Graphical Abstract

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Studies of isolated reaction center preparations of PS II have been pivotal in the development of models of both the coupling between its pigments and the process of charge separation. A number of key puzzles remain in our understanding of these important systems. We apply new and powerful optical techniques (CPL and MCPL) and by means of an exciton analysis, point to key new ideas.
New Perspectives on Photosystem II Reaction Centres

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Abbreviations:
Photosystem II - PSII
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Circular Dichroism - CD
Magnetic Circular Dichroism - MCD
chlorophyll-a - Chl-a
Pheophytin-a - Pheo-a
Supplementary Information - SI
ABSTRACT

We apply the differential optical spectroscopy techniques of Circular Polarisation of Luminescence (CPL) and Magnetic CPL (MCPL) to the study of isolated reaction centres (RCs) of Photosystem II (PS II). The data and subsequent analysis provide insights into aspects of RC chromophore site energies, exciton couplings, and heterogeneities. CPL measurements are able to identify weak luminescence associated with the unbound chlorophyll-a (Chl-a) present in the sample. The overall sign and magnitude of the CPL observed relates well to the Circular Dichroism (CD) of the sample. Both CD and CPL are reasonably consistent with modelling of the RC exciton structure. The Magnetic CPL (MCPL) observed for the free Chl-a luminescence component in the RC samples is also easily understandable, but the MCPL seen near 680 nm at 1.8 K is anomalous, appearing to have a narrow, strongly negative component. A negative sign is inconsistent with MCPL of (exciton coupled) Qy states of either Chl-a or Pheo-a. We propose that this anomaly may arise as a result of luminescence from a transient excited state species created following photo-induced charge separation within the RC. A comparison of CD spectra and modelling of RC preparations having a different number of pigments suggests that the non-conservative nature of CD spectra observed is associated with the ‘special pair’ pigments P_{D1} and P_{D2}. 
Introduction

There have been a number of interesting recent developments in the spectroscopy of Photosystem II (PS II) relevant to the fundamental charge separation processes. These include the identification of vibrational coherence [1] following femtosecond pulse excitation of isolated reaction centres (RCs). There has also been the unexpected involvement of far-red absorbing chlorophylls (chlorophyll-\(f\) and chlorophyll-\(d\)) in charge separation processes in some cyanobacteria [2]. The reduction of the lowest energy (\(Q_y\)) excitation in these chlorophylls significantly reduces the driving potential available in subsequent chemical transformations.

The reaction centre of PS II is unique in its ability to photo-generate the (biologically extreme) oxidative potential needed to drive water-splitting in nature. An extensive range of spectroscopic measurements have been made on PS II over the last three decades [3]. These have been performed on purified protein assemblies sourced from a range of photosynthetic organisms. Although the basic content and layout of the photoactive pigments and redox components contained within PS II is well understood through crystallographic and other studies, there remains a notable lack of consensus regarding some of the fundamental processes, spectroscopic assignments and functions in this critical enzyme. To help resolve some of these issues, we introduce and discuss preliminary results generated by two novel low-temperature optical measurements. These techniques have not previously been applied to the study of RCs. We utilise a relatively simple theoretical model to account for exciton coupling between the photoactive chromophores in RCs, including the significant effects of heterogeneity of their pigment site energies. We interpret these results and relate them to some of the measurements that have been made on RCs made using other spectroscopic techniques.

The minimal PS II assembly capable of oxygen evolution is a PS II core complex, as shown in Fig.1(b). The core complex incorporates D1, D2 and Cytb_{559} protein helices of the RC as well as those of the CP43 and CP47 proximal light harvesting antennae. It also contains a range of other protein components. PS II water splitting capacity is incorporated in the manganese-stabilising protein PsbO which hosts the self-assembled CaMn_{5}O_{5} catalytic centre. A PS II core complex is a large and fragile assembly and many studies have resorted to measurements of the isolated PS II RC [4], a simpler preparation containing only the D1, D2 and Cytb_{559} proteins as shown in Fig. 1(a). An isolated RC retains no catalytic function but performs repetitive charge separation followed by recombination following absorption of a photon. It does not undergo the complex multi-photon redox transformations of a functional PS II core. Although isolated RC preparations have been shown to be variable and heterogeneous [5], the absence of the many metastable photo-chemical steps which plague any study of PS II core complexes means that they have been utilised in many studies. A very wide range of spectroscopies have been performed on isolated RCs. A multi-spectral analysis [6] of a
range of optical spectroscopic measurements on RCs led to the first widely accepted description of the excitonic structure of the reaction centre of PS II. The excitonic assignments proposed have been paralleled and refined by other workers \cite{7} although they have been criticised by one group \cite{8}.

