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Electronic Structure And Dynamics Of Higher-Lying Excited States In Light Harvesting Complex 1 From *Rhodobacter sphaeroides*

Peter D. Dahlberg^{1,‡}, Po-Chieh Ting^{2,‡}, Sara C. Massey², Elizabeth C. Martin³, C. Neil Hunter³, and Gregory S. Engel^{2,*}

¹Graduate Program in the Biophysical Sciences, Institute for Biophysical Dynamics, and the James Franck Institute, The University of Chicago, Chicago, IL 60637

²Department of Chemistry, Institute for Biophysical Dynamics, and the James Franck Institute, The University of Chicago, Chicago, IL 60637

³Department of Molecular Biology and Biotechnology, University of Sheffield, Firth Court, Western Bank, Sheffield S10 2TN, UK

Abstract

Light harvesting in photosynthetic organisms involves efficient transfer of energy from peripheral antenna complexes to core antenna complexes, and ultimately to the reaction center where charge separation drives downstream photosynthetic processes. Antenna complexes contain many strongly coupled chromophores, which complicates analysis of their electronic structure. Two-dimensional electronic spectroscopy (2DES) provides information on energetic coupling and ultrafast energy transfer dynamics, making the technique well suited for the study of photosynthetic antennae. Here, we present 2DES results on excited state properties and dynamics of a core antenna complex, light harvesting complex 1 (LH1), embedded in the photosynthetic membrane of *Rhodobacter sphaeroides.* The experiment reveals weakly allowed higher-lying excited states in LH1 at 770 nm, which transfer energy to the strongly allowed states at 875 nm with a lifetime of 40 fs. The presence of higher-lying excited states is in agreement with effective Hamiltonians constructed using parameters from crystal structures and atomic force microscopy (AFM) studies. The energy transfer dynamics between the higher- and lower-lying excited states agree with Redfield theory calculations.

Graphical abstract

Corresponding Author E-mail and phone: gsengel@uchicago.edu (773-704-0818).

[‡]Authors acknowledge equal contributions to the work

Supporting Information: Supplementary tables, figures, and a discussion of the fitting procedure and LH2 are presented in the supporting information. This material is available free of charge via the internet at http://pubs.acs.org.



Introduction

Photosynthetic organisms have developed a nearly universal strategy in light harvesting, by which they spatially separate the absorption of solar energy and the use of that energy to generate separated charges.¹ Rapid energy transfer from the site of absorption to the site of charge separation is crucial for efficient solar harvesting. In photosynthetic organisms, the absorption and charge separation sites can be separated by nearly one hundred nanometers, yet transfer occurs on the order of tens of picoseconds and with near unity quantum efficiency in low light.²⁻³ To accomplish this feat, they employ specialized pigment-proteins known as antenna complexes to absorb solar energy. Antenna complexes are packed with strongly coupled chromophores. This strong coupling leads to delocalized states with strong transition dipoles and a complex electronic structure that is essential for energy transfer. In this manuscript, we employ two-dimensional electronic spectroscopy (2DES) to observe the electronic structure and dynamics of light harvesting complex 1 (LH1) from *Rhodobacter sphaeroides (R. sphaeroides)* embedded in its native photosynthetic membrane.

R. sphaeroides is a purple bacterium known for its near unity quantum efficiency³ and highly symmetric antenna complexes. R. sphaeroides has two types of antenna complexes: a peripheral complex, light harvesting complex 2 (LH2) and a core complex, light harvesting complex 1, that surrounds the reaction center (RC). An 8-Å resolution structure of an RC-LH1-PufX dimer was obtained from a combination of X-ray crystallography and cryoelectron microscopy.⁴⁻⁵ This structure reveals an S-shaped array of chromophores that intertwines around two RCs. The bacteriochlorophyll (BChl) within the LH1 complex absorb maximally at 875 nm, and are often designated as B875; they share many characteristics with the B850 ring of BChl in LH2. The RC-LH1-PufX complex exists in R. sphaeroides as both a monomer and a dimer, but the dimer is the dominant structure in vivo. Each monomer in the dimer contains 28 BChl *a* molecules bound in alternating α - and β polypeptides (Figure 1).⁴ The electronic coupling between BChl *a* in LH1 is well approximated by dipolar coupling except in the case of nearest neighbors where the Mg-Mg distance is on the order of 10 Å.⁴ The coupling strength of nearest neighbors is estimated to be a few hundred wavenumbers.⁶⁻⁷ This strong coupling leads to the delocalization of excitations across BChl a and the existence of higher-lying excited states. A special pair of BChl a embedded in each RC absorbs at 870 nm and accepts excitations from LH1.8-9 The LH1 complex used in this manuscript was obtained from a mutant of R. sphaeroides that lacked LH2, PufX, and the RC. In the absence of PufX and the RC, the predominant

