Optimal number of pigments in photosynthetic complexes

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Abstract. We study excitation energy transfer in a simple model of a photosynthetic complex. The model, described by the Lindblad equation, consists of pigments interacting via dipole–dipole interaction. The overlapping of pigments induces an on-site energy disorder, providing a mechanism for blocking the excitation transfer. Based on the average efficiency as well as the robustness of random configurations of pigments, we calculate the optimal number of pigments that should be enclosed in a pigment–protein complex of a given size. The results suggest that a large fraction of pigment configurations are efficient as well as robust if the number of pigments is properly chosen. We compare the optimal results of the model to the structure of pigment–protein complexes as found in nature, finding good agreement.

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1. Introduction

Photosynthesis is the main natural process for harvesting the Sun’s energy on Earth, providing a food source for a great variety of organisms ranging from highly evolved photosynthetic systems in higher plants to simpler bacteria [1]. In spite of such diversity, basic underlying principles are shared among the majority of light-harvesting organisms. The initial stage of photosynthesis usually involves multiple pigment–protein complexes (PPCs) that consist of a number of pigment molecules (i.e. chromophores) held in place by a protein cage. PPCs are employed to absorb incoming photons as well as to transport the resulting excitation to the reaction center, where the excitation is used to initiate chemical reactions. The absorption of light, and in particular channeling the absorbed energy to the reaction center, is known to achieve high efficiency [1].

One of the most studied PPCs is the Fenna–Matthews–Olson (FMO) complex that is found in the photosynthetic apparatus of green sulfur bacteria. It is the first PPC for which the atomic structure has been determined [2]. FMO is composed of a large protein envelope that encloses a tightly packed group of seven pigment molecules called bacteriochlorophylls (BChl). In FMO the main role of BChl pigments is actually not to absorb light but instead to transport electronic excitations from the input pigment, which is close to the antenna complex, towards a ‘sink’ pigment that channels excitation to the reaction center. Recently discovered long-lasting quantum coherence in FMO complex [4] gave an additional boost to studies of excitation transport in PPCs. Previously it was namely believed that the transport in PPCs is predominantly of a classical nature. Many aspects of time dependence have been studied [5–7], including the functional role of quantum coherences [7–10], as well as the possibility of transport enhancement via environmental interaction [11–14].

Recent structural analysis [3] suggests the presence of an eighth BChl that is weakly bound to each monomeric unit as an additional input site. Its distance from the core seven pigments is quite large and therefore we do not take it into account in our study.
Also important is the question of the structural characteristics of PPCs that enable efficient excitation transfer, for instance, why a given complex has precisely the shape found in nature. The positions and orientations of pigments, known with high precision from crystallographic measurements, on casual inspection show no clear ordering or organization that would enable an easy classification of efficient configurations. It is also not clear, how special efficient configurations are—whether efficient configurations are a result of a long-lasting process of evolutionary improvement, or, can efficient configurations be readily achieved probing a few random configurations of pigments. Prior to the availability of crystallographic structural data, the average minimal distance between pigment molecules was estimated based on the comparison between the fluorescence yield of *in vivo* and *in vitro* chlorophyll (Chl) solutions [15]. Recently, the efficiency of random configurations within simplified models of PPC was inspected [16–19], suggesting that the efficient configurations are relatively probable. This complies with the results obtained for Photosystem I from plants and cyanobacteria [20, 21], where random orientations of pigment molecules were probed, and high efficiencies of excitation energy transfer (EET) were obtained, irrespectively of the pigment orientation. Also, the PPC configuration was shown to be robust to the removal of a pigment from the complex [20].

A fundamental, yet still unanswered question that we address in this work is, why has a particular photosynthetic complex exactly the specified number of pigments? In other words, what is the optimal number of pigments for a given size of the complex3, or, equivalently, what is the optimal size of the complex for a given number of pigments? For instance, why do we find exactly seven pigments in the FMO complex and not more or less? Using a simple model whose parameters are taken from experiments, that is without any fitting parameters, we calculate the optimal number of photosynthetic pigment molecules for different complex sizes and compare these theoretical predictions with the actual number of pigments found in naturally occurring complexes. We find a very good agreement for a variety of PPCs in different organisms. To judge the optimality we use two criteria: (i) the average efficiency of the excitation transfer, where the averaging is performed over random configurations of pigment molecules and (ii) the robustness of the efficiency to small variations of pigment’s locations. A rationale behind these two choices is that ‘good’ PPCs should have a high efficiency but at the same time should also be robust. A specially ‘tuned’ configuration, that has a very high efficiency but is very fragile, will obviously not work in a natural environment with its changing conditions. Additionally, from an evolutionary perspective, the efficiency should be stable with respect to different foldings of the protein cage. It is advantageous to have a PPC with such a number of pigments that will result in a high average efficiency, i.e. in many close-to-optimal pigment configurations. Thereby, a small change in the environment, be it of a chemical origin or for instance a genetic mutation, will still result in a functional PPC. The robustness of quantum coherence to structural changes in the PPCs has been also found experimentally [22]. The model we use to describe the excitation transfer across the PPC consists of the Lindblad master equation, which describes dipole-coupled pigments with on-site excitonic energies being determined by the distances between disc-like pigments. If two discs come too close, i.e. if they overlap, this effectively rescales their energies, introducing a disorder. We should say that the optimal number of pigments that we predict is quite insensitive to details of the underlying model. Agreement between predictions of our model and naturally-occurring PPCs shows that Nature has optimized PPCs by using just the right number of pigments so that the resulting PPCs are highly efficient and robust at the same time.