Fig. 1

*The structure on the right (b) is a schematic of the crystal structure of the PS II core complex of eukaryotic Cyanidium caldarium* \cite{9}, visualised using PyMol open-source software by Julien Langley. The expanded view (a) of the RC on the left utilises a slightly rotated perspective that enables better visualisation of the key RC pigments. The relevant protein helices are D1 (orange), D2 (cyan) and Cytb\textsubscript{559} (red). Chl-a pigments are shown in green with the special pair pigments P\textsubscript{D1} and P\textsubscript{D2} being darker green. Pheo-a molecules are yellow and the Cytb\textsubscript{559} heme is red. For the PS II core complex (b), the CP43 peripheral light harvesting assembly (left) is mauve whilst the CP47 assembly (right) is blue. Other auxiliary proteins in the core complex are shown in grey. The red arrow displayed in the PS II core complex traces the multi-step electron transfer path from the catalytic CaMn\textsubscript{4}O\textsubscript{5} site to the plastoquinone Q\textsubscript{A} via the secondary donor Tyr\textsubscript{Z}, the primary donor Chl\textsubscript{D1} and the primary acceptor Pheo\textsubscript{D1}. In the RC, Tyr\textsubscript{Z}, the CaMn\textsubscript{4}O\textsubscript{5} and Q\textsubscript{A} are shaded light grey, indicating that they are missing or inactive compared to the same redox component in a core complex. The charge separation and rapid recombination process between Chl\textsubscript{D1} and Pheo\textsubscript{D1} in an RC is indicated by the shorter, red, double-headed arrow.

Circular Polarisation of Luminescence (CPL) measurements, particularly those performed at low temperatures \cite{10}, serve as a natural partner to Circular Dichroism (CD) studies. As fluorescence in the condensed phase most often ensues from the lowest excited state(s) of a molecule (or coupled chromophore assembly) CPL probes the detailed nature of these (photochemically and photophysically) important lowest energy excited states \cite{11}. The low-temperature CPL technique was first developed to study the peripheral light-harvesting antennae of PS II i.e. CP43 \cite{10a} and CP47 \cite{10b}. These measurements helped to clarify excitation transfer processes and, with the assistance of
modelling, provided definitive assignments of site energies and heterogeneity within these relatively complex arrays of ~12-15 Chl-a chromophores.

Magnetic CPL (MCPL) is the luminescence analogue to Magnetic CD (MCD). For either CD or MCD the fractional circular polarisation $\rho_A$ is defined as the ratio of the difference of left and right circularly polarised light, $A_L$ and $A_R$, divided by absorption. This anisotropy ratio has been labelled $\Delta A/A$. The absorbance $A$ is usually defined as the average of absorbance in left and right circularly polarised light. In most cases, $|\rho| << 1$ and thus $A_L\sim A_R$ and $(A_L+A_R)/2$ corresponds to the absorption spectrum. With the definition in (1) the extremal values of $\rho$ are +2 and -2.

$$\rho_A = \Delta A/A = (A_L-A_R)/[(A_L+A_R)/2]$$

(1)

An equivalent definition for the ratio of CPL and MCPL to luminescence spectra $\rho_E$ is

$$\rho_E = \Delta I/I = (I_L-I_R)/[(I_L+I_R)/2]$$

(2)

For a single, purely electronic transition, $\rho_A$ is identical to $\rho_E^{[12]}$. Any mismatch indicates that either (a) the luminescence spectrum may not be fully representative of the ensemble of absorbing species (perhaps owing to heterogeneity) (b) that multiple electronic transitions are involved (c) that the transition is vibronic (i.e. is outside the crude adiabatic Born-Oppenheimer approximation see reference $^{[12]}$ Chapter 3) or (d) that the initially formed excited state changes to a different structure (owing to photophysical or photochemical changes) within the lifetime of the excited state.

MCPL spectra of photosynthetic assemblies have remained unreported until now, although MCD spectra of a number of photosynthetic systems have been reported $^{[13]}$. $\rho_E$ for isolated Chl-a or Pheo-a can be expected to mirror $\rho_A$, with absorption and luminescence known to involve the lowest energy, $Q_y$ state. Exciton coupling and other factors $^{[14]}$ lead to changes in the magnitude of $\rho_A$ for the $Q_y$ state in MCD measurements. The amplitude of $Q_y$ MCD is linear in the applied magnetic field and is also temperature independent, identifying it as an MCD B-term $^{[12]}$. A remarkably strong reduction of $\rho_A$ for the $Q_y$ state MCD of RC-6 near 680 nm was reported some time ago $^{[15]}$. There is a notable similarity to the P680 laser-induced transient changes characteristic of RC-6 charge separation and the so-called ‘MCD deficit’. However, a fundamental understanding of the variations of the corresponding MCD B-term is incomplete $^{[14]}$ and remains a subject of on-going interest and research.

**Method**

*Specialised Spectrometer System*
The single optical spectrometer system utilised was capable of making high-precision measurements of absorption, fluorescence, excitation, CD, MCD, CPL as well as MCPL spectra has been previously described \[16\]. This system has the advantage that simultaneous and/or parallel spectroscopic measurements can be made on a single biochemical sample. Photosynthetic protein preparations can be both variable and fragile. Being able to measure the same sample in the same spectrometer system, without having to thermally cycle or otherwise change the sample is a significant advantage. Protocols, procedures and sample handling systems have been developed that prevent the sample from being degraded by exposure to non-actinic light \[16b, 16c, 17\]. Samples can, if necessary, be loaded whilst only being exposed to light of wavelength longer than 750 nm. The period for which the sample is at ambient temperatures or exposed to air is also minimised.

