Photosynthesis Course Note

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Contents

1	June 29th	1
2	July 1th	2
3	July 6th	3
4	July 8th	3
5	July 13th	4
6	July 15	5
7	License	6

Abstract

This is a note of the course An Introduction to Light Harvesting in Bacteria and Plants. The course is more of a research than of a pedagodic taste: ideas and methods are only mentioned, with reference to the published paper available in the corresponding slides. Therefore, this note aims to provide a guidance to read those slides. Note that the page numbers in this note are the page numbers of each individual PDF, not necessarily the page numbers displayed on the bottom-right corner inside the PDFs.

1 June 29th

In this lecture, the teacher presents backgroud for light-harvesting mechanism in biology systems:

- pp.1 to 9 of PPT General backgroud to Biophysics.
- pp.10 to 18 of PPT Review of several mechanisms of light harvesting in different species.
- pp.19 to 20 of PPT The central topics of this course

- The natrual of exciton in biological light harvesting system. Is it of Frenkel (short-ranged) of Wannier (long-ranged) type?
- The transfer mechanism. (pp.20) It was mentioned that the transfer is much more efficient than predicted by a classical diffusion model.
- How is active self-regulation archieved. (pp.20)

pp.21 displays a observed long-range transport of excitation energy in a biomimetic light-harvesting system. Then, after a review of Quantum Mechanics (**pp. 22 to 32**)¹, the teacher illustrated a model to calculate the excitation energy of a dimer.(**pp.33 to 36**) Note that here it is expected that only one of the dimer is excited, because this energetically eaiser to achieve. Note also that the symble "†" denotes neither conjugation nor creation operator, but simply a label for an excited state's wavefunction.

The calculated excited state energys on page 35 are splitted into two levels. the difference between them, denoted by \mathcal{E} , is expressed as a function of the two transition dipole moments and the speration length of two molecules. Details about the interaction potential V_{uv} can be found on the next lecture.

2 July 1th

Begin with a review of two modes of excitons in organic system $\mathbf{pp.1}$, this lecture proceeds to explain in detail the dipole interaction potential ($\mathbf{pp.2}$ to 3) V_{uv} mentioned in previous lecture. Page 4 to 5 digresses on the difference between the transition dipole moment and the permanent dipole moment. On page 4, the picture shows the transition dipole moment for a Bacteriacholorophyll (the photosynthetic pigments that occur in various phototrophic bacteria). The right side plot illustrates the absorption spectra for Bacteriacholorophyll. Page 5 originally contains an animated display, which is lost in this PDF file, although the content is not important. Page 6 and 7 also digresses on eletronic momenta and oscillator strength.

The following slide comes back to the excitation energy. It states two forms of dimer: the J aggregation (**pp.9 to 10**), and the H aggregation (**pp.11**). The accessible energy level for absorption is determined by conservation of moment. As the result, the J aggregated dimer absorbs less energy than the monomer case, and a red shift will be observed when comparing the absorption spectra. The case of H aggregation is similar. **Page 12 to 13** mentions the transition dipole in more complicated cases.

¹ Interestingly, the teacher mentioned Bohmian Mechanics, an alternative model of Quantum Mechanics that features determinism and nonlocality. A comprehensive discussion of this model could be found here: http://plato.stanford.edu/entries/qm-bohm/

²Reference could be found in J. Phys. Chem. B, Vol. 107, No. 35, 2003.

3 July 6th

This lecture begins with a note on the unbelievable efficiency of biological system. Then it proceeds to review some concepts in quantum statistical mechanics (page 2 to 21). Since we are quite familiar with this subject, it will be omitted from this note. The last page mentioned an important equation for the evolvement of density matrix in an open system, called the "master equation".

