Action Spectroscopy on Dense Samples of Photosynthetic Reaction Centers of \textit{Rhodobacter sphaeroides} WT Based on Nanosecond Laser-Flash \textsuperscript{13}C Photo-CIDNP MAS NMR

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Abstract Photochemically induced dynamic nuclear polarization magic-angle spinning nuclear magnetic resonance (photo-CIDNP MAS NMR) allows for the investigation of the electronic structure of the photochemical machinery of photosynthetic reaction centers (RCs) at atomic resolution. For such experiments, either continuous radiation from white xenon lamps or green laser pulses are applied to optically dense samples. In order to explore their optical properties, optically thick samples of isolated and quinone-removed RCs of the purple bacteria of \textit{Rhodobacter sphaeroides} wild type are studied by nanosecond laser-flash \textsuperscript{13}C photo-CIDNP MAS NMR using excitation wavelengths between 720 and 940 nm. Action spectra of both the transient nuclear polarization as well as the nuclear hyperpolarization, remaining in the electronic ground state at the end of the photocycle, are obtained. It is shown that the signal intensity is limited by the amount of accessible RCs and that the different mechanisms of the photo-CIDNP production rely on the same photo-physical origin, which is the photocycle induced by one single photon.

1 Introduction

Action spectroscopy measures the response of the system to radiation depending on the wavelength and may allow for identifying the light-absorbing chromophore in a...
photo-process by comparing the efficiency of radiation of different wavelengths in driving the process [1–4]. This method has been introduced in 1928 as *exakte Wirkungsspektrometrie* by Otto Heinrich Warburg investigating the inhibition of the *Atmungsferment* (i.e., cytochrome-c oxidase) by carbon monoxide, leading to the Nobel Prize in Medicine in 1931. Photosynthesis of plants and bacteria has been explored by action spectroscopy since the days of Warburg. In general, physiological parameters, such as adenosine triphosphate production, are observed to measure the action [5]. Alternatively, action spectroscopy in photosynthesis has been measured by optical, photothermal and photoacoustic methods [6–8].

Photosynthetic reaction centers (RCs) of purple bacteria of *Rhodobacter (Rb.) sphaeroides* wild type (WT) are well characterized [9]. The cofactor arrangement (Fig. 1) and the kinetics and spin dynamics of the electron transfer (Fig. 2) are well known. In its center, the primary electron donor P, called the special pair, is composed of two Bacteriochlorophyll (BChl) cofactors P_L and P_M. All cofactors are symmetrically arranged while the carotenoid alone breaks this symmetry. Upon illumination, an electron is transferred from the special pair, the primary electron donor P, and a radical pair P^+BPhe^- is formed. It is born in a pure electronic singlet state S0, which evolves coherently into a triplet state T0, whilst the other two triplet states (T_1 and T_-1) are not populated at high magnetic fields. Several mechanisms have been identified transferring the electron polarization to nuclei. In continuous-illumination photochemically induced dynamic nuclear polarization magic-angle spinning nuclear magnetic resonance (photo-CIDNP MAS NMR) experiments [10–13], up to three mechanisms, called three-spin mixing (TSM) [14, 15], differential decay (DD) [16] and differential relaxation (DR) [17, 18] cause the build up of net photo-CIDNP. In laser-flash photo-CIDNP MAS NMR experiments [13, 19–21], transient nuclear polarization (TNP) is observed occurring selectively from the singlet branch of the radical-pair decay since the nuclear polarization of the triplet

![Fig. 1 Structural arrangement of cofactors in the RCs of *Rb. sphaeroides* WT. Cofactors involved into the spin dynamics of the primary radical-pair formation are: the primary electron donor P is formed by two bacteriochlorophyll a molecules, P_L (right) and P_M (left), the (primary) electron acceptor Φ, a bacteriopheophytin a in the active L branch, and the carotenoid (Car). Other cofactors are shown in light grey (Protein Data Bank entry 1MX3)](image-url)
branch is delayed. The occurrence of TNP can also be rationalized in terms of the classical radical-pair mechanism (RPM) [22, 23], which relies on diffusion and diminishes in steady-state experiments [20]. In WT samples, the TNP is caused by an effect called transiently obscuring polarization (TOP). It is caused by the strong paramagnetic interaction of the nearby carotenoid during its triplet state (3Car), which transiently obscures the nuclear polarization of triplet decay branch [20, 21].