3 Note that the size of the PPC might be fixed by external factors like, for instance, the membrane thickness.
2. The model

To calculate the efficiency of the excitation transfer in the PPC, we need equations of motion that describe the dynamics of excitation on multiple chromophores coupled to the environment. It is the ratio of the chromophore–chromophore interaction strength to the chromophore–environment coupling that determines the applicability of the various models. When the interaction between chromophores is small compared to the environmental coupling, the Förster theory [23] is applicable, which leads to a picture of incoherent excitation hopping between chromophores. In the opposite limit of strong inter-chromophore interaction and weak coupling to the environment, the excitation dynamics can be described by a quantum master equation, either the Redfield equation or, employing a secular approximation, the Lindblad equation [24]. For non-perturbative parameter ranges, more advanced methods have been developed [25–27], usually at the expense of higher computational complexity.

Our main optimality criterion, namely the transfer efficiency, is relatively insensitive to details of excitation time dependence. Thus, we are going to use the simplest description of EET with the Lindblad equation, which retains the coherent nature of transport, while still taking environmental interactions into account. We expect that more exact descriptions, which in general enhance excitonic oscillations, lead to similar results due to our averaging procedure. Also, these oscillations appear on the time scale of a few 100 fs, which is much shorter than the time scale of excitonic transfer.

In the following, we will introduce the Lindblad equation for the overlapping disc model, used to describe the excitation dynamics in PPCs. Optimality criteria for the efficiency of PPC configurations, as used in the latter sections, will also be presented.

2.1. Lindblad master equation

The internal dynamics of the system of \( N \) chromophores within a single-excitation manifold is determined by the Hamiltonian of the form [28]

\[
H = \sum_{n=1}^{N} \epsilon_n |n\rangle \langle n| + \sum_{n \neq m = 1}^{N} V_{mn} (|m\rangle \langle n| + |n\rangle \langle m|),
\]

(1)

where a state \(|n\rangle\) represents an excitation on the \( n \)th chromophore site, i.e. the electronic state of the \( n \)th chromophore being in the first excited state. Because EET is sufficiently fast, events with two excitations being present at the same time are rare and it is sufficient to consider only zero and a single-excitation subspace [9]. The coupling \( V_{mn} \) is due to the dipole–dipole interaction between chromophores of the form

\[
V_{mn} = \frac{1}{4\pi \varepsilon_0} \left( \frac{d_m \cdot d_n}{r_{mn}^3} - 3 \frac{(d_m \cdot r_{mn})(d_n \cdot r_{mn})}{r_{mn}^5} \right),
\]

(2)

where \( r_{mn} = x_m - x_n \) is a vector, connecting the \( m \)th and \( n \)th chromophores, \( d_n \) is a transition dipole moment between the ground and the first excited state of the \( n \)th chromophore.

Because the system of chromophores is coupled to the protein and nuclear degrees of freedom, it is described by a reduced density matrix \( \rho \). Decoherence due to environmental interaction, recombination of excitation to the ground state, and transfer of excitation to the sink, are modeled by Lindblad superoperators that augment the von Neumann equation for the time evolution of a density matrix,

\[
\dot{\rho} = -i[H, \rho] + L_{\text{deph}}(\rho) + L_{\text{sink}}(\rho) + L_{\text{recomb}}(\rho).
\]

(3)
To model the effects of the environment, we have taken a simplified picture of purely dephasing Lindblad superoperators (i.e. the Haken–Strobl model), which is believed to capture the basic environmental effects and was used in various previous studies [11–13, 29]. For longer time, relevant for the efficiency of PPC, it has been shown that the description with the Lindblad equation accounts for the main features of the dynamics [30, 31]. The dephasing Lindblad superoperator destroys the phase coherence of any coherent superposition of excitations at different chromophores, and is given by

$$L_{\text{deph}}(\rho) = 2\gamma \sum_{n=1}^{N} \left( |n\rangle \langle n| \rho |n\rangle \langle n| - \frac{1}{2} \{ |n\rangle \langle n|, \rho \} \right),$$

(4)

where a site-independent dephasing rate is given by $\gamma$ and $\{ , \}$ represents the anticommutator. An irreversible transfer of excitation from the $N$th chromophore to the sink $|s\rangle$ is modeled by the Lindblad superoperator

$$L_{\text{sink}}(\rho) = 2\kappa \left( |s\rangle \langle N| \rho |N\rangle \langle s| - \frac{1}{2} \{|N\rangle \langle N|, \rho \} \right),$$

(5)

where $\kappa$ denotes the sink rate. The irreversible loss of excitation due to recombination is given by an analogous term

$$L_{\text{recomb}}(\rho) = 2\Gamma \sum_{n=1}^{N} \left( |0\rangle \langle n| \rho |n\rangle \langle 0| - \frac{1}{2} \{|n\rangle \langle n|, \rho \} \right),$$

(6)

with a site-independent recombination rate given by $\Gamma$. The ground state of a chromophore system (a state without any excitation) is represented as $|0\rangle$. Note that $L_{\text{sink}}$ and $L_{\text{recomb}}$ can be equivalently represented by an antihermitian Hamiltonian at the expense of non-conserved density matrix probability [12], avoiding the need for an additional sink and the ground state.

The relevant environmental parameters going into the Lindblad equation (3) are dephasing strength $\gamma$, recombination rate $\Gamma$ and the sink rate $\kappa$. We use standard values inferred from experiments and used before [12], sink rate $\kappa = 1 \text{ ps}^{-1}$, recombination rate $\Gamma = 1 \text{ ns}^{-1}$ and dephasing rate at room temperature $\gamma = 300 \text{ cm}^{-1}$. The dephasing rate, being a product of the temperature and the derivative of the spectral density, can be estimated by using the experimentally determined parameters of the spectral density (as done e.g. in [12]). This value approximately agrees with the optimal dephasing rate at which transfer is most efficient [11, 12]. We note that the results shown depend very weakly on the actual value of the dephasing rate as long as it is of the same order of magnitude as $\gamma = 300 \text{ cm}^{-1}$. In appendix A we show that the values of $\gamma = 150 \text{ cm}^{-1}$ and $600 \text{ cm}^{-1}$ give almost the same optimal PPC size.