In any optical spectrometer developed to measure very small levels of dichroism, in either absorption or luminescence, there is often a problem with variable baselines, offsets and other artefacts. When the sample utilised is a low-temperature frozen glass, these problems can become severe. Typically, artefactual dichroism is induced by unintended birefringence of the components in the optical pathway. This birefringence can be induced by even small levels of strain in the optical components of the system (lenses/windows/polarisers) or in the sample itself. For CD (or equivalently MCD) measurements, the presence of residual signals can be identified by (non-zero values of) dichroism measurements in regions of no absorption. For the corresponding CPL measurements, there is no such recourse, as residual circular polarisation of (no) luminescence cannot be monitored. For MCPL, a very useful procedure is to reverse the applied magnetic field, which reverses the sign of the MCPL signal but not that of any CPL (or artefactual signals).

For CPL measurements to be of value, it is critical to develop and apply procedures that can quantify the presence of artefacts, so as to eliminate or at least minimise them. Such procedures were pioneered in the CPL papers on CP43 and CP47 \[10\]. In the current work, the linearity of the MCD/MCPL signal with applied magnetic field as well as the reversal of its sign with applied field was verified, to the extent that the signal-to-noise available in spectra allowed.

A detailed description of the properties and biochemical preparation of RC-6 and RC-6 samples sourced from spinach has been described in detail \[18\]. The samples used for optical experiments were prepared in a similar way to those for the ΔFLN experiments above, except that they were diluted to ~55% with a 1:1 glycerol:ethylene glycol mixture (v/v) as glassing agent. Specialised strain-free, cylindrical fused-quartz sample cells of 12 mm diameter and 0.2 mm to 1 mm in pathlength were used. These cells have also been described \[16c, 17\].
Results and Discussion

CPL and MCPL Spectra

In an analysis of CPL and MCPL it is most helpful to be able to compare luminescence, CPL, MCPL spectra with corresponding absorption, CD and MCD spectra. A detailed comparison of CD and CPL spectra was critical in the analysis of spectra of CP43 \cite{10a} and CP47 \cite{10b}. We report CPL and MCPL data only for RC-6, as it appears to be more photo-stable than RC-5 and less heterogeneous. Clear evidence of RC-5 heterogeneity is provided by a characteristic ~10% reduction of its Cytb\textsubscript{559} chromophore, whereas in RC-6 Cytb\textsubscript{559} is fully oxidised \cite{19}. Additionally, RC-6 has been the RC system studied most in reports of coherence in transient pump-probe experiments \cite{1a}.

Fig 2. compares absorption, fluorescence, CD and MCD spectra of RC-6 with CPL and MCPL measurements. All spectra were made on the same sample or sample preparation. They were all taken on the same spectrometer system and in the same superconducting magnet cryostat. Luminescence, CPL and MCPL spectra were taken simultaneously, as were absorption, CD and MCD spectra. CD and MCD spectra were taken before and after the strong laser illumination necessary to record CPL and MCPL spectra to quantify any changes in the sample. These were found to be minimal.

Spectra were also taken with a reversed magnetic field, which must reverse the sign of a genuine MCPL signal (as it does for the MCD signal) but leaves the CPL (and equivalently the CD) and corresponding luminescence (or absorption) unchanged. Subtraction of these ‘normal’ and ‘reversed field’ spectra provides the MCPL spectrum whilst addition of the two spectra provides the CPL spectrum. CPL spectra recorded at zero applied field were seen to be identical to those determined by the addition of normal and reverse field spectra. The amplitude of the luminescence spectrum decayed slightly upon extended laser illumination at 1.8 K without a significant change of shape in the spectrum. This intensity change effect was corrected for when averaging over many spectra. Averaging was necessary to improve signal-to-noise to a useful level.

The CPL and MCPL spectra in Fig. 2 were averaged over ~ 4-10 scans. As mentioned above, CPL and MCPL spectral components are additive. To exclude artefacts associated with any gradual degradation of the sample, spectra were recorded with the negative field spectrum
taken first and compared with pairs where the positive magnetic field spectra were taken first. This precaution was taken as MCPL spectra of RC-6 appeared to be distinctly anomalous in the 680 nm region (Fig. 2).