After decay of the carotenoid triplet, this pool of nuclear polarization becomes visible in the NMR experiment again by extinguishing the opposite nuclear polarization of the singlet branch, and the TNP decays. The enhancement of signal intensity and selectivity of the nuclei allows for detailed studies on structure and spin dynamics of the photosynthetic cofactors active in the formation of the primary radical pair in various RCs [24–28].

Also the excitation transfer processes in RCs of Rb. sphaeroides WT are well known. The four BChl cofactors and the two bacteriopheophytin (BPhe) cofactors are excitonically strongly coupled, allowing for fast and efficient excitation transfer towards P occurring on the time scale of some 100 fs and with a quantum yield close to unity [29–33]. The carotenoid cofactor also functions as antenna, however, excitation transfer is slower (∼2 ps) and the quantum yield is lower (∼75%) [34]. Hence, in optically thin samples, the action spectrum is almost identical with the absorption spectrum. In optically dense samples, shielding of the interior sample volume by the pigments in the front layers of the RCs, which are directly exposed to the laser pulses, blocks light transmission through the sample at the absorption frequencies and, therefore, the activation of the pigments in the inner parts of the sample. In addition, light scattering limits the penetration depth, especially of blue light. Under these conditions, the law of Lambert and Beer cannot be applied.
In photosynthetic RC of *Rb. sphaeroides*, bleaching is expected to cause an additional effect, since light absorption leads to the formation of radical pairs and these radical pairs have a different absorption spectrum. The main difference between the absorption spectrum of the $\text{P}^{+}\text{BChl}^{-}$ radical pair and that of the electronic ground state is the total bleaching of the special-pair absorption at 860 nm once a radical pair is formed, while the other absorption bands undergo only little changes [35]. Hence, irradiation into the special-pair absorption may allow for substantial bleaching, especially under highly saturating laser pulses, which are used here and therefore result in a deeper penetration of the light into optically dense samples. Since NMR experiments with classical induction coils obtain the strongest signal from the center of the coil, an increase of penetration depth may have a strong effect on the obtained light-induced signals. For all photo-CIDNP MAS NMR experiments, as have been conducted till date, induction coils and samples in MAS rotors with diameters of 4 or 7 mm have been applied. In this study, we investigate the near-infrared (NIR) action spectrum of the solid-state photo-CIDNP effect in RCs of *Rb. sphaeroides* WT in order to explore the way of light penetration into the samples.

**2 Materials and Methods**

2.1 RC Preparation

$\delta$-Aminolevulinic acid (ALA) is a precursor for biosynthetic tetrapyrrole synthesis. *Rb. sphaeroides* WT was grown under anaerobic conditions in a medium containing 1.0 mM [4-$^{13}$C]-ALA, leading to a specific $^{13}$C isotope label pattern on the BChl and BPhe cofactors (Fig. 3) [36]. The total incorporation ($P_{\text{tot}}$) was calculated by the weighted sum:

![Diagram of Bacteriochlorophyll a molecule with isotope label pattern biosynthetically introduced by feeding with 4-$^{13}$C-$\delta$-aminolevulinic acid (4-ALA). The bacteriopheophytin a (BPhe) cofactors are labeled according to the same pattern.](image)
\[ P_{\text{tot}} = \sum_{n=0}^{8} \frac{n}{8} P_n, \]

where \( n \) stands for the number of labels present in an isotopomer and \( P_n \) is the corresponding fraction of labeled BChl estimated from the isotopic labeling pattern detected with the mass spectrometer [36]. The extent of \(^{13}\text{C}\) incorporation has been determined by gas chromatography–mass spectrometry to be 60 ± 5\%. The RCs were isolated by the procedure described in Ref. [36]. The quinone-depletion in the WT RC is reported in Ref. [11]. The samples were loaded into optically transparent colorless 4-mm sapphire rotors. The samples contained 10 mg of RCs in a volume of about 70 \( \mu l \) and scattered light significantly. The optical density was 1.35 at 865 nm after dilution by a factor 500. Using an extinction coefficient of 75 \( \text{mM}^{-1} \text{cm}^{-1} \) [12], a sample concentration of \( \sim 1 \text{ mM} \) has been calculated.

