V = V_{00} \exp(\sum k \omega_k) \approx 2V_{00} \approx 140 \text{ cm}^{-1}$ under the full 
 environmental spectral density $P_{\text{exp}}(\omega)$, including the 55 
 intra-pigment modes shown in black in Fig. 2a. The renormalised 
 electronic coupling $V \approx 140 \text{ cm}^{-1}$ yields a best fit to experimentally 
 measured absorption spectra, as shown in black in Fig. 2c, when all the 
 $M = 55$ intra-pigment modes are considered in simulations. 
 The energy-gap between absorption peaks is gradually reduced 
 from an excitonic splitting $\Delta = 2V \approx 280 \text{ cm}^{-1}$ to 
 $\Delta' \approx 2V_{00} \approx 140 \text{ cm}^{-1}$, as the number $M$ of the lowest-frequency 
 intra-pigment modes considered in simulations is increased from 
 20 via 40 to 55 (see Fig. 2a, c). The electronic coupling $V \approx 70 \text{ cm}^{-1}$ 
 estimated based on the coarse-grained low-frequency spectral 
 density cannot reproduce the experimental results when the full 
 spectral density is considered in simulations, as shown in Fig. 2d. 
 The energy-gap between absorption peaks shown in Fig. 2c, d can 
 be quantitatively well described by the splitting of 0-0 transitions, 
 $2V_{00} \approx 2V \exp(-\sum k \omega_k)$, implying that the effective couplings 
 $V_{01}$ between 0-0 and 0-1 transitions, schematically shown in 
 Fig. 3b, are not strong enough to modify the energy-gap between 
 low-energy absorption peaks of WSCP. However, the weak $V_{01}$ 
 couplings can redistribute the transition dipole strength from 0-0 
 to 0-1 transitions and significantly modify the high-energy part 
 of absorption spectra, which cannot be described by conventional line 
 shape theory (see Supplementary Note 10).

**Multi-mode vibronic mixing in exciton basis.** In contrast to 
 WSCP, the bare excitonic splitting of SP is of the order of the 
 typical vibrational frequencies of the intra-pigment modes and 
 the resulting redistribution of oscillator strengths and shifts of 
 optical lines are much more difficult to predict. To qualitatively 
 estimate these effects, we consider second-order perturbation 
 theory starting from the full Hamiltonian $H = H_0 + H_1 + H_{\text{exc}}$ 
 in the single-exciton manifold. In that case, the vibronic mixing 
 is induced by the relative motion of the intra-pigment modes with 
 identical frequency $\omega_k$, described by $b_k = (b_{1,k} - b_{2,k})/\sqrt{2}$, as the 
 center of mass motion, described by $B_k = (b_{1,k} + b_{2,k})/\sqrt{2}$, 
 merely induces the homogeneous broadening of absorption line 
 shapes without affecting exciton dynamics (see Supplementary 
 Note 1). Hence, we can discard the center-of-mass part of the 
 total Hamiltonian to find $H = H_0 + H_1$ where

$$H_0 = H_1 + H_{\text{exc}} + \cos(\theta) \sigma_z \sum_{k=1}^{55} \omega_k \sqrt{s_k/2} (b_k + b_k^*) ,$$

$$H_1 = -\sin(\theta) \sigma_y \sum_{k=1}^{55} \omega_k \sqrt{s_k/2} (b_k + b_k^*) .$$

Here $\theta = \tan^{-1}[2V/(\varepsilon_1 - \varepsilon_2)]$, while $\sigma_+ = |E_+\rangle \langle E_+| + |E_-\rangle \langle E_-|$ and 
 $\sigma_- = |E_+\rangle \langle E_-| - |E_-\rangle \langle E_+|$ are the Pauli matrices in the exciton basis. 
 The Hamiltonian $H_0$ is diagonalised by the polaron 
 transformation using second order perturbation theory 
 starting from the full Hamiltonian $H_{\text{exc}} = H_0 + H_{\text{exc}}$, of the order of 
 $s_k \approx 0.01$, the vibronic mixing is dominated by contributions from the single 
 excitation subspace where it leads to eigenstates of $H_{\text{exc}}$ of the form

$$\psi_{\pm} = a_{\pm,0} |E_+, 0\rangle + \sum_{k=1}^{55} a_{\pm, s_k} |E_+, s_k\rangle ,$$
\( \Delta k \) so that they cannot simply be ignored on the basis of being off-resonant.

When the excitonic splitting is larger than the vibrational frequencies, \( \Delta > \omega_k \), the situation is reversed (see Fig. 4b), resulting in an increased vibronic splitting \( \Delta' \) compared to the bare excitonic splitting \( \Delta \). This case cannot be described by the splitting of 0-0 transitions, since the effective coupling \( V_{00} = V \exp(-\sum_k \beta_k) \) is smaller in magnitude than a bare electronic coupling \( V \) for arbitrary HR factors defined by \( \beta_k \geq 0 \). This implies that the mixing of 0-0 and 0-1 transitions can result in two absorption peaks with an energy gap \( \Delta' \) being larger than the bare excitonic splitting \( \Delta \).

**Special pair in bacterial reaction center** The photosynthetic reaction center which drives exciton dissociation into free charges consists of the SP and four additional pigments\(^5^3\). The SP is a strongly coupled dimeric unit with an electronic coupling estimated to be \( V = 625 \text{ cm}^{-1} \), a difference in mean site energies of \( \langle \epsilon_1 - \epsilon_2 \rangle = 315 \text{ cm}^{-1} \) and consequently a bare excitonic splitting of \( \Delta \approx 1290 \text{ cm}^{-1} \). These electronic parameters have been estimated based on a best fit to absorption, linear dichroism, and hole burning spectra of bacterial reaction centers using conventional line shape theory\(^5^3\). In what follows, we neglect the order of magnitude weaker electronic coupling of the SP to the four additional pigments and do not aim to reproduce experimentally measured absorption spectra of the whole bacterial reaction centers and re-estimate electronic parameters. Rather we concentrate on the effect of multi-mode vibronic mixing on the SP and its consequences regarding the nature and lifetimes of excitonic coherence and long-lived oscillatory signals in 2D electronic spectra.

While in WSCP the excitonic splitting is far detuned from high-frequency modes, the situation is markedly different for the SP. Here the environmental spectral density contains high-frequency intra-pigment modes both above and below the bare excitonic gap, as shown in black in Fig. 5. The smaller frequency differences between vibrational modes and excitonic splitting and the varying sign of their detuning makes the effect of multimode mixing harder to predict analytically. Indeed, the perturbation procedure for obtaining Eq. (9) will be inaccurate for a larger number of modes. The vibronic splitting can be estimated beyond the perturbation theory by numerically diagonalising the Hamiltonian \( H = H_0 + H_I \) in Eqs. (5), (6), leading to \( \Delta' \approx 1744 \text{ cm}^{-1} \) (see Supplementary Note 7). This estimate is in line with numerically exact simulated results where the energy-gap between absorption peaks is approximately \( 1710 \text{ cm}^{-1} \) (see 780 and 900 nm peaks in Fig. 5b, corresponding to \( |\psi_+\rangle \) and \( |\psi_-\rangle \), respectively) and the oscillatory dynamics of excitonic coherence is dominated by \( 1755 \text{ cm}^{-1} \) frequency component (see Fig. 5c). We note that the difference between excitonic and vibronic splittings is significant, of the order of \( \Delta' - \Delta \approx 465 \text{ cm}^{-1} \), and this shift cannot be described by conventional line shape theory where multi-mode vibronic mixing is ignored and as a result the
energy-gap between absorption peaks is reduced to the excitonic splitting (see the inset in Fig. 5b).