2.2. Overlapping discs model

Because we want to study the dependence of efficiency on the size of the PPC, keeping the number of pigments fixed, we have to account for the size-dependence of the Hamiltonian. On-site energies and interaction strengths in equation (1) will be determined from the geometry of the pigments configuration. By changing the PPC’s size, two gross effects are at play. Firstly,

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4 The values entering the Lindblad equation (3) should be in units of frequency, e.g. $\text{s}^{-1}$. The conversion from inverse centimeters $\text{cm}^{-1}$, as used traditionally in spectroscopy, is given by $\omega [\text{cm}^{-1}] = 200\pi c \tilde{\nu}$, where $c = 3 \times 10^8$ and $\tilde{\nu}$ is value in $[\text{cm}^{-1}]$. 

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as the distance between pigments is reduced, the dipole–dipole interaction between pigments gets larger, enhancing the transfer of excitation among them; this effect is already taken into account by the \( \sim 1/r^3 \) dependence of \( V_{nm} \) in equation (2). Secondly, as the chromophores get even closer together, approaching the distances comparable to the extent of the chromophore electronic orbitals, the Hamiltonian (1) is not sufficient for the description of EET anymore, because the effects due to the electronic orbital overlap become important. A detailed analysis of the processes that take place as chromophores get close together would require advanced quantum chemistry methods and is out of the scope of this paper. However, the main effect can be effectively taken into account by appropriately rescaling the parameters of equation (1). Because the pigment molecules will be deformed, their excitation energies will also change. As the on-site energies \( \epsilon_n \), being of the order of few eV, are about \( \sim 100 \) times larger than \( V_{nm} \), even a small relative change in \( \epsilon_n \) can have a large effect. Effectively, close or even overlapping chromophores will therefore result in widely different values of the on-site energies \( \epsilon_n \) at different sites, i.e. in a disorder. Thus there are two competing factors that determine the optimal size of PPCs: reducing the inter-chromophore distance increases the EET, while the overlapping of chromophores introduces a disorder that effectively suppresses the EET.

On-site energy has an additional contribution due to the local environment (e.g. because of pigment–protein interactions), which is however of the order of \( \sim 10^2 \) cm\(^{-1} \) and is usually much smaller that the on-site disorder due to pigment overlap, which is proportional to the unperturbed on-site energy of \( \sim 10^4 \) cm\(^{-1} \). Therefore, the effects of pigment–protein interaction were neglected when obtaining the results presented in the main text. To verify whether neglecting the on-site disorder due to protein interactions is justifiable, we have also calculated the optimal size for \( N = 7 \) pigments with a random on-site disorder added to each random sample of chromophores. The results (see appendix B) show that a disorder of such magnitude indeed has no gross effect on the results.

Each chromophore in our model is represented as a disc—a thin cylinder—of radius \( r \) and height \( a \). Each disc is supposed to represent an approximate size of the electronic cloud of the orbitals involved in the EET (highest occupied electronic orbital, lowest unoccupied electronic orbital). We have estimated the height to be \( a = 1 \) Å, while the cylinder’s radius is taken as \( r = 4 \) Å. The size of this cylinder, in comparison to a BChl pigment molecule, can be seen in figure 1(a). Radius \( r = 4 \) Å is chosen so that it contains the 16 closest non-hydrogen atoms to the Mg atom located in the center of the pyridine ring. For the given locations and orientations of the discs, we then determine if there are any overlaps between discs. If two discs overlap (for an example see figure 1(b)) we rescale the radius of one of them to the new radius \( r_n \) so that they do not overlap anymore but instead only touch. After eliminating all overlaps, we end up with disc’s radii \( r_n \) (for an example see figure 1(c)).

Provided the radius of the \( n \)th disc \( r_n \) is different from the non-overlapping size \( r \), we have to appropriately rescale the on-site energy \( \epsilon_n \). If the effective size of the electronic cloud is reduced from \( r \) to \( r_n \), the kinetic energy of the electron increases by a factor \( r^2/r_n^2 \). We therefore estimate that the energy of excitation on a resized disc will also scale quadratically with its size, giving the on-site energy dependence

\[
\frac{\epsilon_n}{\epsilon^{(0)}} = \left( \frac{r}{r_n} \right)^2 - 1,
\]

where \( \epsilon^{(0)} \) is the excitation energy of non-deformed pigments, i.e. the energy difference of the two lowest electronic states on a pigment. In FMO \( \epsilon^{(0)} \) is approximately \( \epsilon^{(0)} \approx 12300 \) cm\(^{-1} \).

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Figure 1. Illustration of the simple model of PPC. (a) BChl molecule with omitted phytol tail, enclosed by the cylinder as used in simulations ($a = 1 \text{Å}$, $r = 4 \text{Å}$). (b) Discs, randomly positioned in a sphere, prior to the reduction of radii of overlapping discs. Green discs at the top/bottom represent the input site/output site. (c) Same sample as in (b) after the shrinking of overlapping discs.

The overall offset of on-site energies is irrelevant for the dynamics in the model, therefore we shift all energies by $\epsilon^{(0)}$. Such quadratic on-site energy scaling can be rigorously shown under an assumption that the electronic eigenfunctions of the rescaled pigment are just the rescaled eigenfunctions of the original pigment of radius $r$. Let orbitals $\psi_i$ be the eigenfunctions of the Hamiltonian $H(x) = T(x) + U(x)$, where $T(x)$ is the kinetic energy operator and $U(x)$ is a confining potential. The on-site energy of a given chromophore $\epsilon_n$ is the difference between the energy of the ground and the first excited state, $\epsilon_n = E_2 - E_1$. Assuming that eigenstates $\psi_i$ are just scaled to a smaller volume, $\psi_i^\ast(x) = \lambda^{3/2} \psi_i(x/\lambda)$, scaling the on-site energies from equation (7) is obtained by a comparison of the eigenvalue equations for the original eigenstate $H \psi_i = E_i \psi_i$ and the scaled eigenstate $H^* \psi_i^\ast = E_i^* \psi_i^\ast$, where the scaled confining potential in $H^*$ has to be $U^*(x) = U(x/\lambda)/\lambda^2$. The dipole strength of the chromophores is similarly scaled linearly with the radius $r_n$ of the cylinder,

$$d_n = \frac{r_n}{r} d,$$

where $d$ is the bare transition dipole moment of the original chromophore of size $r$, and $d_n$ is the scaled dipole strength of the resized disc. This can be justified on the same grounds as the scaling of on-site energies, by inserting the rescaled wavefunction $\psi_i^\ast(x)$ into the expression for a transition dipole matrix of a relevant chromophore transition, $d = \langle \psi_e | e \cdot x | \psi_g \rangle$.

There are different possibilities of how to precisely resize the discs in order to avoid overlaps. While different procedures lead to different on-site energies, the determined optimal complex size changes little. The results in the main text were obtained by sequentially inspecting each pair of discs, only resizing the disc having a greater radius afterwards, while keeping the other disc intact. We have verified other resizing procedures, for instance, resizing both discs...
in a pair to the same size. Such resizing effectively reduces the disorder of on-site energies, as even strongly overlapping pigments will have identical on-site energies. Nevertheless, for such a resizing procedure, the determined optimal radius of PPCs are within 2 Å of the values obtained by the resizing procedure used throughout the paper, and are thus within the error bounds of the model.

To summarize, in our overlapping disc model the matrix elements of $H$ are calculated for the given PPC configuration (positions, as well as disc and dipole orientations) by first resizing all overlapping discs, obtaining rescaled radii $r_n$ and then scaling the dipole strengths and on-site energies according to equations (7) and (8).

2.3. Optimality criteria

We have already introduced equations of motion that govern the dynamics of excitation on chromophores, as well the overlapping disc model that allows us to determine the Hamiltonian for a given configuration of chromophores. What is left is the criteria that will enable us to determine whether a given configuration of chromophores is efficient in terms of the EET. The efficiency of the PPC complex is characterized by the probability that the excitation, initially localized on the input site, will be funneled to the reaction center through the output site. For an example of time evolution see appendix C. The efficiency in the model is not unity because excitation can be lost. The probability that the excitation will be transported to the reaction center can be expressed as

$$\eta = 2\kappa \int_0^\infty dt \rho_{NN}(t),$$

which will be used as our main efficiency criterion. Closely related is the average transfer time, which signifies the speed of transfer of excitation to the reaction center, and is expressed as

$$\tau = \frac{2\kappa}{\eta} \int_0^\infty dt t \rho_{NN}(t),$$

with smaller transfer times being better.

As an additional viability criterion of PPC, the robustness of the efficiency to the static disorder will also be inspected. Dynamic disorder due to thermal motion is already effectively described by the dephasing Lindblad terms in equation (3). Static disorder due to structural changes of PPC, for instance due to changes in the biological environment, like temperature, electric charges, etc, should be treated separately. A given configuration of the pigments in the PPC is robust to the static disorder if random displacements of pigments from the original locations do not induce large changes in the PPC’s efficiency $\eta$ (or equivalently, the average transfer time $\tau$). To put it on a more quantitative ground, we define the pigment configuration robustness $\sigma_\eta(x)$ for a given configuration of pigments with positions $x = (x_1, x_2, \ldots, x_N)$, as a standard deviation of efficiency $\eta$ when the pigment coordinates are varied in the neighborhood of the original positions,

$$\sigma_\eta^2(x) = \int (\eta^2(x + y) - \bar{\eta}(x))^2 w(y) \, dy, \quad \bar{\eta}(x) = \int \eta(x + y) w(y) \, dy.$$  

We omitted the disc and dipole orientations from the definition of the robustness of the pigment configuration to simplify the expressions. However, no qualitative differences are to be expected if orientations are also varied when inspecting the robustness.
The probability density $w(y)$ defines the *neighborhood* of a given configuration, and is localized around the original location of the pigments. The most straightforward choice for the distribution $w(y)$ is a product of uncorrelated normal distributions at each pigment location,

$$w(y) = (2\pi \sigma^2)^{-N/2} \exp \left( \frac{y \cdot y}{2\sigma^2} \right),$$  \hspace{1cm} (12)$$

where $\sigma$ defines the size of the neighborhood in which the robustness is being probed. With a given probability distribution, robustness $\sigma_\eta$ is a function of the original pigment locations $x$ and the size of deviations from the original locations $\sigma$. In the limit of small deviations, $\sigma \to 0$, the expression can be simplified to

$$\sigma^2_\eta(x) = \sigma^2 \sum_{i=1}^{3N} \eta_i(x_i)^2, \quad \eta_i(x_i) = \frac{\partial \eta}{\partial x_i} \bigg|_{x_i'=x_i},$$  \hspace{1cm} (13)$$

where the sum goes over all the components of the pigment coordinates. Robustness $\sigma_\eta$, in the limit of small pigment displacements, is thus proportional to the amplitude of pigment displacements.

### 3. Optimal number of pigments: the case of the Fenna–Matthews–Olson complex

In this section, the efficiency and robustness of random configurations within the model will be considered on an example of the FMO complex. The FMO consists of $N = 7$ BChl pigments. On-site energy for BChl was chosen to be $\epsilon^{(0)} = 12\,300 \text{ cm}^{-1}$, which is within the range of the on-site energies for BChls in the FMO as determined in the literature \cite{32–34}. The strength of the transition dipole moment $d = |d|$ was taken as $d^2 = 26 \text{ D}^2$ (note that the published values for $d$ from the calculations and experimental data vary considerably \cite{35}). The on-site energy $\epsilon^{(0)}$ and dipole strength $d$ used hold for BChl pigments in general and therefore the results presented are expected to also be valid for other PPCs containing BChls, not just for the FMO complex.

To determine the optimal size of PPC for a given number of chromophores (or equivalently, the optimal number of chromophores for a given size), we considered two criteria based on the overall behavior of the efficiencies and robustness of random configurations. In the following, the motivation for choosing such optimality criteria will be given, and the results for the case of the FMO will be presented.

### 3.1. Average efficiency

We shall use the average efficiency $\langle \eta \rangle$, averaged over random positions and orientations of pigments enclosed in a predefined volume. The reason for using random averaging with a uniform distribution is twofold: first, the high average EET efficiency under uniform averaging will mean that there are many different configurations that have high efficiency, i.e. high efficiency is globally robust. Choosing averaging over random configurations therefore offers insight into how special efficient configurations of chromophores are within the space of all configurations. The second reason is that we have *a priori* no knowledge of what would be the appropriate measure for possible pigment configurations under, say, different protein cage...

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6 The dipole moment is given in units of debye (D). The conversion of the interaction $V_{mn}$ to units of cm$^{-1}$ is obtained by conversion $\frac{D^2}{4\pi\varepsilon_0} = 5030 \text{ cm}^{-1} \text{ Å}^3$. 

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foldings due to, for instance, mutations. A uniform measure represents in this case a ‘least-information’ distribution.\footnote{We have to note that we also checked other distributions, for instance a uniform distribution on the surface of a sphere of radius $R$, and obtained practically the same results. For instance, the difference in the position of the maximum in figure 2 was within our error estimate of 1 Å (seen as an ‘error’ band in figure 3).} Using configurations sampled according to a uniform distribution over the chromophore positions within a ball of radius $R$ and the random orientations of dipoles and discs, the average efficiency is calculated. Formally, it can be written as

$$
\langle \eta \rangle = \int_{R} \eta(X) w_{\text{conf}}(X) \, dX,
$$

where $X$ contains the positions and orientations of the chromophore discs and dipoles (apart from the positions of the input and output sites, which are fixed on the poles of the sphere), and $w_{\text{conf}}(X) \propto 1$ signifies a uniform distribution of the chromophores inside the sphere. Observing the dependence of the average efficiency $\langle \eta \rangle$ for different numbers of chromophores and different radii $R$ of the enclosing sphere, we can determine the optimal number of chromophores for a given radius $R$, or equivalently, the optimal radius $R_{\text{opt}}$ for a given number of chromophores.

To obtain more detailed information about the efficiencies of a random configuration, we also observed the probability distribution over efficiencies $p_{R,N}(\eta)$, defined as $p_{R,N}(\eta) = \int_{R} \delta(\eta(X) - \eta) w_{\text{conf}}(X) \, dX$. For the number of chromophores as found in FMO ($N = 7$),

**Figure 2.** Probability density $p_{R,N}(\eta)$ for $N = 7$ for a range of sphere radii $R$ (density plot in the background of the left plot; contours connecting equal values of $p_{R,N}$ are also shown). The solid blue curve is the average efficiency $\langle \eta \rangle$. On the right a close-up of $\langle \eta \rangle$ and the average transfer time $\langle \tau \rangle$ (dotted curve, right axis) around the maximum of $\langle \eta \rangle$ is plotted.
Figure 3. (a) The optimal enclosing radius $R_{\text{opt}}(N)$ of the sphere for a different number of chromophores $N$, based on the maximum of the average efficiency $\langle \eta(R) \rangle$. The width of shading (i.e. $\approx \pm 1 \, \text{Å}$) denotes a range of $R$ for which the average efficiency is within 1% of the maximal value. The vertical dashed line marks the case of the FMO with $N = 7$ chromophores. (b) The structure of the FMO complex as determined by the spectroscopic data, is enclosed in the optimal sphere of the radius $R \approx 16 \, \text{Å}$. The structural data was obtained from PDB entry 3EOJ [3].

the probability distribution over efficiencies $p_{R,N}(\eta)$ is shown in figure 2. When going from large radii $R$ to smaller, configurations tend to get more efficient, which is expected as the chromophores are closer to each other, thus increasing the dipole coupling. However, as $R$ is reduced even further, overlapping of chromophores gets more probable, causing an on-site energy disorder. This leads to the localization of excitation on chromophores not connected to the sink site. Such configurations have low efficiency. Therefore, as $R$ gets smaller the distribution $p_{R,N}$ becomes bimodal, with a lower efficiency mode due to overlapping configurations and a high efficiency mode for non-overlapping configurations.