4 July 8th

In the previous lecture, the master equation for density matrix 3 could lead to problematic result. When the density matrix propagates as described by the master equation, the eigenvalues for it might evolve into comprising negative values. This is because the added terms are not rigorously derived, but just an approximation. On **page 2** the teacher mentioned a specific case. Therefore, one introduces the **Lindblad equations** (**pp.3**). This equation ensures that the density matrix $\rho \geq 0$ (i.e. having all positive eigenvalues). However, this operator sometimes breaks down the symmetries of the system. That is, a system started with translation symmetry at t_0 might not have translation symmetry at $t > t_0$.

The teacher wrote an example for the V_m term on ppt on blackboard: Considering the case of a harmonic oscillator.

$$H = w_0 a^{\dagger} a$$

If we add an interaction between the oscillator and other excitons:

$$A^{\dagger}A(a^{\dagger}+a) = \text{extra} \otimes \text{bath}$$

Then $A^{\dagger}A$ is V_m in ppt.

Next, the author provided a specific model of energy transfer. ⁵

pp.5 shows diagramatically the exciton transport process. External influence to the system includes trapping, decaying and disspation. Here k_t characterizes the trapping rate. And k_d characterizes the decay rate.

pp.6 of PPT explains some notations. Here τ_n characterize the life time of exciton in on state n. ⁶ We hope that the total life time $\langle t \rangle = \sum \tau_n$ is small, because the shorter an exciton is fixed on a state, the more random the system is and the more likely that the exciton is trapped.

Following are a series of toycases that has been calculated.

pp.8 to 9 are the $\rho(t)$ for these simple cases.

 $^{^3\}mathrm{Reference}$ not found.

 $^{^4}$ Digression: in the article PRE 65 056120, we can find something about the entanglement between two oscillators.

⁵ **Note**: This section's material could be found in *Excitonic energy transfer in light-harvesting complexes in purple bacteria*. *JChemPhys*_136_245104.pdf

⁶ Here $\rho_n \equiv \rho_{nn}$.

pp.10 of PPT ⁷ This ring with 16 sites shows that there is always a peek Max population in the opposite site(See upper-right plot). This is because there is always two channels of equal length for excitons to get to the opposite site.

pp.11 of PPT is a two ring case, with 8 sites on each ring. The initial configuration is that excitons are evenly excited in the left ring. The right ring comprised of trapping sites.

pp.12 of PPT is another two ring case, with a change in the right ring's number of trapping sites. It shows that a asymmetric design would somehow imporve the efficiency, since the plot on the right has a maximum region while the left one has none. The next slide shows similar result.

Caution: I am quite confused by the following content, from pp.16 to the end of PPT. Erogo the following notes are note well organized. It is advised that one should look at the original literature instead.

pp.15 to 16 shows a more advanced calculation of excited states.

pp.16 of **PPT** Calculating the absorption spectrum, showing that only one level is acceptable.

pp.17 of PPT Changing the configuration of laser could result (in generally) more acceptable levels. However, these laser configurations are not practically achievable.

 ${f pp.19}$ of ${f PPT}$ Using Frankel-Exciton Model and find very good fit, balabala.

5 July 13th

In this lecture, we start with a review of simple harmonic oscillators in quantum mechanics (**pp.1 to 8**). Then turn to the introduction of Glauber coherent states.

Glauber Coherent States (defined on pp.9) is the eigenstate of the annilation operator:

$$A |\alpha\rangle = a |\alpha\rangle \tag{5.0.1}$$

These states maintain their coherence usually longer those number eigenstates. There are other features of the Glauber coherent state, mentioned in **pp.9** to **10**.

The next few deductions will give us a very physical picture of Glauber coherent states. First, express Glauber coherent state as a linear combination of number eigenstates (**pp.11 to 14**). (It is also found that different coherent states are note orthogonal (**pp.15**). Also, the overlap of

 $^{^7}$ Could be found on Efficient energy transfer in light-harvesting systems. Cao and Silbey et al New J Phys (2010)

two coherent state decays expoentially as the difference of the eigenvalues

Digression: usualy, the coherent state representation is much more numerical efficient in computer simulation than the number representation.

Next, we explore several interesting properties of these coherent states:

- Expectations (pp.16 to 17) It is found that $\langle X \rangle = \sqrt{2}\lambda \text{Re}(\alpha)$, and $\langle P \rangle = \frac{\sqrt{2}\hbar}{\lambda} \mathrm{Im}(\alpha)$. Or: α is related to $\langle \alpha | X | \alpha \rangle$ and $\langle \alpha | P | \alpha \rangle$. Hence a real α means the state $|\alpha\rangle$ is a static state with $\langle P \rangle =$ 0. Moreover, this state is the lowest energy eigenstate of a simple harmonic oscillator with the origin shifted from 0 to α .
- Variance (pp.18 to 20) It is also found that the variance $(\Delta X)^2 =$ $\frac{\lambda^2}{2}$, this is exactly the same as the variance of the gound state of simple harmonic oscillator. And $(\Delta P)^2 = \frac{\hbar^2}{2\lambda^2}$. From the two expression, we see that any coherent states have the same minimal uncertainty: $\Delta X \cdot \Delta P = \frac{\hbar}{2}$
- Time Evolution (pp.21) Using the energy eigenstate of simple harmonic oscillator, we can find the time evolution of coherent state as $\alpha(t) = e^{-i\omega t}\alpha_0$. It shows that α is really like a classical oscillator and the norm $|\alpha|^2$ is conserved. This is so cool!

The next slide (pp.22) summarizes the classical feature of Glauber coherent state.

Then we turn to its application in biophysics.

Polaron dynamics in LH1/LH2 We first introduces polaron (pp.24 to 25). The polaron model: Polaron is short for polarized electron. In a polarized crystal, an moving electron will deform the lattice. Then polaron is the electron and those deformed part put it together. This seems quite like the copper pair picture in BCS theory. In the light-harvesting system, since only a very small amount of light is captured, the exciton concentration is low. And we can use the polaron picture for electrons.

Next we write out the Hamiltonian for the LH1/2. (pp.26) The phonon H_{ph} term contains 16 phonons for 16 pigments in LH1. The exciton term H_{ex} assumes that the interaction is only nearst neighbour interaction. The exciton-phonon interaction $H_{\text{ex-ph}}$ is essentially the usual electron-phonon interaction term in solid state physics.

July 15 6

Here we first introduces a new term in the Hamiltonian, $H_{\text{ev-ph}}^{\text{off-diagonal}}$

Taking a linear approximation of the phonon band. We will find a Hunag-Rhys factor S in $\sum_q g_q^2 \omega_q = S\omega_0$.

We will use a time-dependent variational calculation. Then we introduces a Davydov ansatz. Note that

$$e^{A+B} = e^A e^B e^{\frac{1}{2}[A,B]}$$

And using this relationship, it could be found that the $|\beta\rangle$ is actually a Glauber coherent state.

note that the Davydov ansatz is not translational invariant.

Next we project the Davydov ansatz into bloch states, and get the Toyozawa's ansatz. This Toyozawa ansatz has a central position. Using thise, we get the energy band.

Explain for the energy band.

Shows a Holstein polaron's phase diagram. Outside the thin region, there is always a unique solution (global minimum for the variational parameters). Below the region, large polaron. Above, small polaron. Inside the solid region, there will be two solutions coexisting. The the solution obatined depeands on the initial configuration. The physical picture is that, changing the value of J and g, the size of polaron will change sharply when one transverse the dashed line.

Shows some variational parameters calculated. The strange wedge in the picture, is the mode of q=0 mode ($\beta_{q=0}=g$). It is called the Goldston mode, decoupled from the system. This explains that the constant value inside these plots, when q=0.

When we calculate the off-diagonal coupling. There will be two wedges Then frequency plot. The labels are (J, g, ϕ) , where ϕ is the off-diagonal strength.

Another approach, called the Munn-Silbey Method, in which one first do a strange transformation and

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