Fig. 2
The upper part of the l.h. panel shows overlays of RC-6 absorption and luminescence spectra (grey and black, respectively) taken at 1.8 K. Corresponding CD and MCD spectra (light blue and light green) are compared with CPL and MCPL (blue and green) spectra in the lower section. MCD and MCPL spectra are normalised to an applied magnetic field of 1 T. The r.h. panels show CPL (blue) and MCPL (green) ($I_L-I_R$) spectra overlaid with the luminescence spectrum scaled arbitrarily as $(I_L+I_R)/2$. To match the peak height of the luminescence spectrum at 685 nm, the CPL spectrum ($I_L-I_R$) has been multiplied by 1600. Similarly, the MCPL at 1 T has been multiplied by 20000, to match the luminescence intensity in the 670 nm region. The r.h. scale displays $\rho_E$, the ratio of CPL to luminescence intensities (pale blue) and MCPL at 1 T to luminescence intensities (pale green) as defined in Eqn. (2). Luminescence spectra were taken by illuminating with a Verdi G laser at 532 nm and an excitation fluence of ~100 mW/cm$^2$.

The absorption, CD and MCD spectra shown in Fig. 2 are in good agreement with those previously reported $^{7a, 19-20}$. The main features of the absorption and CD spectra have been accounted for by modelling $^{6-7}$ the exciton coupling interactions between the pigments in the
RC. Utilising the exciton mechanism, and allowing coupling only between Q_y excitations, leads the Q_y CD to be constrained to show overall equal positive and negative components, i.e. summing to zero net area. The CD generated by this mechanism is called conservative. However, both RC-6 and RC-5 display a similar and significant net positive CD component. This non-conservative behaviour has been previously noted but cannot be easily accounted for by the (much smaller) CD of isolated Chl-a or to coupling to other porphyrin excitations.

The $\rho_A$ of RC-6 in MCD varies significantly across the Q_y absorption region (Fig. 3). The spectral region dominated by the ChlZ pigments (~660 – 675 nm) has a value of $\rho_A$ in a 1 T field of ~0.4-0.6 x 10^{-3} (Fig. 3). This is close to the value seen for Chl-a in ether at room temperature of $\rho_A$=0.38 x 10^{-3} [21]. In contrast and as previously noted [15], the MCD spectrum of isolated RCs shows a very strongly reduced MCD amplitude in the 680 nm region. We have previously noted that the reduction of this MCD to absorption ratio, also called the ‘MCD deficit’, was phenomenologically related to the role the Chl-a performed in a photosynthetic assembly [2a, 22]. Pigments involved in charge separation have a significantly reduced MCD. A theoretical understanding of this ‘MCD deficit’ is still lacking [14b].

We emphasise again that for any given purely electronic transition, the CD (or MCD) $\rho_A$ and $\rho_E$ values are identical. They are constrained to be so [12] as they involve the same transition dipole matrix elements. Luminescence at low temperatures in the condensed phase normally
arises only from the lowest excited state of a chromophore or excitonically coupled assembly of chromophores. Therefore, the $\rho_E$ value in CPL (or MCPL) is expected to be constant across a spectral profile, if luminescence arises from a single electronic state. Fig. 2 shows, very clearly, that $\rho_E$ is not constant in RC-6 for either the CPL or MCPL spectra.

The sign of the main RC-6 CPL feature is positive, as expected. It has the same sign as the lowest energy CD feature. The CPL $\rho_E$ is $\sim0.7 \times 10^{-3}$ and reasonably constant across the main luminescence profile. Its magnitude is also in reasonable agreement with the CD to absorption ratio $\rho_A \sim 1 \times 10^{-3}$ in the low energy region of absorption (Fig. 3). The remaining discrepancy seen here is likely to be, at least partly, due to the non-correlated heterogeneity of pigment site energies. The quantum efficiency of fluorescence in RC-6 is around 10% and those sites that emit may not be equivalently represented in absorption.

The region at shorter wavelengths (660 nm to 675 nm) exhibits no CPL to within the signal-to-noise available. Luminescence in this region can be attributed to a small amount of ‘free’ Chl-a present in the sample, these pigment(s) are not excitonically coupled to other pigments. Absorption of this (monomeric) Q_y component will then inherently exhibit only very weak (monomeric) CD and thus have correspondingly weak CPL.

The CPL $\rho_E$ in the region beyond 700 nm gradually vanishes. This region corresponds to the (non-resonantly excited) vibrational sideband luminescence region of RC-6. We have shown that corresponding Q_y vibrational sideband CPL in CP43 is weak, owing to the process of vibronic localisation [10a]. Thus, we attribute the reduction seen in CPL for the long wavelength region in Fig. 2 to the same vibronic localisation effect.

In comparison to the CPL spectrum, the MCPL spectrum of RC-6 exhibits a radically different profile to that of the luminescence spectrum. The MCPL spectrum in Fig. 2 has been scaled to match the luminescence intensity in the 670 nm region, this requires a factor of $2 \times 10^4$. This factor determines the MCPL $\rho_E$ at 1T applied field near 670 nm to be $\sim0.5 \times 10^{-3}$. The value of $\rho_E$ appears to be $\sim 0.3 \times 10^{-3}$ in the spectral range from 690 nm to 720 nm, but the data is relatively noisy. In the spectral range from 675 nm to 685 nm the value of $\rho_E$ drops markedly and even becomes negative at 680 nm.
The 680 nm luminescence feature has $\rho_E = -0.3 \times 10^{-3}$. It has only half the width of the main luminescence feature and contributes $\sim 10\%$ of the total luminescence (see SI for details). It’s CPL appears to be negligible. There are no known examples where the MCD (and equivalently MCPL) of a $Q_y$ chlorin excitation is negative and there is no clear mechanism whereby a chlorin $Q_y$ excitation could exhibit a negative MCD or MCPL. The $Q_y\rho_A$ in 1 T MCD of RC-6 drops to $0.15 \times 10^{-3}$ at 682 nm and thus exhibits the strongest ‘MCD deficit’ to be identified, but $\rho_A$ always remains positive (Fig. 3).