Low efficiency of overlapping configurations therefore leads to a maximum in the average transfer efficiency $\langle \eta(R) \rangle$ at the optimal radius $R_{\text{opt}}$. The average transfer efficiencies at the optimal radius are rather high, e.g. for $N = 7$ in figure 2 it is $\langle \eta(R_{\text{opt}}) \rangle \approx 0.95$ with a large fraction of the configurations having an even larger efficiency than the average. Thus within the model, high efficiency is not due to finely tuned pigment positions and orientations, but occurs for the majority of pigment configurations for the parameters estimated to be relevant in PPCs. For the FMO case with $N = 7$, the optimal radius was estimated to $R_{\text{opt}} \approx 16 \, \text{Å}$, which fits the actual configuration of pigments very well (see figure 3). The average transfer time $\langle \tau \rangle$ is also minimal at $R = R_{\text{opt}}$ (see figure 2). The optimal average transfer time of $\sim 30 \, \text{ps}$ is so large due to the contribution of very inefficient configurations of chromophores. Looking at the average transfer time of the 5% of the most efficient configurations, we get the value of 5 ps, which is comparable to the transfer times as determined using different models of the FMO in [12–14, 36].

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The estimated optimal radius is quite insensitive to small variations of input parameters, e.g. dipole moment $d$, chromophore disc radius $r$ or its thickness $a$, or the scaling of on-site energies and dipole strengths of resized discs. For instance, decreasing the disc’s radius to $r = 3.5 \, \text{Å}$ decreases $R_{\text{opt}}$ by $\approx 2 \, \text{Å}$, changing the disc’s thickness to $a = 0.5$ or $1.5 \, \text{Å}$ changes $R_{\text{opt}}$ by $\approx \pm 2 \, \text{Å}$, while changing quadratic energy scaling to a linear or cubic one again changes $R_{\text{opt}}$ by $\approx \pm 2 \, \text{Å}$. Similarly, changing the dephasing rate $\gamma$ by a factor of 2 changes $R_{\text{opt}}$ by $\approx \pm 2 \, \text{Å}$, see appendix A. Details of the disc resizing procedure also change $R_{\text{opt}}$ for less than $2 \, \text{Å}$, as the extreme case of resizing each overlapping disc pair to the same size reduces $R_{\text{opt}}$ by $\approx -2 \, \text{Å}$.

The optimal radius of the enclosing sphere was obtained from the average efficiency over all random configurations within a sphere. However, even though the evolutionary drive to more efficient configurations might not be very strong if the majority of configurations are already efficient, some optimization is still to be expected. Thus one might argue that the optimal enclosing volume of the natural PPCs should be determined considering only the ensemble of more optimal configurations. We will denote such averages with $\langle \eta \rangle_p$ where $p$ specifies a portion of the most efficient configurations that should be taken into account when calculating the average (e.g. $\langle \eta \rangle_{0.05}$ is the average of $\eta$ over $5\%$ of the most efficient configurations, as shown in figure 4). As $p$ is reduced, the overall value of the average efficiency $\langle \eta \rangle_p$ will increase. The increase will be more pronounced in the region of $R < R_{\text{opt}}$, where the distribution is bimodal. The location of the maximum of $\langle \eta \rangle_p$ will be thus moved to smaller values of $R$, indicating more densely packed chromophore configurations. However, as we will see in the next subsection, the robustness of such densely packed configurations deteriorates very quickly, supporting our choice of estimator for the $R_{\text{opt}}$.

Note that the overlaps between the pigments and protein cage are not considered explicitly in the model. If overlaps with the protein cage would be taken into account, $R_{\text{opt}}$ would represent the size of a protein cage, whereas in our model without pigment-protein overlaps, $R_{\text{opt}}$ is the...
size of a sphere that contains all pigment centers. For instance, looking at figure 3(b), we can see that the sphere with $R_{\text{opt}}$ contains all of the pigment centers, while parts of a few pigments still protrude in the bounding sphere. If overlaps of pigments with the protein cage were taken into account explicitly, $R_{\text{opt}}$ would be approximately by a disc radius $r = 4 \, \text{Å}$ larger, i.e. in the corresponding figure, the bounding sphere would enclose all pigments completely.

3.2. Robustness

The robustness of the PPC configurations to static disorder should also be taken into account when determining whether a given configuration of pigments is feasible, as the conditions in which the PPCs operate are subject to constant environmental changes. In the previous subsection, we have inspected the probability distribution of the efficiencies $\eta$ over random configurations, showing that the majority of random configurations achieve a relatively high efficiency when the enclosing volume is optimal. In this subsection, we will present an analogous analysis of the robustness of random configurations, in particular of those with high $\eta$. We shall show that highly efficient configurations in small enclosing $R$ are very fragile.

We have defined robustness of efficiency $\sigma_\eta$ in equation (11). In simulations we have displaced the pigment positions according to a normal distribution with a width of $\sigma = 0.1 \, \text{Å}$, which is small enough to quantify the robustness in the neighborhood of a specific configuration, while larger than displacements due to thermal vibrations, which are already effectively described by the Lindblad equation. We are specifically focusing on a subset of the most optimal configurations in terms of $\eta$. The average robustness of the subset of the optimal configuration is denoted as $\langle \sigma_\eta \rangle_p$, where $p$ specifies a fraction of most optimal configurations in terms of EET efficiency $\eta$. As an example, we will consider the robustness of the top 5% of efficient configurations $\langle \sigma_\eta \rangle_{0.05}$. The dependence of the average robustness on the radius of the enclosing sphere $R$ is shown in figure 4. The average efficiency of optimal configurations $\langle \eta \rangle_{0.05}$ is also shown in the figure. While the average efficiency of the top 5% of optimal configurations continues to rise as the enclosing sphere radius $R$ is reduced, we can see that the average robustness $\langle \sigma_\eta \rangle$ worsens very quickly as the $R$ drops below $R_{\text{opt}}$.