### 2.2 Absorption spectrum

For measuring of the absorption spectrum of isolated RCs of \( \textit{Rb. sphaeroides} \) WT, \( \sim 20 \mu g \) of RC has been taken out of the MAS NMR rotor and diluted 500 times in tris-hydroxymethyl-aminomethane (Tris) and lauryldimethylamine-N-oxide (LDAO) buffer. The concentration was \( \sim 0.02 \text{ mM} \) (OD = 1.35 cm\(^{-1}\), extinction coefficient of 75 \( \text{mM}^{-1} \text{cm}^{-1} \)). The absorption spectrum has been measured at room temperature.

### 2.3 Setup

The nanosecond-laser flash photo-CIDNP setup has been described in detail elsewhere [19, 20]. Into this setup, an optical parametric oscillator (OPO 532/300, GWU-Lasertechnik, Erftstadt, Germany) tunable between 680 and 1050 nm has been incorporated. Laser power for all experiments has been 28.2 mJ and the beam diameter matched the inner diameter of the MAS rotor of about 3 mm.

### 2.4 MAS NMR measurements

The NMR experiments were performed on an AV-200 NMR spectrometer (Bruker Biospin, Karlsruhe, Germany). Time-resolved nanosecond-flash photo-CIDNP MAS NMR measurements have been performed as described in Refs. [19–21] by changing the delay time between laser light pulse excitation and NMR detection systematically in three steps from 0 to 10 and 100 \( \mu s \).

In all experiments, the signal has been detected with a polarization extinction technique allowing for fast scanning [19], using two-pulse phase modulation (TPPM) proton decoupling [37]. The cycle delay was 300 ms. The temperature has been 233 K at a spinning frequency of 8 kHz. Artificial line broadening of 40 Hz has been applied. The number of scans was 120 k per spectrum. The time-resolved photo-CIDNP MAS NMR spectra were referenced to the \(^{13}\text{COOH}\) response of solid tyrosine-HCl at 172.1 ppm.
3 Results and Discussion

3.1 NIR action spectroscopy with time-resolved photo-CIDNP MAS NMR

The NIR absorption spectrum of diluted RCs of *Rb. sphaeroides* WT is shown as solid line in Fig. 4. The broad absorption band at longest wavelengths (860 nm) is the first electronic transition of the primary donor P, which is of $Q_y$ character (for review, see Ref. [9]). The signals at 803 and 756 nm arise from the $Q_y$ transitions of the two accessory BCHls and the BPhe cofactors, respectively. The weaker bands at 598 and 524 nm originate from the $Q_x$ transitions of the two accessory BCHls and the BPhe cofactors, respectively. Features around 500 nm are caused by the carotenoid [33, 38]. The large signal around 400 nm, which is not completely shown, is composed by the Soret bands of several cofactors.

Action spectroscopy requires a systematic variation of the excitation wavelength. Here, we report the dependence of the $^{13}$C photo-CIDNP MAS NMR intensities on the excitation wavelength at investigating 15 excitations wavelengths in the range from 720 till 940 nm. In Fig. 5, all the spectra have been collected using a delay time between the laser pulse and the NMR detection pulse of 100 μs. This set of experiments has been performed in order to understand the correlation between photo-CIDNP built up by TSM and DD as a function of the light wavelength. Trace A in Fig. 5 is obtained at 940 nm, i.e., at wavelengths longer than the most red-shifted absorption band. Trace B in Fig. 5 is measured at the maximum of the $Q_y$ absorption of the special pair (860 nm). Trace C (820 nm) in Fig. 5 is recorded between the absorption maxima of the special pair and the accessory BCHls. Trace D (803 nm) in Fig. 5 is obtained at the maximum of the accessory BCHl absorption.

![Fig. 4](image-url)

*Fig. 4* The solid line is an ultraviolet–visible-NIR spectrum of purified RCs of *Rhodobacter (Rb.) sphaeroides* WT obtained at room temperature. The spectrum is normalized to the highest signal intensity at 860 nm (left scale). The peak at 860 nm is assigned to the bacteriochlorophylls of P. The peaks at 803 and 756 nm are assigned to the accessory bacteriochlorophylls and the bacteriopheophytins, respectively. The dotted line is the action spectrum of photo-CIDNP intensities of C-19 of P$_L$ (159.8 ppm) obtained 100 μs after the laser flash and plotted as a function of the wavelength. The action spectrum is normalized to the highest signal intensity obtained with an excitation wavelength at 860 nm (right scale)*
Trace E (780 nm) in Fig. 5 is measured in between the absorption maxima of the accessory BChls and the BPheos. Trace F (756 nm) in Fig. 5 is detected at the maximum of the BPheo absorption, and trace G (720 nm) in Fig. 5 is registered in the low-absorbing region beyond the Q_y transitions.

3.2 Characteristics of action spectrum

The dotted line in Fig. 4 shows the action spectrum as obtained for the C-19 resonance of the P_L cofactor at 160.1 ppm (Table 1). The photo-CIDNP intensities have been quantified for the two signals at 159.8 ppm (assigned to C19 of P_L) and 164.0 ppm (assigned to C6 of P_L). Since all light-induced signals are of the same photochemical origin, the ratio between them remains constant over the entire range of excitation frequencies. Only the value at 940 nm is different since the signal-to-noise ratio is too low for proper determination.

The action spectrum is essentially a flat but slightly tilted line with a single maximum at about 860 nm. Once all the RCs are excited regardless of whether excitation happens to be in the absorption maximum of P860 or at the periphery, then the full photo-CIDNP signal amplitude is achieved. For this interpretation, further evidence is obtained by analysis of the shape of the broad action signal around 860 nm. It shows a sharp increase at its flanks and constancy in its central area. Such behavior is typical for optical saturation of the signal. At conditions where the signal intensity is limited by the number of RCs which can be converted to a radical pair, the signal cannot further increase by increase of absorption.
However, the maximum is clearly correlated with the absorption of P, while the absorptions of the other bands do not cause any feature into the action spectrum. They can still contribute to the photo-CIDNP signal by energy transfer to P860 and subsequent charge separation. But unlike the 860 nm absorption band, these bands are not bleached, given the relatively long laser pulse width of 8 ns as compared to the internal energy transfer times of 100–200 fs. Thus, these bands result effectively in a kind of shielding of the lower sample layers and reduce the overall amount of radical pairs being formed. Note that RCs in the front layers are already in the P860\textsuperscript{+}BPhe\textsuperscript{−} state and cannot form a second radical pair during the laser pulse width again, and so it cannot increase the photo-CIDNP signal accordingly.

The featureless baseline in the regions of strong absorption demonstrates that the sample is in the regime of high optical density and corresponds with photo-CIDNP induction on the sample surface only.

While there is no characteristic feature in the photo-CIDNP action spectrum around the absorption bands of the accessory BCHls and the BPheos, the absorption of the special pair induces a clear effect on the action spectrum. The general appearance of the action spectrum is clearly distinguished from the absorption spectrum of the electronic ground state. For an optically thin sample, a coincidence of both types of spectra would be expected. The features of the action spectrum indeed agree with the absorption spectrum of the primary radical pair transient,

| Data set | Trace in Fig. 6 | \( \lambda_{\text{exc}} \) (nm) | TSM/DD 100 \( \mu \)s | Ratio C19/C6 |
|----------|---------------|--------------|-----------------|-------------|
|          |               |              | C6-P\textsubscript{L} | C19-P\textsubscript{L} |
| 1        | A             | 940          | 0.17            | 0.24        | 1.42        |
| 2        |               | 920          | 0.56            | 0.65        | 1.16        |
| 3        |               | 900          | 0.83            | 0.92        | 1.11        |
| 4        |               | 890          | 0.82            | 0.97        | 1.19        |
| 5        |               | 880          | 0.94            | 0.99        | 1.05        |
| 6        |               | 870          | 0.88            | 1.00        | 1.14        |
| 7        | B             | 860          | 0.84            | 0.93        | 1.11        |
| 8        |               | 850          | 0.89            | 0.97        | 1.09        |
| 9        |               | 840          | 0.91            | 0.96        | 1.05        |
| 10       | C             | 820          | 0.74            | 0.78        | 1.06        |
| 11       | D             | 803          | 0.50            | 0.48        | 0.97        |
| 12       | E             | 780          | 0.45            | 0.46        | 1.04        |
| 13       | F             | 756          | 0.50            | 0.49        | 0.99        |
| 14       |               | 740          | 0.51            | 0.53        | 1.03        |
| 15       | G             | 720          | 0.49            | 0.54        | 1.11        |

Signal enhancement by TSM and DD is measured by the intensity of the signal at 159.8 ppm assigned to C19 of P\textsubscript{L} and by intensity of the signal at 164.0 ppm assigned to C6 of P\textsubscript{L}, measured at a delay time \( \Delta \) of 100 \( \mu \)s.
showing strong bleaching around 860 nm [35], while there is no significant change of photo-CIDNP intensities for the other cofactors. Therefore, we conclude that the action spectrum mainly reflects the transient bleaching of the sample which allows for deeper penetration of light. Hence, it is the strength of its bleaching which distinguishes P in the long living radical pair state from the other cofactors and allows for its direct excitation under conditions of optically dense samples. This finding is in line with the strong photo-CIDNP MAS NMR signal obtained with 532-nm excitation [19–21], occurring at an absorption wavelength having low absorption and therefore great penetration depth.

3.3 Ratio of TNP- to TSM/DD-induced nuclear polarization

Figure 6 shows time-resolved photo-CIDNP MAS NMR spectra obtained at three selected excitation wavelengths (traces A–C) with three delay times between the laser pulse and the NMR detection pulse of 0, 10 and 100 μs (spectra shown in black, grey and light grey, respectively). Comparing the three traces, the spectral

![Fig. 6](image_url)

**Fig. 6** $^{13}$C photo-CIDNP MAS NMR spectra of RCs of 4-ALA $^{13}$C-isotope labeled *Rb. sphaeroides* WT collected using laser pulses at wavelengths of 860 nm (A), 803 nm (B) and 756 nm (C), respectively. The absorption maxima of the donor BChls correspond to trace A, of the accessory BChls to trace B and of the BPhe to trace C. Traces A, B and C show the superposition of three spectra having different time delays between the light pulse and the NMR detection pulse (black 0 μs, grey 10 μs, and light grey 100 μs). All the spectra have been collected at a magnetic field of 4.7 T and a temperature of 233 K. The laser pulse length was 8 ns and the laser power was 28.2 mJ in all experiments.
evolution from 0 to 100 μs appears not to change, as suggested by the constancy of all spectral features and their intensity ratios.

While the spectra obtained at 0 μs (black) are mainly caused by the TNP due to broadening of the signals of the carbons of the donor by the nearby molecular triplet of the carotenoid (TOP), the spectrum at 100 μs (light grey) fully represents buildup by TSM and DD. For quantification of the contributions of the different enhancement mechanisms, we took the signal at 160.1 ppm, assigned to C19 of PL measured at a delay time of 0 μs for quantification of the TNP/TOP effect. Signal enhancement via the TSM/DD mechanisms, also quantified by the signal intensity of the signal at 160.1 ppm assigned to C19 of PL measured at a delay time of 100 μs.

Table 2 Analysis of photo-CIDNP MAS NMR intensities measured at different excitation wavelengths λ_{exc} and induced by different signal enhancement mechanisms

| Data set | Peak area TNP/TOP 0 μs | Peak area TSM/DD 100 μs | Ratio TSM/DD to TNP/TOP |
|----------|------------------------|------------------------|-------------------------|
| Number   | Trace in Fig. 6 | λ_{exc} (nm) |                        |                         |
| 7        | A                     | 860                    | −0.2                    | −1.0                    | 0.5 |
| 11       | B                     | 803                    | −0.1                    | −0.5                    | 0.5 |
| 13       | C                     | 756                    | −0.1                    | −0.5                    | 0.5 |

Signal obscured by TOP is measured by the intensity of the signal at 159.8 ppm assigned to C19 of PL measured at a delay time of 0 μs. Signal enhancement via the TSM/DD mechanisms is measured by the intensity of the signal at 159.8 ppm assigned to C19 of PL measured at a delay time of 100 μs.

4 Outlook

A quantification of the effect, for example, according to the theory of Kubelka and Munk [39], may lead to an exact characterization of the optical properties of dense RC samples as well as of the sensitivity distribution within NMR coils. In particular, such knowledge would allow optimizing photo-CIDNP MAS NMR signals by using specific excitation wavelengths.
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