![Luminescence and MCPL spectra](image)

**Fig. 4**
Luminescence (black, grey) l.h. scale with CPL (blue, light blue) and MCPL at 5T (green, light green) r.h. scale, of RC-6 at 1.8 K and 77 K respectively.

When MCPL spectra are taken at 77 K, the puzzling negative MCPL feature seen at 1.8 K is not evident (see Fig. 4). The main luminescence peak becomes weaker at 77 K but apart from some broadening, changes little. There is even less change in the free Chl-a luminescence component and its MCPL is also unchanged.

Whereas chlorins display a positive MCD/MCPL of their (lowest energy) $Q_y$ state, tetrapyrroles in general, as characterised by the four-orbital model of Gouterman $^{[23]}$, can have a very wide range of magneto-optical properties, including instances $^{[24]}$ where the lowest excited state shows a strong negative MCD (and thus MCPL). The observation of a strongly negative, relatively sharp MCPL feature in RC-6 with little CPL and no analogous feature in absorption, suggests the possibility that the unique excited state photoactivity of RC-6 may create a short-lived excited state transient species responsible for the feature observed. We propose that by 77 K the lifetime of the transient species has become shorter and thus contributes less to the luminescence.

Exciton Modelling of RCs
The electronic structure of RCs can be modelled using an exciton coupling model. The pairwise interaction $\hat{V}_{ab}$ couples the transition dipole moments $\tilde{\mu}_{a,b}$, each centred at a Chl-a or Pheo-a, separated by $r_{ab}$.

$$\hat{V}_{ab} = (\tilde{\mu}_a \cdot \tilde{\mu}_b) - 3(\tilde{\mu}_a \cdot r_{ab})(\tilde{\mu}_b \cdot r_{ab}) r_{ab}^{-3} \quad (3)$$

For all 8 pigments in an RC, an 8 x 8 Hamiltonian matrix is created. The diagonal elements are the (uncoupled) site energies of each pigment. The off-diagonal components $\hat{V}_{ab}$ are the coupling terms between each pair of pigments. The magnitude and direction of the Qy transition dipoles can be experimentally determined for isolated, monomeric Chl-a and Pheo-a. Here, we use the same transition dipoles and recently-determined coupling matrix $\hat{V}_{ab}$ as in reference [25] (also shown in Table S3). Structural data is obtained from the x-ray crystal structure reported in reference [26] to provide the relative distances and orientations of the transition dipoles.

Each site energy is attributed a Gaussian-shaped spectral width to account for inhomogeneous broadening. However, variations in the site energy of one pigment are not correlated with those of other pigments. This leads to the situation where the relative site energies of pigments can have a range of realisations with markedly distinct electronic characteristics. A similar phenomenon occurs [27] for the series of inorganic complexes Ir(bpy)$_x$(phen)$_{3-x}$, $x=0-3$ where two distinct luminescences (dual luminescence) are observed in frozen solution spectra of the $x=1$ and $x=2$ cases. In RCs, inhomogeneous broadening is comparable to differences in site energy and the relative order of the two lowest-energy exciton eigenstates can be ‘inverted’ compared to normal for a subset of realisations [28].

To fully account for non-correlation, a large number of realisations ($>10^6$) need to be included to effectively model spectra. This is done by varying site energies over their inhomogeneous (and non-correlated) distributions and then adding each contribution to a total spectrum, each realisation being weighted by its probability. We approximate the influence of phonon wings at 0 K by attributing a Gaussian width of $30 \text{ cm}^{-1}$ to each exciton eigenstate. We do not account for thermal motions by coupling excitations to the thermal bath. This needs to be done to account for the temperature dependence of spectra in more detailed calculations. We further reduce the time needed for the computational modelling by noting that coupling terms between the ChlZ pigments and other pigments are very small. Neglecting coupling to ChlZ pigments reduces the size of the Hamiltonian matrix to be repetitively diagonalised to 6 x 6.
Fig. 5  
The upper panel shows the 1.8 K absorption spectrum of RC-6 (grey trace) along with the absorption line-shape (dotted black trace) calculated using the parameters described in the text and the SI. The lower panel shows the simultaneously measured experimental CD spectrum (blue trace) and the computed CD spectrum (dotted blue trace). In the upper panel, the green envelope is the distribution of lowest energy exciton realisation, whilst the smaller dark grey envelope is the distribution of the lowest energy exciton of the inverted population, i.e. where the lowest exciton is dominantly PheoD2. The light grey envelope is the calculated absorption profile of the inverted population. The light grey envelope in the lower panel is the corresponding calculated CD of the inverted population.