The rapid worsening of EET robustness with the reducing sphere radius suggests that even if the PPC configurations occurring in nature are indeed optimized in terms of the pigment positions and orientations, the excessive stacking of pigments is not favored as it makes PPC configurations very sensitive to any displacements of pigments. The transition from a robust to a non-robust regime takes place at a radius comparable to $R_{\text{opt}}$ at which the average efficiency $\langle \eta \rangle$ has a maximum. This is not surprising, as both the efficiency of the configurations and the robustness of the configurations are strongly influenced by the overlapping of pigments, which gets more pronounced for $R \lesssim R_{\text{opt}}$.

We have presented the results for the robustness $\sigma_\eta$ of the 5% of most efficient configurations, with pigment displacements $\sigma = 0.1 \, \text{Å}$. The general characteristics of $\langle \sigma_\eta \rangle_p$ however do not quantitatively change for different $p$ (the case of $p = 0.15$ is also shown in figure 4) or displacements $\sigma$. Most importantly, the radius $R$ at which the robustness of configurations drops significantly takes place at approximately the value of $R_{\text{opt}}$. The same robustness behavior is also observed in the limit of infinitesimal robustness from equation (13) where $\sigma \rightarrow 0$. 

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4. Optimal pigment numbers in other pigment–protein complexes

In the previous section we calculated the optimal size $R$ or the optimal number of pigments for the FMO complex. We also demonstrated that a large portion of chromophore configurations has a high efficiency when the enclosing volume is properly chosen ($\sim R_{\text{opt}}$). Additionally, the robustness of configurations to chromophore displacements starts to deteriorate quickly once the enclosing volume is reduced below $R_{\text{opt}}$. Based on these two observations, we argue that the enclosing volume of PPCs occurring in nature should be close to the optimal volume, as determined by our simple model. In this section we will present similar results for the PPCs containing Chl chromophores.

We compare results of the model to the structure of the PPCs from the Photosystem II (PSII) [37], found in cyanobacteria, algae and plants. PSII consists of multiple functional units, which are either part of the outer light-harvesting antenna or the inner core, to which excitations are funneled. In the light-harvesting antenna we will consider the light-harvesting complex II (LHCII), while in the core we will focus on the PC43 and PC47 complexes that funnel excitations to the reaction center and thus have a similar role as the FMO complex in bacteria. A monomeric unit of LHCII contains 14 chlorophyll molecules (8 Chl-$a$ and 6 Chl-$b$), while CP43 and CP47 contain 13 and 16 Chls, respectively. The model parameters for the sink rate, dephasing and recombination are kept the same as in the FMO case, while the transition dipole strength and on-site energies are different for Chl molecules. The transition dipole moment of Chl molecules is chosen as $d^2 = 15$ D$^2$ and the on-site energy $\epsilon^{(0)} = 15 300$ cm$^{-1}$, where values were taken according to [38] (we take the average between the values for Chl-$a$ and Chl-$b$). For the CP43 and CP47 complexes we have simulated random configurations of 13 and 16 chromophores enclosed into a sphere as the actual chromophore positions are distributed relatively uniformly in all directions. The shape of LHCII is however significantly elongated in one direction. We therefore choose the cylindrical volume, having only one additional parameter that has to be provided, i.e. the ratio between the cylinder radius $R_c$ and cylinder height $A$.

Based on the positions of the LHCII chromophores we have estimated the ratio of the two to be $R_c/A = 0.34$.

The CP43 and CP47 primarily play a role of an exciton wire, making the model with the input site and output site at the opposite sides of the sphere applicable. The optimal radius as predicted by the model is $R \approx 18$ Å for CP43 and $R \approx 20$ Å for CP47. As the LHCII also has to transport excitations from adjacent complexes, we have also determined the optimal shape of LHCII, with the input and output sites located at the opposite sides of the enclosing cylinder. With the ratio $R_c/A$ fixed, we have varied the height $A$ of the cylinder and determined the optimal height to be $A_{\text{opt}} \approx 43$ Å.

In addition to acting as an excitation wire, CP43 and CP47 complexes are also directly involved in the absorption of photons, in which case the role of the input site can be taken by any chromophore site. This is an even more common scenario in the LHCII complex, whose primary role is the absorption of photons. To verify whether the findings about the optimal enclosing volume are also valid when the main purpose of the PPC is the absorption of photons, we randomly placed the input site inside the enclosing geometry for each configuration in a random ensemble. The general characteristics of the distribution over efficiencies $p_{R}(\eta)$ do not change considerably, however, the distribution is somewhat shifted to the higher efficiencies because in many random configurations the input site is considerably closer to the output site than the diameter of the enclosing volume. This results in the optimal size of the enclosing...
Figure 5. (a)–(c) Various PPCs enclosed in optimal geometries as described in the main text. The structural data was obtained from PDB entries 1RWT (LHCII) [39] and 3ARC (CP43, CP47) [40]. In (c) the enclosing geometries of two additional monomeric units of the LHCII trimmer are also shown. (d) Plot showing the dependence of the average efficiency $\langle \eta \rangle$ on the size of the enclosing geometry. The upper panel shows the case of CP43 (green) and CP47 (blue), while the lower panel shows the case of LHCII. The solid curves are for the fixed input site, while the dotted line is for the randomly placed input site. Vertical dashed lines mark the sizes of enclosing volumes as used in panels (a)–(c).