**Long-lived multi-mode vibronic coherence.** The considerable size of the multi-mode mixing effects on excitonic energy gaps suggest a possibly significant influence on coherent excitonic dynamics. The coarse-grained spectral density shown in red in Fig. 5a, which corresponds to a vibrational lifetime of $\gamma_k = (50 \text{ fs})^{-1}$, yields short-lived oscillator strengths of excitonic coherence $\rho_{\text{exc}}(t) = \langle \hat{E}_- | \hat{\rho}_k(t) | \hat{E}_+ \rangle$ with $\hat{\rho}_k(t)$ denoting reduced electronic density matrix (see red line in Fig. 5c). Even if a few intra-pigment modes near-resonant with excitonic splitting are selected to be weakly damped, $\gamma_k = (1 \text{ ps})^{-1}$, the vibronic mixing with the large number of remaining strongly-damped modes, $\gamma_k = (50 \text{ fs})^{-1}$, suppresses the lifetime of excitonic coherences, making the resulting dynamics essentially identical to that where all the modes are strongly damped (see Supplementary Note 7 for detailed analysis of multi-mode vibronic mixing). In sharp contrast, when the picosecond lifetime of actual intra-pigment modes is considered, $\gamma_k = (1 \text{ ps})^{-1}$, the excitonic coherence dynamics is dominated by long-lived oscillations with frequency $\Delta_k \approx 1755 \text{ cm}^{-1}$, associated with the vibronic coherence between $|\psi_+\rangle$ and $|\psi_-\rangle$ states (see black line in Fig. 5c).

In 2D electronic spectroscopy, the third-order nonlinear optical response of molecular systems is measured by using a sequence of femtosecond pulses with controlled time delays. As is the case of pump probe experiments, electronically excited state populations and coherences can be created by a pair of pump pulses, and the molecular dynamics in the electronic excited state manifold can be monitored by controlling the time delay $T$ between pump and probe. The additional time delay between two pump pulses enables one to monitor the molecular dynamics as a function of excitation and detection wavelengths for each waiting time $T$. The optical transitions induced by the pump pulses can also create vibrational coherences in the electronic ground state manifold, making it challenging to extract the information about coherent electronic dynamics from multidimensional spectroscopic data.

Our numerically exact simulations of the SP demonstrate that long-lived oscillatory signals in 2D electronic spectra can originate from purely vibrational coherences or from vibronic coherences induced by multi-mode mixing. The latter have been ignored in previous numerical studies which considered only a few intra-pigment modes quasi-resonant with excitonic splitting and neglected all the modes that are far detuned from excitonic transitions as they were deemed to have a negligible effect. However, the correct assessment of the nature of oscillatory 2D signals requires the computation of 2D spectra under the influence of the full spectral density. In order to make such computation feasible, in Supplementary Note 8, we provide an approximate master equation for vibronic dynamics, which takes into account multi-mode mixing effects and quantitatively reproduces numerically exact absorption line shape of the SP. Figure 5d shows the resulting rephasing 2D spectra at waiting time $T = 0$ in the presence of inhomogeneous broadening. The 2D lineshape, shown as a function of excitation and detection wavelengths, is dominated by a diagonal peak excited and detected at 900 nm which coincides with the position of the main absorption peak (see Fig. 5b). To investigate the excited state coherence between vibronic eigenstates $|\psi_+\rangle$ and $|\psi_-\rangle$, which induce the absorption peaks at 780 and 900 nm, respectively, we focus on a cross-peak R12 marked in Fig. 5d. Figure 5e shows the transient of the cross-peak as a function of the waiting time $T$ where the oscillatory 2D signals originating from electronic ground state manifold, shown in red, are comparable to those of excited state signals, shown in blue. The ground state signals consist of multiple frequency components below $1600 \text{ cm}^{-1}$, corresponding to the vibrational frequencies $\omega_k$ of underdamped intra-pigment modes, as shown in Fig. 5f. It is important to note that the excited state signals include a long-lived oscillatory component with frequency $\sim 1800 \text{ cm}^{-1}$, which is not present in the ground state signals and cannot originate from purely vibrational effects as they exceed the high-frequency cut-off of the environmental spectral density (see Fig. 5a). This component must therefore originate from long-lived vibronic coherence due to multi-mode mixing. The long-lived oscillations at $\Delta \sim 1800 \text{ cm}^{-1}$ frequency cannot be described by coarse-grained environment models where only a few intra-pigment modes near-resonant with the excitonic splitting $\Delta = 1300 \text{ cm}^{-1}$ are weakly damped ($\gamma_k = (1 \text{ ps})^{-1}$), while all the other intra-pigment modes are strongly damped ($\gamma_k = (50 \text{ fs})^{-1}$) or neglected ($\gamma_k = 0$) in 2D simulations (see Supplementary Note 8). Our results demonstrate that while some oscillatory components in 2D spectra can originate from purely vibrational motions, long-lived 2D oscillations can also be the result of a strong vibronic mixing of excitons with a large number of underdamped intra-pigment modes.