Fig. 5 shows a comparison of calculated and experimental absorption and CD spectra of RC-6, using the parameters given in Tables S2 and S3. Both experimental and theoretical spectra are similar to those previously presented [6b, 7b, 29]. However, we have introduced a number of new features to the analysis.

Firstly, the site energy and width of ChlZD1 is not parameterised but experimentally determined by means of the subtraction of spectra of RC-6 and RC-5 scaled per reaction centre. The coupling between ChlZ pigments and other RC pigments is small (Table S3) and thus the excitonic effects of removing it on other RC pigments is minimal. It has been shown that the MCD magnitude of ChlZD1 is very similar to that of Chl-a in ether and that the RC-6 minus RC-5 spectral profile provides a good model for the Qy spectrum of an isolated Chl-a with known axial coordination [1c]. ChlZD2 was taken to have the same width as ChlZD1 and its spectral position adjusted to reproduce the experimental absorption profile.
Secondly, the amplitude of the calculated CD spectrum in Fig 5 has been determined using the appropriate constants that connect absorption and CD spectra to calculated rotational strength. The CD is not arbitrarily scaled, as has been done in previous comparisons of experimental and calculated spectra. This assists in identifying regions of the CD spectrum that may be responsible for the overall non-conservative characteristics seen. Particularly useful is a quantitative comparison of the CD spectra of RC-6 and RC-5, when scaled per PS II reaction centre (Fig 6 and Fig S2). This establishes that ChlZ_{D1} contributes minimally to the overall CD. This result is consistent with the fact that ChlZ_{D1} interacts only weakly with other pigments in the RC.

Thirdly, we have identified a single set of site energies and widths that is able to account for absorption and CD spectra of both RC-6 and RC-5 reasonably well. Absorption and CD for each Q_y exciton realisation are shown in Fig. 6. Also shown are the absorption and CD of exciton realisations where Pheo_{D2} dominates the lowest energy exciton, i.e. for the inverted population. As the same site energies and widths are used for all pigments except for (uncoupled) ChlZ contributions, calculated absorption and CD spectra of the interacting pigments are the same for RC-5 and RC-6. The lowest-energy, inverted-population exciton has an intensity of 14% of the lowest exciton for all realisations. This result is in agreement with previous modelling [28] of the absorption spectrum of RC-5.

![Fig. 6](image_url)

The l.h. panels and r.h. panels show absorption (upper) and CD (lower) spectra of RC-6 and RC-5 respectively. Experimental absorption and CD spectra (solid grey lines for absorption and solid blue lines for CD) were measured simultaneously. Calculated spectra (dotted lines) use the parameters...
shown in Tables S2 and S3. The brown lines are the (uncoupled) ChlZ absorption components. The filled lineshapes are the six exciton components. The centre panel shows calculated absorption and CD spectra of the 6 coupled pigments common to RC-6 and RC-5 for the inverted population, i.e. where the lowest exciton is dominated by PheoD2. Calculated absorption and CD of the inverted population have been scaled by a factor of 4.

The modelling presented in Figs. 5 and 6 highlights the effects of non-correlated heterogeneity in the spectroscopy of RC-6 and RC-5. Close to 15% of all realisations give rise to a predominantly PheoD2 exciton at lowest energy rather than a ChlD1-based exciton. Absorption and CD spectra of the inverted population are distinctly different to that of the entire population. It seems noteworthy that the MCPL anomaly and the MCD deficit both occur in this inverted population spectral region.

Calculated CD of the second and third excitons (nominally identified as involving PheoD2 and PheoD1) do not exhibit the same profile in absorption and CD (Fig. 6). The CD is seen to change sign across an exciton realisation in some cases. In PS II core complexes, site energy heterogeneity is far smaller and thus the effects of non-correlation become more subtle [30].

RC-6 and RC-5 show a very similar non-conservative CD component amplitude. The CD difference spectra shown in Fig. S2 reveal that the change in CD in passing from RC-6 to RC-5 is close to conservative. This may reflect relatively small shifts and linewidth changes in the PheoD2/PheoD1 region between 675 nm and 681 nm for the two systems. The calculated exciton CD in this region is quite sensitive to small changes. There is virtually no change in the CD in the 650 nm to 670 nm region although the absorption change, owing to the removal of ChlZD1, is significant. Thus, both the excitonic and non-conservative components of the CD in RCs, arise principally from the six central pigments.

The similar amplitude of the calculated and experimental CD in the 680 nm region in RC-5 and RC-6 leads us to suggest that ChlD1/PheoD1 exciton coupling is responsible for the main positive peak seen. This is in spite of the strongest coupling term in PS II reaction centre being between Pd1 and Pd2 (Table S3). Pd1/Pd2 dominate the highest energy exciton (dark blue envelope of Fig 6). Reducing the CD contributions of the Pd1/Pd2 excitons (Fig. S3 provides a full set of pigment distributions for each exciton) we can, at least phenomenologically, account for much of the non-conservative nature of the CD seen. It is not clear how this procedure could be theoretically justified but may be related to the fact that only the Pd1-Pd2 pair of pigments experience direct electronic overlap. This direct overlap also gives rise to the optical charge transfer band [31] in PS II core complexes.