Volume being somewhat larger, $R_{\text{opt}} \approx 20$ Å for CP43 and $R_{\text{opt}} \approx 22$ Å for CP47. For the LHCII we have moved the output site to the midpoint on the side between the top and bottom of the cylinder, where the actual output site is supposedly located [41]. For such geometry and
previously used $R_c/A = 0.34$, we have obtained the optimal height of the enclosing cylinder at $A \approx 47 \text{ Å}$.

The optimal enclosing values obtained from the model (averaged between the case for a fixed input site and random input site) were compared to the actual configurations of pigments as obtained from spectroscopic data, and are shown in figures 5(a)–(c), showing good agreement. For the spherical geometries, we have centered the sphere of optimal radius $R_{\text{opt}}$ to the arithmetic mean of locations of BChl/Chl centers. For the LHCII, where a cylinder was used as the enclosing geometry, the cylinder axis was determined such that $\sum_i N r_{i\perp}^2$ was minimal, where $r_{i\perp}$ is the distance from the cylinder axis to the position of $i$th Chl center. Interestingly, the three cylinder axes do not lie in a plane but are instead tilted at an angle $15^\circ$ to the plane containing the three cylinder centers. It is not known if this plays any functional role.

5. Conclusion

We have studied the efficiency of the EET in protein–pigment complexes for random configurations of pigments. The Hamiltonian part of the Lindblad master equation is determined from the geometry of the pigment configurations. If pigments are too close, so that they overlap, this introduces a disorder into on-site energy, effectively inhibiting excitation transport. Fixing the enclosing volume in which pigment molecules are located, we have calculated the average efficiency over random pigment configurations as well as the robustness of efficiency to variations of pigment locations. Doing this, we have determined the optimal number of pigments for a given size of the complex. Even though the model is an oversimplification of actual processes that take place in nature, the statistical predictions obtained from the model are robust to its variations.

Comparing a theoretically predicted optimal number of pigments with several naturally-occurring complexes, we find good agreement. This might indicate that PPCs are not optimized just to have the highest possible efficiency—in fact, efficient configurations are quite common—but instead to be robust to variations in pigment locations. Namely, it turns out that configurations optimized for the highest efficiency, that is those with specially tuned positions and dipole orientations, are very sensitive to small perturbations. The number of pigments in nature is therefore chosen in such a way that the probability of having efficient configurations that are at the same time also robust is the highest.

The presented findings could be in principle verified experimentally by modifying the structure of known PPCs and probing the efficiency of excitation transfer. For the FMO complex the structure was already changed by a mutation of the genes encoding the structure of BChls, as well as by substituting the carbon $^{12}\text{C}$ atoms with $^{13}\text{C}$ [22]. A comparison of excitonic spectra revealed no distinctive differences in the dynamics of excitations, complying with the hypothesis that configurations are not highly tuned but instead very robust. An additional intriguing possibility would be to inspect the characteristics of FMO with mutated protein cages, modifying the positions and orientations of pigment molecules. One could also compare our predictions for the optimal sizes (e.g. figure 3(a)) with other complexes occurring in nature.

Appendix A. Dependence on the dephasing rate $\gamma$

In the simplified model used, environmental interaction is described by the dephasing rate $\gamma$, having the same value for all chromophore sites. In principle, environmental interaction
requires more involved equations of motion (e.g. HEOM [7]), taking into account the spectral density of the environmental modes. However, due to the crude nature of the model, a simplified description is expected to account for the main environmental effects influencing the efficiency of the excitation transfer (see e.g. [30, 31] for a more detailed comparison of approaches). The adequacy of a simple Lindblad-type description of the dynamics for the determination of optimal size is also justified due to the high robustness of the results to the actual choice of the dephasing value $\gamma$, as seen in figure A.1(a), where $R_{\text{opt}}$ only changes for $\sim \pm 2$ Å as the dephasing rate $\gamma$ is changed by a factor of 2. The optimal size $R_{\text{opt}}$ of the PPC is somewhat smaller as the dephasing rate gets stronger, which is expected as a larger dephasing rate enables transfer across the sites with a greater on-site energy mismatch, getting increasingly common in more compact configurations of chromophores.

Appendix B. Random on-site disorder

To verify whether the effects of the local chromophore environment, due to e.g. pigment–protein interaction, can affect the findings about the optimal PPCs sizes, we have amended the Hamiltonian in equation (1) with random on-site disorder $\epsilon_{\text{rand}}^n$ of the magnitudes as found in naturally occurring PPCs (i.e. on-site energy differences in the order of $\sim 100 \text{ cm}^{-1}$). The values of disorder for each realization of random PPCs were calculated according to the Gaussian distribution with variance $\sigma_{\text{rand}}$. The results are shown in figure A.1(b). In the region $R < R_{\text{opt}}$, where the average transfer efficiency is strongly affected by disc overlaps, an addition of random on-site energy disorder has no noticeable effect. The effect is more pronounced for $R > R_{\text{opt}}$ where an overlapping of discs is not the limiting factor of the transfer efficiency any more. The estimated optimal size $R_{\text{opt}}$ however, is not changed considerably by the addition of the random on-site disorder.

Appendix C. Time evolution of site populations

To provide some insight into the temporal dynamics of the excitation transport, we present the time evolution of site populations for two different realizations of PPCs within the Lindblad

Figure A.1. (a) Average efficiencies $\langle \eta \rangle$ for different values of dephasing $\gamma = 150, 300, 600 \text{ cm}^{-1}$. (b) Average efficiencies $\langle \eta \rangle$ for different values of random on-site energies disorder $\sigma_{\text{rand}}$. 

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model. In figure C.1(a), time evolution for the FMO Hamiltonian from [6] is shown, and in figure C.1(b), the time evolution of randomly generated PPC of radius R = 18 Å. The values of dephasing, sink rate and recombination rate are the same as used in the main text.

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