**Discussion**

Employing numerically exact methods and an analytical theory, we have investigated exciton-vibrational dynamics under the complete vibrational spectrum that has been estimated in earlier experiments. We considered two paradigmatic regimes. The first regime, represented by an excitonic dimer in WSCP, is characterized by an excitonic splitting that is smaller than vibrational frequencies of intra-pigment modes. In this case, the main effect of vibronic coupling to the intra-pigment modes is a reduction of the dipole strength of 0-0 transitions of monomers and of their effective coupling strength $V_{00}$ that determines the splitting between absorption peaks in the low-energy spectrum. A second important effect concerns the modulation of the vibrational sideband of optical transitions by a vibronic mixing between 0-0 and 0-1 transitions. Although the vibronic mixing is not strong enough to modulate the low-energy part of absorption spectra of WSCP, it can induce a notable dipole strength redistribution between 0-0 and 0-1 transitions, which cannot be described by approximate theories where the vibronic mixing is ignored.

In the second regime, represented by the SP of the photosynthetic reaction center of purple bacteria, the excitonic splitting is located in the middle of the high frequency part of the intra-pigment vibrational spectrum. In this case, the splitting between main absorption peaks can be even larger than the bare excitonic splitting, due to multi-mode vibronic mixing effects. This regime is found to be particularly suitable for the discovery of new long-lived quantum coherences in photosynthesis. We found that the coherence time of excitonic dynamics is not simply governed by the lifetime of quasi-resonant intra-pigment modes. Rather it is determined by the lifetimes of individual intra-pigment modes involved in a multi-mode vibronic mixing. This implies that approximate theoretical models based on coarse-graining of the high frequency part of the vibrational environments may underestimate the lifetime of excitonic coherences and could be inappropriate to analyze quantum coherences observed in nonlinear experiments on photosynthetic systems. In addition, our results demonstrate that even if the frequency $\Delta$ of oscillatory 2D signals is not well matched to one of the vibrational frequencies $\omega_k$ of intra-pigment modes, the long-lived 2D oscillations can be vibronic in origin, rather than being purely electronic, as is the case of the SP where $\omega_k \lesssim 1600 \text{ cm}^{-1} < \Delta \approx 1800 \text{ cm}^{-1}$. This implies that the origin of long-lived oscillatory 2D signals cannot be identified based only on a comparison of the frequency spectrum of nonlinear signals with the vibrational frequency spectrum of underdamped modes. Hence,
we contend that previously ignored multi-mode vibronic effects must be included in the interpretation of nonlinear spectroscopic signals before the current debate regarding the presence and nature of long-lived quantum coherences in pigment-protein complexes can be settled conclusively.

Our results suggest the possibility that the energy transfer dynamics between electronic states, such as excitons and charge-transfer states, could be governed by the multi-mode nature of the total vibrational environments, rather than a few vibrational modes quasi-resonant with electronic energy-gaps (see Supplementary Note 9). The generality of the methods employed here also suggests that our results have a broad scope and can be of relevance in a wide variety of scenarios involving strong hybridization of electronic and vibrational degrees of freedom, such as recent observations of nonadiabatic dynamics in cavity polaritonics68,69. We expect that renormalization effects considered here may open entirely new toolboxes for vibrational reservoir engineering with possible applications in information technologies and polaritonic chemistry.

Data availability
The simulated absorption and 2DES data generated in this study are provided in the Source Data file. The data used in this paper are also available from the authors upon request. Source data are provided with this paper.

Code availability
The codes used in this work are available from the authors upon reasonable request.

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