5 Conclusions
A comparison of CD and MCD spectra with measurements of CPL and MCPL spectra show that the CPL is compatible with the observed CD. An anomalous feature appears in the MCPL spectrum near 681 nm. This feature is not present in 77 K spectra, which suggests that the anomaly may be due to the formation of a metastable emitting species having very different electronic properties to that of a chlorin.

Exciton modelling of absorption and CD spectra of RC-6 and RC-5 has been improved. A set of site energies, for which the ChlZD1 value was determined from experiment was introduced and applied to both RC-6 and RC-5. A more recent set of structural parameters and a more refined set of coupling energies were also utilised in the modelling. The presence of a substantial (14%) inverted population in the exciton realisations, for which the lowest exciton is predominantly PheoD1 and not ChlD1 is consistent with an earlier analysis [28].

A comparison of CD spectra of RC-6 and RC-5 establishes that ChlZD1 has little contribution to the Qy CD of RC-6 or RC-5. This can also be reasonably assumed to be the case for ChZD2. The non-excitonic component of the overall CD in the Qy region can be qualitatively accounted for by neglecting the CD associated with the PDI-PD2 special pair exciton realisations.

This work provides clues as to the origin of the non-conservative Qy CD in RCs. Additionally, MCPL data points to the creation of a metastable species in the excited state of RC-6, which puts interpretation of ps and fs transient experiments on RCs in question.

**Supplementary Material**
Document provided.

**Conflicts of Interest**
None

**Acknowledgements**
The technical support of Keith Jackman is gratefully acknowledged. Julien Langley kindly provided the graphics for Figure 1. Martyna Judd was responsible for determining the correct conversion from rotational strength to CD amplitude and assisted in numerous discussions. We recognise the support of the Australian Research Council through grants DP110104565 and DP150103137 (E.K.).

**References**


[27] E. Krausz, J. Higgins, H. Riesen. The apparent "dual emitter" characteristics of iridium bipyridine phenanthroline complexes, [Ir(bpy)x(phen)3-x]3+ for x = 1,2. Inorganic Chemistry. 1993, 32, 4053.


Supplementary Information

New Perspectives on Photosystem II Reaction Centres

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**SI Global Fitting of Luminescence, CPL and MCPL spectra**

Attempts were made to simultaneously fit the luminescence, CPL and MCPL spectra with three Gaussian profiles. These correspond to the main luminescence peak at 685 nm, a broader free Chl-α component centred near 671 nm and an anomalous, relatively narrow peak at 681 nm. The latter is needed mainly to account for the negative MCPL feature. The main luminescence peak lineshape is not well accounted for as it clearly deviates from Gaussian, but a reasonable account of the overall spectral profiles can be provided, which enables an estimate of the relative areas of the components in the luminescence, CPL and MCPL spectra. These are provided in Fig S1 and Table S1.

![Luminescence, CPL and MCPL spectra](image)

**Fig. S1**
Luminescence (black) CPL (light blue) and MCPL at 1 T (light green) of RC-6 at 1.8 K, scaled by factors of 1, 1600 and 20000 together with the global fit (thicker dashed curves) to three Gaussians (See Table S1)

**Table S1**

<table>
<thead>
<tr>
<th>Position nm/ (FWHM cm⁻¹)</th>
<th>Luminescence</th>
<th>CPL</th>
<th>MCPL</th>
</tr>
</thead>
</table>
| 671/(150)                | Amplitude=3.7 x 10⁻³  
Area=1.8 x 10⁻²  
Relative area=0.066 | Amplitude=0  
ρₑ=0 | Amplitude=5.4 x 10⁻⁶  
1T ρₑ=0.28 x 10⁻³ |
| 681/(100)                | Amplitude=9.8 x 10⁻³  
Area=3.1 x 10⁻³  
Relative area=0.115 | Amplitude=0  
ρₑ=0 | Amplitude=-1.2 x 10⁻⁵  
1T ρₑ=-0.26 x 10⁻³ |
| 685/(190)                | Amplitude=3.6 x 10⁻²  
Area=0.22  
Relative area=0.819 | Amplitude=2.5 x 10⁻⁵  
ρₑ=0.69 x 10⁻³ | Amplitude=1.7 x 10⁻⁵  
1T ρₑ=0.1 x 10⁻³ |
**SI Exciton Modelling of Absorption and CD spectra of RC-6 and RC-5**