structure of the membrane-embedded LH1 is likely a closed, but empty ring.¹⁰⁻¹¹ Highresolution AFM topographs of membrane-embedded LH1-only complexes showed that they are 9.9 ± 0.7 nm in diameter, smaller than the 11.2 ± 0.6 nm measured for monomeric RC-LH1 core complexes with $16 \alpha_1 \beta_1 BChl_2$ LH1 subunits.¹² Accordingly, the LH1-only complex was assigned to be a ring of $15 \alpha_1 \beta_1 BChl_2$ subunits.¹¹

A closed ring, unlike an open ring, has an additional circular symmetry that leads to a series of pairwise degenerate electronic states. The manifold of states in a closed ring of LH1 adopts a labeling scheme to reflect these degeneracies. The lowest and highest energy states are the only non-degenerate states and, in the case of an LH1, are composed of 15 $\alpha_1\beta_1BChl_2$ subunits; these states are labeled k = 0 and k = 15. All other states are pairwise degenerate and are labeled $k = \pm 1...\pm 14$. In a perfectly symmetric monomeric ring, the transition dipole for most excited states becomes vanishingly small, so the only states with finite transition dipole strength are $k = \pm 1$ and $k = \pm 14$, with $k = \pm 1$ being the dominant transition dipole. However, in nature, this perfect circular symmetry is never realized due to disorder. In the dimeric RC-LH1-PufX complex, the "S" shape breaks the circular symmetry and therefore the degeneracies. Observation of the higher-lying excited states and their lifetime in the photosynthetic membrane can inform on the nearest-neighbor coupling strengths, site energies and coupling to the bath, thus providing great insight into the electronic structure of LH1 *in vivo*.

Previous ultrafast spectroscopy studies have been performed on detergent isolated preparations of LH1. Fluorescence anisotropy data suggest the energy transfer between BChl *a* molecules occurs on 110 fs and 400 fs timescales.¹³ Three-photon echo peak shift, pump-probe, and transient grating experiments resolved four dominant vibrational modes and two decay lifetimes.¹⁴⁻¹⁵ Pump-dump-probe and pump-probe experiments have characterized the exciton equilibration between the low-energy states as taking place on a 100 fs timescale.¹⁶⁻¹⁷ To date, none of these ultrafast studies have analyzed simultaneously the higher-lying excited states and their coupling to the strongly allowed $k = \pm 1$ states. In this paper, we present a 2DES waiting time series of monomeric LH1-only complexes embedded in the photosynthetic membrane from *R. sphaeroides*. We find evidence of higher-lying excited states and resolve waiting time dynamics. We use this information to refine the site energies and nearest-neighbor couplings within the B875 band of chromophores and find good agreement between calculated and experimentally observed energy transfer rates.

Experimental Methods

Preparation of LH1-Only Membranes

A mutant containing only LH1 complexes, designated L3 (*puc1BA pufLMX*), was created by genomic deletion of *puc1BA* and *pufLMX* genes using the methodology described in Mothersole et al.¹⁸ The cells were cultured semi-aerobically in the dark at 30 °C. LH1-only membranes were obtained by disrupting the cells using a French press at 14,000 PSI. A slow spin was performed (12,000 RPM JA 30.STI for 20 minutes) to remove large cellular debris. The supernatant was diluted to an optical density of 0.3 at 875 nm in a 200-µm path length quartz cell (Starna Cells Inc.), which was used for analysis.

Two-Dimensional Electronic Spectroscopy

Two-dimensional electronic spectroscopy correlates excitation energies (λ_{τ}) and stimulated emission/ground state bleach/excited state absorption energies (λ_t) as a function of an ultrafast time delay. Spectrally and temporally resolved signals acquired from this spectroscopic technique are beneficial for determining the electronic structure of systems with dense absorption spectra, such as those of antenna complexes.¹⁹ The theory and experimental design are described elsewhere;²⁰⁻²⁴ here, the working principle and information content of a 2DES spectrum is summarized. 2DES is a third-order optical technique involving three ultrafast laser pulses interacting with a sample. The third-order polarization, created by the three light-matter interactions, generates a photon echo signal in a phase-matched direction. The signal is heterodyned with a fourth pulse known as the local oscillator and is spectrally resolved in a spectrometer, and the heterodyned signal is collected on a camera. A 2DES spectrum is a frequency-frequency correlation map, acquiring extra spectral resolution by Fourier transforming across the coherence time (τ , time delay between pulses 1 and 2) axis. 2DES spectra are collected as a function of the waiting time delay (T, time delay between pulses 2 and 3). 2DES has advantages of both spectral and temporal resolution, correlating energy transfer between excited states with different energies while obtaining lifetimes of excited states.

2DES spectra in this study were acquired using the GRadient-Assisted Photon Echo Spectroscopy (GRAPES) technique.²⁵⁻²⁸ A Ti:Sapphire regenerative amplifier (Coherent Inc.) was used to generate a 30 fs FWHM pulse centered at 800 nm with a 5 kHz repetition rate and a power of 2 W. The pulse was focused through argon gas (4 psi) and then again through air to obtain a ~140 nm broadband pulse. An SLM-based pulse shaper (Biophotonics Solutions) compressed the pulse to 15 fs and shaped the spectrum producing the excitation spectrum shown in Figure 1, which was sufficiently broad to excite both the low energy and high energy states simultaneously. The power for pulses 1-3 was attenuated to 33 μ J/cm² (1.37 ×10¹⁴ photons/cm²). Pulse 4 was attenuated an additional 2 orders of magnitude. The coherence times were sampled from 0 to 200 fs in 0.9 fs steps using the gradient-assisted method. The echo signal was spectrally resolved and heterodyne detected on a high-speed CMOS camera (Phantom Miro M310) triggered at 1/100th the repetition rate of the regenerative amplifier. The spectral phase was determined using separately acquired pump-probe spectra, see Figure S1.²⁹ To extract timescales of waiting time dynamics, each waiting time trace in the phased data was fit to a model function with a biexponential decay plus an offset to recover both early-time dynamics and vibrational relaxations. The residuals from curve fitting at each point in the 2DES map were Fourier transformed to obtain information on vibrational frequencies associated with waiting time dynamics.

Effective Hamiltonian of LH1

Model Hamiltonians were constructed to reproduce the experimental results and revise physical parameters such as nearest neighbor coupling, site energies, and spectral density. Our sample contains a mixture of structures containing 14, 15, and 16 $\alpha_1\beta_1BChl_2$ subunits in both open and closed ring configurations. However, prior AFM studies indicate that the most prevalent structure of LH1 in the absence of PufX and the RC is a closed ring of 15

 $\alpha_1\beta_1BChl_2$ subunits.¹⁰⁻¹¹ Therefore, we constructed three separate effective Hamiltonians to account for the ill-defined structure of monomeric LH1 in the absence of PufX and the RC.

The first Hamiltonian, $H_{closed}^{(15)}$, describes a closed ring of 15 subunits. Because no crystal structures exist with this geometry, a model was constructed with 30 equally spaced BChl *a* in the XY plane arranged in a ring with a radius of 5 nm. The Q_y dipole orientations were alternated between pointing at 203° and 21° in the XY plane. The orientation of the dipoles was taken from the average over the first 20 chromophores in the dimeric crystal structure starting from one end of the S-shaped band.⁴ The site energies were made to alternate between 12150 cm⁻¹ (823 nm) and 12450 cm⁻¹ (803 nm) and the nearest neighbor coupling was approximated as 435 cm⁻¹. These values were comparable to previous studies and reproduced features in the absorption spectra.⁶ The coupling for all other chromophores was calculated assuming purely dipolar interaction. A dipole strength of 8.3 Debye was adopted

for all BChl *a* following the calculations in §ener et al.⁶ The second Hamiltonian, $H_{open}^{(14)}$, was calculated in the same manner except the distances and orientations of chromophores were taken directly from one half of the LH1 in the dimeric RC-LH1-PufX crystal structure shown in Figure 1 which consists of a nearly-closed ring of 14 $\alpha_1\beta_1BChl_2$ LH1 subunits.⁹

The third Hamiltonian, $H_{closed}^{(16)}$, was constructed in the same manner using the monomeric RC-LH1 structure from a similar organism, *Thermochromatium tepidum*.³⁰ This 3.0 Å-resolution structure has a closed ring of 16 $\alpha_1\beta_1BChl_2$ subunits that is slightly elliptical. The ellipticity breaks some of the degeneracies that arise from a closed ring.

The eigenstate energies and dipole strengths for $\rm H_{open}^{(14)}, \rm H_{closed}^{(15)},$ and $\rm H_{closed}^{(16)}$ are listed in Table S1, and stick spectra for all the Hamiltonians can be seen in Figure 2 along with the linear absorption spectrum of LH1-only membranes after the removal of Mie scatter. A comparison of the stick spectra to a 77 Kelvin absorption spectrum is made in SI Figures S2 and S3. All three Hamiltonians give rise to a strongly allowed band of excitonic states at low energy around 875 nm. This feature is the well-known B875 absorption band of LH1. In the cases of $H_{closed}^{(15)}$, and $H_{closed}^{(16)}$, the transition dipole strength is carried almost exclusively by the k±1 states that have a transition energy of 11443/11401 cm⁻¹ (874/877 nm). $H_{open}^{(14)}$ provides a similar electronic structure, with the majority of the oscillator strength being carried by the three lowest energy states, E1-3, with transition energies of 11396-11445 cm⁻¹ (877-874 nm). All three Hamiltonians also produce weak higher-lying excited states with transition energies at approximately 13195 cm⁻¹ (760 nm). These higher-lying excited states, in analogy to the higher-lying B850* states in LH2, will be referred to as B875*. The absorption spectrum is mirrored in the excitation fluorescence spectrum (Figure S4), indicating electronic coupling between the B875* states and the B875 states. Without further analysis, it would be difficult to separate this absorption feature from free BChl a in the sample, which absorbs at 770 nm and overlaps B875*, or from vibrationally excited B875 states. Both free BChl a as well as vibrationally excited states would likely relax to the B875 states before fluorescing. Therefore, the assignment of this feature is not conclusive based

on linear absorption or fluorescence spectra alone and we therefore employ to 2DES and Redfield theory for an unambiguous assignment.

Results and Discussion

The real portion of the phased rephasing 2DES spectra of LH1-only membranes at early waiting times, shown in Figure 3, reveal more features than the linear absorption spectrum. The strongest feature in the phased 2DES spectra appears as the near-diagonal peak at $\lambda_{\tau} = 870 \text{ nm}$, $\lambda_t = 880 \text{ nm}$. The offset from the diagonal is likely due to overlapping excited state absorption and stimulated emission. The absolute-valued spectra for the same waiting times are shown in Figure S5 and show a peak centered at $\lambda_{\tau} = 875 \text{ nm}$ for more details on the fitting procedure, $\lambda_t = 875 \text{ nm}$. A regression of the phased data to a bi-exponential function yields two time constants of $\tau_1 = 55 \pm 2$ fs and $\tau_2 = 1130 \pm 26$ fs (Figure S6). Similar dynamics have been reported previously in LH1 and in the B850 ring from LH2 and have been attributed to vibrational relaxation, protein reorganization and spectral diffusion. ¹⁴

A diagonal peak near $\lambda_{\tau} = \lambda_t = 770$ nm corresponding to B875* is not visible due to its low intensity relative to the B875 diagonal peak. In third-order experiments, the signal intensity scales with the fourth power of transition dipoles, whereas a signal in first-order experiments, such as that in a linear absorption spectrum, scales with the square of transition dipoles. Based on the linear absorption spectrum, we expect the relative strength of the B875* diagonal peak to be ~(0.05)², or 0.25%, of the intensity at the B875 diagonal peak, which is too weak to be observed. The signal intensity of a cross peak in a 2DES spectrum is the result of two interactions with each of the corresponding diagonal features. Based on the linear absorption spectrum, the intensity at $\lambda_{\tau} = 770$ nm and $\lambda_t = 880$ nm would be 5% of the B875 diagonal feature if energy transferred from B875* to B875. The cross peak observed between $\lambda_{\tau} = 770$ nm and $\lambda_t = 880$ nm has a maximum of 0.023 in the normalized spectra (Figure 4 and Figure S7).

The position and amplitude of the cross peak indicate that the states observed in the linear absorption spectrum around 770 nm are transferring excitations to B875. Nevertheless, the existence of coupling and transfer could be due to vibrationally excited B875 states or free BChl *a* that are loosely coupled to the B875 states, rather than transfer from higher-lying B875* states. These possibilities can be excluded by analysis of the waiting time dynamics. The cross peak between B875* and B875 grows in rapidly at early waiting times. A representative waiting time trace at $\lambda_{\tau} = 770$ nm and $\lambda_t = 880$ nm is shown in Figure 5. We observe that the waiting time trace reaches its maximum within 100 fs. When a biexponential function is fit to the data, the trace gives a fast rise of 31 ± 3 fs and a slow decay of 1280 ± 100 fs (see the SI for more details on the fitting procedure). Analysis of the complete cross peak region can be seen in Figure 5b. The rapid timescale for the growth of this cross peak excludes any possibility of energy transfer from free BChl *a* to the B875 states, because any energy transfer between free BChl *a* and LH1 would occur on a picosecond timescale due to weak coupling.

Fourier analysis in the waiting time domain is further utilized to elucidate the origin of the cross peaks. In 2DES, vibrational states give clear oscillations in the waiting time domain that result from vibrational coherences.³¹⁻³³ A vibrational coherence is a superposition between two vibrational states within the same electronic excited state. In phased 2DES spectra, we observe excited state absorption features in the upper off-diagonal region of the

map (Figure 3). From the waiting time trace, we see clear oscillations from this region that persist for more than 2 ps (Figure 6a), which is on the appropriate timescale for vibrational coherences at room temperature. Fourier transforming the residual from exponential fitting yields four distinct vibrational modes at 96, 288, 558, and 728 cm⁻¹ (Figure 6b). These vibrational modes are also observed in the Raman spectrum of LH1 and free BChl a.^{15, 34-36} When the cross peak between B875* and B875 is analyzed, no clear oscillations indicative of vibrational coherences can be resolved (Figure 6 c and d). Thus, we attribute the cross peak on the lower off-diagonal region to be energy transfer from higher-lying B875* states to lower-lying B875 states.

Redfield theory was further used to validate the rapid energy transfer rates observed from B875* to B875. The sub-100 fs transfer rate is similar to results obtained in LH2 from B850* to B850.³⁷ The rapid energy transfer is largely due to the co-localization of the excitations on the same ring of chromophores. Figure S8 shows the transition dipole weight for the B875 and B875* states for the three Hamiltonians. In addition to spatially overlapping states, the rapid rate of energy transfer is also due to high-energy bath modes, some of which can be seen in the vibrational coherences in Figures 6 and S9. In order to properly weight these high-energy bath modes, we used a log-normal spectral density given by

$$J(\omega) = \frac{s}{\sigma\omega\sqrt{2\pi}} e^{-[\ln(\frac{\omega}{\omega_c})]/2\sigma^2},$$
 (1)

with a high cutoff frequency, ω_{c} of 1000 cm⁻¹.³⁸ The Huang-Rhys factor, *S*, was 0.5 and the standard deviation, σ , was 0.8 cm⁻¹ Using this spectral density, the population transfer time was calculated following the procedure outlined by M. Reppert.³⁹ The lifetime for $H_{closed}^{(15)}$, calculated as the transfer time from $k_{\pm 14}$ to k0 and $k_{\pm 1}$, was found to be 51 fs. Similarly, the lifetime for $H_{closed}^{(16)}$ calculated as the transfer time from k_{16} to k_0 and $k_{\pm 1}$, was found to be 54 fs. Lastly, the lifetime for $H_{open}^{(14)}$, calculated as the transfer time from E_{28} to E_{1-3} , was found to be 42 fs. All values are within a factor of two of the observed energy transfer lifetimes of 30-40 fs.

Conclusion

The rapid growth of the cross peak and lack of vibrational coherences in the waiting time frequency domain firmly assigns the B875* states as electronic in origin. Their rapid transfer to B875 mirrors the dynamics found in the B850* and B850 states of LH2 and is in agreement with the effective Hamiltonians and Redfield theory calculations of population transfer rates. The role of B875* remains uncertain. The energy of B875* and its proximity to the RC and LH2 raises the possibility of it being an intermediate state as energy is transferred from the peripheral antenna to the RC, though this is unlikely. Due to the short lifetime of B875*, direct transfer to the RC would not compete with relaxation to the B875 ring despite the strong spectral overlap of the H and B peaks of the RC with the B875*

states. Also, the distance between B800 and B875 makes energy transfer from B800 to B875* an unlikely pathway. While other mechanisms such as super-transfer or moderation of energy transfer based the state of the reaction center are conceivable, the presence of the B875* states is likely an informative byproduct of the strong coupling between chromophores in the B875 ring, which is essential for biological function.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Figure 1.

Left: The 8 Å-resolution structure of a monomer of LH1 complex in *R. sphaeroides* reconstructed from the RC-LH1 dimeric crystal structure found in PDB ID: 4V9G.¹⁴ The protein backbone (shown in gray) holds 28 BChl *a* (shown in green) tightly in space. *Right:* The absorption spectrum of LH1-only membranes is shown in black. The two visible features near 765 nm and 875 nm correspond to B875* and B875, respectively. The normalized laser excitation spectrum used for the 2DES experiments is shown in shaded gray.



Figure 2.

Stick spectra from $H_{open}^{(14)}, H_{closed}^{(15)}$, and $H_{closed}^{(16)}$ are overlaid with the absorption spectra of LH1-only membranes after the removal of Mie scatter. The stick spectra are in good agreement with the absorption spectra, recapitulating the strong B875 states and the weak B875* states.



Figure 3.

Representative phased rephasing 2DES spectra of LH1-only membranes at waiting times of T = 0 fs, T = 50 fs, T = 100 fs and T = 500 fs. All spectra are normalized to the signal maximum across all waiting times.



Figure 4.

Phased rephasing 2DES spectra of the cross peak between B875* and B875 (region enclosed by the red rectangle in the 2DES spectrum above) in LH1-only membranes showing stimulated emission around $\lambda_{\tau} = 770$ nm and $\lambda_{t} = 880$ nm at waiting times of T = 0 fs, T = 50 fs, T = 100 fs and T= 500 fs, respectively. All spectra are normalized to the signal maximum across all waiting times.



Figure 5.

Representative waiting time trace and lifetime map at the B875*-B875 cross peak of LH1only membranes show rapid energy transfer within 40 fs. (a) The black curve is the average waiting time trace of LH1-only membranes at λ_{τ} = 770 nm and λ_{t} = 880 nm (indicated by black diamond in the spectrum to the right) of 10 scans acquired in rapid succession over the course of 2 hours. The width of the gray shading is the standard error (1 σ) representing our uncertainty in the mean. The red trace is a regression of the data to a model bi-exponential function. (b) Lifetime map for the rapid component of the bi-exponential around λ_{τ} = 770 nm and λ_{t} = 880 nm. The saturation level of the map is weighted by both the signal strengths at T = 500 fs for each pixel. The contour curves represent the phased 2DES spectrum at waiting time of 100 fs.



Figure 6.

Waiting time traces from two locations in the 2DES spectra and their conjugate waiting time frequency traces show no clear vibrations at the cross peak between B875* and B875. All black traces in the figures represent an average of 10 scans. The width of the gray shading is the standard error (1 σ) representing our uncertainty in the mean. (a) Waiting time trace of LH1-only membranes at λ_{τ} = 895 nm and λ_{t} = 830 nm showing oscillations due to vibrational coherences at this location. (b) Average power spectrum of the residuals from fitting the same waiting time trace in (a) for each of the 10 scans (resolution is 11 cm⁻¹). These peaks are known vibrational modes of BChl *a*.³⁴ (c) Waiting time trace from the B875*-B875 lower cross peak at λ_{τ} = 770 nm and λ_{t} = 880 nm. (d) Average power spectrum of the residuals from fitting the same waiting time trace in (c) for each of the 10 scans.