*Table S2*

<table>
<thead>
<tr>
<th>Site Energy (nm)</th>
<th>FWHM (cm(^{-1}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chl(_{D1})</td>
<td>680</td>
</tr>
<tr>
<td>Pheo(_{D2})</td>
<td>676</td>
</tr>
<tr>
<td>Pheo(_{D1})</td>
<td>669</td>
</tr>
<tr>
<td>P(_{D1})</td>
<td>665.5</td>
</tr>
<tr>
<td>P(_{D2})</td>
<td>665.5</td>
</tr>
<tr>
<td>Chl(_{D2})</td>
<td>669</td>
</tr>
<tr>
<td>ChlZ(_{D1})</td>
<td>672</td>
</tr>
<tr>
<td>ChlZ(_{D2})</td>
<td>665</td>
</tr>
</tbody>
</table>

*The site parameters used for RC5 and RC6 are identical, except that RC5 lacks ChlZ\(_{D1}\) at 672 nm. The spectral position and width of ChlZ\(_{D1}\) was taken from the subtraction \(^{[1]}\) of scaled spectra of RC-6 and RC-5.*

---

*Table S3*

**PSII RC Coupling matrix\(^{[2]}\)**

<table>
<thead>
<tr>
<th>Coupling (cm(^{-1}))</th>
<th>Chl(_{D1})</th>
<th>Pheo(_{D2})</th>
<th>Pheo(_{D1})</th>
<th>P(_{D1})</th>
<th>P(_{D2})</th>
<th>Chl(_{D2})</th>
<th>ChlZ(_{D1})</th>
<th>ChlZ(_{D2})</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chl(_{D1})</td>
<td>0</td>
<td>-2.18</td>
<td>43.51</td>
<td>-27.32</td>
<td>-46.77</td>
<td>3.54</td>
<td>1.67</td>
<td>-0.09</td>
</tr>
<tr>
<td>Pheo(_{D2})</td>
<td>-2.18</td>
<td>0</td>
<td>1.55</td>
<td>12.61</td>
<td>-2.99</td>
<td>41.65</td>
<td>-0.19</td>
<td>-2.57</td>
</tr>
<tr>
<td>Pheo(_{D1})</td>
<td>43.51</td>
<td>1.55</td>
<td>0</td>
<td>-3.96</td>
<td>15.06</td>
<td>-2.37</td>
<td>-2.52</td>
<td>-0.18</td>
</tr>
<tr>
<td>P(_{D1})</td>
<td>-27.32</td>
<td>12.61</td>
<td>-3.96</td>
<td>0</td>
<td>158</td>
<td>-41.83</td>
<td>0.45</td>
<td>0.58</td>
</tr>
<tr>
<td>P(_{D2})</td>
<td>-46.77</td>
<td>-2.99</td>
<td>15.06</td>
<td>158</td>
<td>0</td>
<td>-22.04</td>
<td>0.6</td>
<td>0.61</td>
</tr>
<tr>
<td>Chl(_{D2})</td>
<td>3.54</td>
<td>41.65</td>
<td>-2.37</td>
<td>-41.83</td>
<td>-22.04</td>
<td>0</td>
<td>-0.05</td>
<td>1.8</td>
</tr>
<tr>
<td>ChlZ(_{D1})</td>
<td>1.67</td>
<td>-0.19</td>
<td>-2.52</td>
<td>0.45</td>
<td>0.6</td>
<td>-0.05</td>
<td>0</td>
<td>0.15</td>
</tr>
<tr>
<td>ChlZ(_{D2})</td>
<td>-0.09</td>
<td>-2.57</td>
<td>-0.18</td>
<td>0.58</td>
<td>0.61</td>
<td>1.8</td>
<td>0.15</td>
<td>0</td>
</tr>
</tbody>
</table>

*Transition Dipole strengths were taken\(^{[2]}\) as 5.47 D for chl-a and 4.25 D for pheo-a.*
**SI Quantitative Absorption and CD comparison of RC-6 with RC-5**

Fig. S2

The top and bottom panels compare the absorption (grey) and CD (blue) spectra of RC-6 (thick) and RC-5 (thin) respectively. Spectra are scaled to two pheo-a molecules per RC, i.e. the same number of reaction centres. Also shown are difference spectra (dotted lines), as well as the modelled absorptions of ChlZ$_{D1}$ (dark brown) and ChlZ$_{D2}$ (light brown).
### SI Pigment Distribution of RC excitons

![Figure S3](image)

Fig. S3

The pigment distribution of the six excitons of RC-6 and RC-5 in Fig. 6 of the main text are shown in the six panels of the figure. Each filled and coloured profile is one of the excitons. The coloured lines within each profile shows the contribution of each coupled pigment. The colour coding used for pigments in Exciton 6 applies to all excitons.

The pigment distributions shown in Fig. S3 identify which pigments contribute significantly to each exciton. For example, ChlD1 dominates Exciton 1 and Exciton 6 is an equal mixture of P_{D1} and P_{D2}. Note that the distribution of each pigment varies and may not have the same lineshape as the exciton itself. This is due to the non-correlated heterogeneity of pigment energies. A less detailed pigment distribution is provided in Fig. 4 of the pioneering 2008 paper\(^3\) by the Renger group.

### Supplementary References:

