RESEARCH

Chlorophyll triplet states in thylakoid membranes of *Acaryochloris marina.* **Evidence for a triplet state sitting on the photosystem I primary donor populated by intersystem crossing**

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Abstract

Photo-induced triplet states in the thylakoid membranes isolated from the cyanobacterium *Acaryocholoris marina*, that harbours Chlorophyll (Chl) *d* as its main chromophore, have been investigated by Optically Detected Magnetic Resonance (ODMR) and time-resolved Electron Paramagnetic Resonance (TR-EPR). Thylakoids were subjected to treatments aimed at poising the redox state of the terminal electron transfer acceptors and donors of Photosystem II (PSII) and Photosystem I (PSI), respectively. Under ambient redox conditions, four Chl *d* triplet populations were detectable, identifable by their characteristic zero feld splitting parameters, after deconvolution of the Fluorescence Detected Magnetic Resonance (FDMR) spectra. Illumination in the presence of the redox mediator N,N,N′,N′-Tetramethyl-p-phenylenediamine (TMPD) and sodium ascorbate at room temperature led to a redistribution of the triplet populations, with T_3 ($|D|=0.0245$ cm⁻¹, $|E|=0.0042$ cm⁻¹) becoming dominant and increasing in intensity with respect to untreated samples. A second triplet population (T₄, |*D*|=0.0248 cm⁻¹, |*E*|=0.0040 cm⁻¹) having an intensity ratio of about 1:4 with respect to T₃ was also detectable after illumination in the presence of TMPD and ascorbate. The microwave-induced Triplet-minus-Singlet spectrum acquired at the maximum of the |D|–|E| transition (610 MHz) displays a broad minimum at 740 nm, accompanied by a set of complex spectral features that overall resemble, despite showing further fne spectral structure, the previously reported Triplet-minus-Singlet spectrum attributed to the recombination triplet of PSI reaction centre, ${}^{3}P_{740}$ [Schenderlein M, Çetin M, Barber J, et al. Spectroscopic studies of the chlorophyll *d* containing photosystem I from the cyanobacterium *Acaryochloris marina*. Biochim Biophys Acta 1777:1400–1408]. However, TR-EPR experiments indicate that this triplet displays an *eaeaea* electron spin polarisation pattern which is characteristic of triplet sublevels populated by intersystem crossing rather than recombination, for which an *aeeaae* polarisation pattern is expected instead. It is proposed that the observed triplet, which leads to the bleaching of the P_{740} singlet state, sits on the PSI reaction centre.

Keywords Photosystem I · Reaction centre · Triplet state · Electron spin polarisation · Optically detected magnetic resonance · Electronparamagnetic resonance · *Acaryochloris marina*

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Introduction

Acaryochloris marina is an unusual cyanobacterium (Miyashita et al. [1996](#page-17-0)) which harbours Chlorophyll (Chl) *d* as the main chromophore bound to the core complexes of both Photosystem I (PSI) and Photosystem II (PSII). Chl *a*, that is otherwise the almost ubiquitously dominant pigment in the core complexes of oxygenic reaction centres (RC), is still present in *A. marina* with stoichiometries varying from 1–2 to 2–4 molecules per RC (Hu et al. [1998;](#page-16-0) Akiyama et a. [2002](#page-15-0); Itoh et al. [2007;](#page-16-1) Tomo et al. [2007,](#page-18-0) [2008](#page-18-1), [2011\)](#page-18-2). The electronic transition of Chl *d* is about 15–20 nm red-shifted with respect to that of Chl *a* in organic solvents (Holt and Morley [1959;](#page-16-2) Nieuwenburg et al. [2007;](#page-17-1) Niedzwiedzki and Blankenship [2010](#page-17-2)). Similar red-shifts of the electronic transition are present in the Chl (*a*)/*d* binding protein complexes of *A. marina* (Schiller et al. [1997;](#page-18-3) Hu et al. [1998;](#page-16-0) Itoh et al. [2007](#page-16-1); Tomo et al. [2008](#page-18-1), [2011\)](#page-18-2), with respect to those of the organisms in which Chl *a* is the predominant chromophore. The bulk, constitutive, replacement of Chl *a* with Chl *d* in *A. marina* can then be seen as an adaptive strategy to the environmental niche in which this organism strives, since it experiences substantial fltering of incident sunlight by other photosynthetic organisms, resulting in a large enhancement of far-red/near infrared radiation (Kühl et al. [2005](#page-16-3)).

Since Chl *d* represents more than 95% of the chlorophyll pool in this organism, this implies an average reduction of approximately 100 meV in the excited state energy at disposal for photochemical conversion. Such decrease in the maximal utilisable photon energy poses, in turn, questions regarding the minimal energetic requirements for oxygenic photosynthesis, in conjunction with the efficiency of light conversion in photosystems operating with a lower energy headroom. For this reason, the role of the residual amount of Chl *a* present in both PSI and PSII of *A. marina* has been associated not only to light harvesting but also to photochemical and/or successive electron transfer (ET) reactions (Kumazaki et al. [2002](#page-16-4); Schlodder et al. [2007](#page-18-4); Itoh et al. [2007;](#page-16-1) Tomo et al. [2007,](#page-18-0) [2008](#page-18-1), [2011;](#page-18-2) Ohashi et al. [2008](#page-17-3); Renger and Schlodder [2008\)](#page-17-4). Similar questions concerning the minimal energetic requirement for photochemical energy conversion apply also to a more recently discovered class of cyanobacteria that are capable of conditionally activating the synthesis of both Chl *d* and, to a larger extent, Chl *f*, an even further intrinsically red-shifted chromophore. The biosynthesis of long-wavelength absorbing Chls is elicited when these organisms are exposed to heavy fltering of incident radiation by other photosynthetic organisms and/or environmental factors, through the so-called FaRLiP response (Gan et al. [2014](#page-16-5); Ho et al. [2017](#page-16-6)). Furthermore, the accumulation of Chl *d* and *f* is accompanied by a large reshaping of the photosynthetic apparatus involving the synthesis of specifc isoforms of both PSI and PSII subunits as well as of the phycobilisome antenna. Yet, the maximal extent of Chl *d* and *f* accumulation is about 10% of the total Chl pool, so that Chl *a* remains always the dominant pigment in cyanobacteria capable of the FaRLiP response (Gan et al. [2014;](#page-16-5) Ho et al. [2017](#page-16-6)). Whereas the role of the intrinsically red-shifted Chl *d* and, especially, Chl *f* as antenna pigments upon the induction of the FaRLiP transition is generally accepted, their possible involvement in photochemical reactions in PSII (Allakhverdiev et al. [2016](#page-15-1); Nürnberg et al. [2018;](#page-17-5) Mascoli et al. [2020](#page-17-6); Zamzam et al. [2020](#page-18-5)) and especially PSI, remains controversial, since evidence both in favour (Kaucikas et al. [2017](#page-16-7); Nürnberg et al. [2018\)](#page-17-5) and against (Cherepanov et al. [2020](#page-15-2); Kato et al. [2020\)](#page-16-8) their participation in primary energy conversion has been presented. The very extensive and constitutive replacement of Chl *a* with Chl *d* in *A. marina* may then be understood as an extreme case of adaptation to incident light having a highly enriched component in the far-red portion of the spectrum.

The possible role of Chl *a* in the Chl *d*-dominated photosystems of *A. marina* and associated bioenergetic reconfgurations are briefy reviewed in the following paragraphs.

Photosystem I

In *A. marina* PSI clear evidence for the involvement of Chl *d* in photochemical conversion was originally gathered by transient optical spectroscopy at room temperature (RT) showing that the main bleaching associated with the oxidation of the terminal electron donor peaked at 740 nm, thereafter labelled as P_{740} (Hu et al. [1998](#page-16-0)). This is a remarkable red-shift with respect to the bleaching observed in canonical PSI, which falls at approximately 700 nm (Kok [1961](#page-16-9); Ke [1973;](#page-16-10) Webber and Lubitz [2001\)](#page-18-6), with subtle variations between species especially in the associated fne spectral structure (Witt et al. [2003;](#page-18-7) Nakamura et al. [2011](#page-17-7)). Further confirmation of the radical cation of P_{740} being located on Chl *d* molecules came from FT-IR (Sivakumar et al. [2003](#page-18-8); Hastings and Wang [2008](#page-16-11)) and hyperfne-resolved electron paramagnetic resonance (EPR) spectroscopy (Mino et al. [2005](#page-17-8)) investigations. These evidences were later confrmed by structural models (Hamaguchi et al. [2021;](#page-16-12) Xu et al. [2021\)](#page-18-9) showing a pair of Chl *d/*Chl *d'* (the latter being the epimer) molecules forming a dimer at the interface of the PsaA and PsaB reaction centre (RC) subunits, in analogy to Chl *a/*Chl a' composing P_{700} . Transient absorption measurements performed at low temperature (5–10 K) indicate that both the cation (P_{740}^+) , under ambient redox conditions, and the triplet state $({}^{3}P_{740})$, populated under reducing conditions, display, even in place of signifcant diferences in the satellite bands, a main bleaching at about 735–740 nm (Schenderlein et al. [2008\)](#page-18-10) and hence likely involve either the same Chl(s) *d*

giving rise to the diferential absorption spectrum at RT, or a pool of Chls strongly interacting with them.

Surprisingly, the structural model of *A. marina* PSI identifed the presence of Pheophytin *a* (Pheo *a*, the Chl *a* free base) at the so-called eC_3 position, in a symmetrical arrangement in both RC subunits (Hamaguchi et al. [2021](#page-16-12); Xu et al. [2021](#page-18-9))*.* These cofactors, which are Chl *a* molecules in canonical PSI, can be identifed with, or be part of, the socalled A_0 electron acceptor (reviewed in Brettel [1997](#page-15-3); Santabarbara et al. [2005](#page-18-11)). The involvement of Pheo *a* in either primary charge separation or charge stabilisation reactions is consistent with the previous investigation by Kumazaki and coworkers (Kumazaki et al. [2002](#page-16-4)) which attributed a transient bleaching centred at approximately 680–685 nm to a Chl *a* acting as an electron acceptor. Since Pheo *a* has an intrinsically higher redox potential than Chl *a* by approximately 100–200 mV, depending on the solvent characteristics (Kobayashi et al. [2013](#page-16-13)), the substitution can be rationalised in terms of a compensation for the lower energy of the excited state, P_{740}^* with respect to P_{700}^* . This may explain the similarity in photon trapping kinetics reported for the isolated PSI of *A. marina* both by ultrafast transient absorption (Kumazaki et al. [2002\)](#page-16-4) and fuorescence lifetime studies (Mi et al. [2003\)](#page-17-9) with respect to Chl *a* binding PSI of other cyanobacterial species (reviewed in (Gobets and van Grondelle [2001](#page-16-14)).

The nature of the Chls in the so-called eC_2 position, located between the P_{740} Chl *d* dimer and the Pheo *a* molecules, is less well established because of difficulties in assigning unambiguously the Chls at the current model resolution. Hence, although the authors assigned these molecules also to Chl *d*, there remains a possibility for Chl *a* being part of the RC.

The structure of *A. marina* PSI does not show any other significant differences in the arrangement of the other cofactors involved in electron transfer reactions. This is in agreement with previous functional spectroscopic studies indicating that the rate of charge recombination between P_{740}^{+} and either the reduced terminal iron-sulphur cluster acceptors $F_{A/B}$ (Hu et al. [1998\)](#page-16-0) or the semi-phylloquinones, *A*[−], both at RT (Hu et al. [1998](#page-16-0)) and at cryogenic temperatures (Schenderlein et al. [2008\)](#page-18-10), were in line with those of canonical PSI. Only the kinetics of A_1^- oxidation by the successive acceptor, the iron-sulphur cluster F_X appeared to be somewhat slower (biphasic, 355 ns and 88 ns (Santabarbara et al. [2012](#page-17-10))) than in canonical PSI (250–300 ns and 10–25 ns, for reviews see: Brettel [1997;](#page-15-3) Santabarbara et al. [2005](#page-18-11), [2010\)](#page-18-12). The similarity of both forward electron transfer (ET) and recombination kinetics therefore indicates a substantial conservation of the energetic properties of the PSI acceptor side in *A. marina*, considering that the redox midpoint potential of the terminal donor P_{740}^+ (Bailleul et al. [2008](#page-15-4); Schenderlein et al. [2008](#page-18-10)) is also equivalent to that of

P⁺₇₀₀, *i.e.* +425–460 mV (Nakamura et al. [2011\)](#page-17-7). Besides the nature of the pigments composing the RC, the other main diference revealed by the structures between canonical PSI and that of *A. marina* is the reduced number of antenna molecules, with a total of about 75 Chl *d* in the latter with respect to about 90 Chl *a* in the former.

Photosystem II

The nature of the cofactors involved in photochemical reactions in *A. marina* PSII and, in particular, the specifc role of Chl *d* have not been fully established yet. Nonetheless, evidence has been reported in favour of the primary electron donor being either Chl *d* or a Chl *d*/Chl *a* heterodimer (Schlodder et al. [2007](#page-18-4); Renger and Schlodder [2008;](#page-17-4) Ohashi et al. [2008;](#page-17-3) Tomo et al. [2007](#page-18-0), [2011](#page-18-2)). The mean excited state energy for the pigment constituting the RC has been on this basis estimated to correspond to a photon of between 713 nm (Tomo et al. [2007,](#page-18-0) [2011\)](#page-18-2) and 725 nm (Schlodder et al. [2007;](#page-18-4) Renger and Schlodder [2008;](#page-17-4) Itoh et al. [2007](#page-16-1)), that is about 15–30 nm red-shifted with respect to 685 nm in canonical Chl *a*-binding PSII. This is equivalent to a decrease of about 60–100 meV at disposal for photochemical reactions, a loss comparable to that experienced by *A. marina* PSI. Substantial compensation for the decrease in the RC excited state energy appears to be provided by a higher redox potential of the active Pheo *a* acceptor that has been titrated at about – 480 mV in manganese depleted *A. marina* PSII (Allakhverdiev et al. [2010\)](#page-15-5) that, after correction, would yield a potential of ~ -410 mV for an intact core complex (Allakhverdiev et al. [2011\)](#page-15-6). The Pheo *a* redox potential in the intact Chl *a*-binding PSII of *Synechocystis sp.* PCC6803 was determined to be -540 mV (Allakhverdiev et al. 2010) in a comparative study. On the other hand, only relatively subtle diferences between the canonical PSII and the Chl *d-*dominated supercomplex of *A. marina* were observed for forward ET and recombination reactions between the quinone acceptors Q_A and Q_B and the donor side (Razeghifard et al. [2005](#page-17-11); Renger et al. [2007;](#page-17-12) Cser and Vass [2007](#page-15-7); Viola et al. [2022](#page-18-13)). Results from kinetic studies are in substantial agreement with direct redox titration of Q_A in manganese depleted PSII of *A. marina*, for which a corrected midpoint potential of ~ –70 mV was estimated (Allakhverdiev et al. [2011\)](#page-15-6), with respect to ~ –140 mV in Chl *a*-binding PSII. Based on the Pheo *a* titration data, the energy diference with Q_A would be therefore approximately 60 meV lower than in canonical PSII, but in both cases, larger than~325 meV. The substantial, although not integral, conservation of the energetic properties of both the acceptor side and the cofactors involved in water splitting might be necessary for proton shuttling through the membrane and for chemical catalysis, respectively.

*Light‑induced damage of A. marina PSII***.**

PSII embedded in the thylakoids of *A. marina* has been shown to be more susceptible to light-induced damage with respect to both canonical PSII and the Chl *a*/*f*-binding PSII of *Chroococcidiopsis thermalis* PCC 7203 (Viola et al. [2022\)](#page-18-13), one of the best-characterised cyanobacterial strain displaying the FaRLiP response. The sensitivity of Chl *d* binding PSII of *A marina* was shown to correlate with an enhanced light-induced singlet oxygen $({}^{1}O_{2})$ sensitisation. This can be in principle rationalised by a larger propensity to populate the RC triplet state either due to the smaller energy gap between Q_A and Pheo *a* (Allakhverdiev et al. [2011](#page-15-6); Viola et al. [2022](#page-18-13)) that shall facilitate the repopulation of the precursor radical pair from which the triplet originates, or to diferent, but yet to be established, energetics causing a marked increase of the lifetime of the [*P*⁺ ⁷²⁵*Pheo*−] radical pair precursor.

The study of the photo-induced triplet states in *A. marina* photosystems, particularly in a relatively intact environment such as the thylakoid membranes, can be helpful in understanding the role of Chl *d* and Chl *a* in photochemical reactions. The triplet states represent specifc spectroscopic markers of the RC pigments, where the triplet population occurs, typically, through the charge recombination mechanism (for reviews see: Budil and Thurnauer [1991](#page-15-8); Lubitz [2002](#page-17-13)). Although triplet–triplet migration can take place, it is a much slower process than singlet energy transfer, at least at low temperatures. The triplet state is then mainly localised on the chromophore on which it is initially populated, or nearby. The study of RC localised triplets, in turn, aids in the identifcation of the cofactors involved in charge separation reactions. Furthermore, Chl triplet states are known to be involved in light-induced photo-oxidative damage, especially to PSII (Durrant et al. [1990](#page-16-15); Vass and Styring [1993](#page-18-14); Telfer et al. [1994](#page-18-15); Telfer [2014](#page-18-16)), since they can sensitise ${}^{1}O_{2}$ production via the interaction of O_{2} that is a triplet ground state (Krasnovsky [1982](#page-16-16), [1994](#page-16-17); Krasnovskiĭ [2004](#page-16-18)). The energy level of ³Chl *d*, albeit lower than that of ³Chl *a*, is sufficient to interact with O_2 (Neverov et al. [2011;](#page-17-14) Hartzler et al. [2014\)](#page-16-19). Singlet oxygen sensitisation is independent of whether the Chl triplet is populated via the recombination mechanism in the RC or directly by intersystem crossing (ISC) from the singlet state. ISC is an active and signifcant decay pathway in Chls (Bowers and Porter [1967](#page-15-9)) that is typically considered to take place in light harvesting pigments. 3 Chl in the antenna are efficiently quenched by triplet–triplet energy transfer to carotenoids (Car). The Car triplet is too low in energy to interact with oxygen and hence, this state can decay harmlessly to the ground state (Wolff and Witt [1969](#page-18-17); Mathis et al. [1979](#page-17-15); Kramer and Mathis [1980](#page-16-20); Siefermann-Harms [1987\)](#page-18-18). Nonetheless, even residual or imperfect quenching by Car, can lead to the presence of

potentially harmful ISC populated ³Chl (Santabarbara et al. [2001](#page-18-19), [2002b;](#page-18-20) Santabarbara and Jennings [2005](#page-18-21)).

The triplet population mechanisms can be discriminated performing time-resolved EPR (TR-EPR) experiments, since the spin polarisation pattern of the light-induced EPR spectrum of ³ Chls is *eeeaaa or eaeaea* for ISC and *aeeaae* for the recombination mechanism (Budil and Thurnauer [1991](#page-15-8); Lubitz et al. [2002\)](#page-17-16). EPR spectroscopy also allows to estimate key features characterising the triplet state, such as the zero feld splitting (zfs) parameters that depend on the chemical nature of the cofactor but can also be modulated by the surroundings that provide solvation and coordination.

On the other hand, most EPR approaches do not give any direct information on the optical properties of the cofactors carrying the triplet state. A notable exception is represented by Optical Detected Magnetic Resonance (ODMR) in zeromagnetic feld, a technique that allows to directly detect the changes in the optical properties of a system due to the change in triplet populations, elicited by resonant microwave frequencies between a couple of triplet sublevels. ODMR is intrinsically a bi-dimensional technique, providing a microwave spectrum when monitoring at a fxed observation wavelength, and an optical Triplet-minus-Singlet (TmS) spectrum, when sitting at a resonance frequency and scanning the observation wavelength. ODMR has superior resolution for zfs determination compared to EPR, because of the absence of an applied magnetic feld, allowing to resolve triplet populations which would be otherwise indistinguishable by EPR, and to simultaneously characterise their associated optical properties (Carbonera [2009\)](#page-15-10). Hence, both ODMR at 1.8 K and TR-EPR (between 10 and 100 K) were here applied in order to characterise the photo-induced Chl triplets in the photosystems embedded in the thylakoid membrane of *A. marina*.

Materials and methods

Growth conditions

A. marina cells were cultivated in a modifed BG11 medium, in which distilled water was replaced by artifcial sea-water (SW-BG11). Cells were initially grown in 250 mL and, when necessary, transferred to 1 L cylindrical fasks, containing 150 and 650 mL of liquid medium, respectively. In both cases the cultures were maintained under continuous aeration (4 L/min) and illumination, provided by 15 W incandescent lamps, giving an average photon fux density of 35 µmoles of photon $m^{-2} s^{-1}$ at the sample level, in a temperature-controlled room at 21 °C.

Thylakoid membranes purifcation

Cells were harvested by centrifugation at 9700x*g* for 5 min in a SS34 rotor (Sorvall), washed in 50 mM HEPES/NaOH pH 7.5 under the same centrifugation conditions and the pellet suspended in the same buffer. For thylakoid membranes preparation, 1–1.2 g of cells were disrupted using a TissueLyzer (Qiagen) in the presence of 212–300 μm glass beads (cells to beads ratio \sim 1:2). ε -aminocaproic acid and phenylmethylsulfonyl fuoride (PMSF) were used as proteases inhibitors. Broken material was frst fltered through Mylar cloth and residual cell debris and glass beads were further removed by centrifugation (11150x*g* for 5 min). The resulting supernatant was centrifuged at 140000x*g* in a 50.2Ti rotor (Beckman) for 45 min and the pellet, containing the thylakoid membranes was suspended in a small volume of 50 mM HEPES/NaOH bufer (pH 7.5), frozen in liquid nitrogen and stored at –80 °C until use. Pigments concentration in thylakoids was estimated by pure methanol extraction using the extinction coefficient of 63.69 mg/mL at 697 nm (Li et al. [2012\)](#page-17-17) in an Uvidec 510 (Jasco) spectrophotometer. All procedures were performed at 4° C and under dim light or darkness.

Measurement conditions

Immediately before measurements, the sample was diluted to a fnal concentration equivalent to 100 μg mL−1 Chl *d* in a bufer containing 150 mM HEPES/NaOH bufer (pH 7.5), 100 mM KCl, 5 mM $MgCl₂$, and a final concentration of 60% v/v of glycerol to obtain a transparent glass at low temperature. For treatments with redox mediators (at the appropriate conditions as further reported in the text and fgures), freshly prepared solutions of the reagents, N,N,N′,N′-Tetramethylp-phenylenediamine (TMPD), sodium ascorbate and sodium dithionite, were initially added to the concentrated thylakoid suspension, in the absence of glycerol, and incubated in the dark, on ice, for at least 15 min, after which a bufered mediators-containing glycerol solution was added to the samples to reach the fnal desired ratio of cryoprotectant/water. All buffers and mediator solutions were degassed and flushed with nitrogen before use.

Sample illumination was performed using a 150 W projector, fltered through 10 cm of water and a heath mirror (Ealing 35–6865). Illumination was either performed at RT or at 77 K, by immersing the sample in an unshielded Dewar vessel.

Dark-adapted and pre-illuminated samples were then immediately transferred to a pre-cooled cryostat at 60 K for ODMR measurement or frozen in liquid nitrogen in the measuring tube for EPR measurements.

Optically detected magnetic resonance apparatus

ODMR experiments were performed in a home-built apparatus which has been previously described in detail (Carbonera et al. [1992;](#page-15-11) Santabarbara et al. [2002a;](#page-17-18) Agostini et al. [2018](#page-15-12)), with some modifcations. In brief, the amplitude modulated microwave feld, generated by a HP8559b sweep oscillator equipped with the HP83522a plug-in device, by a Sco-Nucletudes 10-46-30 TWT amplifer was transmitted trough a semi-rigid coaxial cable to the sample held in a slow-wave helix with a pitch of about 2 mm, acting as broad band resonator. The experiments were performed with a maximum output power of 500 mW. The amplitude modulation frequency was set, if not otherwise diferently indicated, at 33.3 Hz, and selected in order to maximise the ms-lifetime Chl triplet signals.

Optical excitation was provided by a 250 W tungsten lamp (Philips) led by a stabilised power supply. For fuorescencedetected measurements (FDMR) the sample was excited by a combination of broadband flters, consisting in a solution of $CuSO₄ 0.1 M$ (path-length 5 cm), and a heath mirror (Ealing). The light was focused onto a flat cell $(5,0\times1,2 \text{ mm})$ placed in a liquid helium cryostat (Oxford instruments, mod. Spectromag 4) which was pumped to reach the operating temperature of 1.8 K. The sample was kept with the normal to its plane rotated 40° about the cryostat axis, and the emission was collected at 90° through the appropriate cuton or interference filters (FWHM \sim 10 nm) or combinations thereof, with a photodiode (Centronic mod, OSI 5 K). The signal was initially detected with a phase sensitive lock-in amplifer (EG&G mod. 5210), digitised and averaged with a computer-controlled LeCroy LT364 oscilloscope triggered by the microwave sweep oscillator.

For absorption-detected measurements (ADMR) the same apparatus was used, but the beam was fltered through a 10 cm water and was focused on a fat cell (1.2 mm pathlength) placed perpendicular to the light beam. The transmitted light was analysed by a Jobin–Yvon HR250 monochromator, equipped with the Data-Link plug-in controlled by a PC, and acquired by the same photodiode used for the fuorescence experiments. The signal is presented as the ratio of the amplitude modulated and the continuous components $(\Delta I/I)$ of the photodiode signal corresponding to $\Delta A/A$ for small changes in transmission that are generally not larger than 10^{-5} .

Deconvolution analysis of FDMR/ADMR spectra

Deconvolution analysis in terms of Gaussian sub-bands of the fuorescence emission spectra was performed using a global convolution routine, employing a laboratory-developed software, as previously described (Santabarbara et al. [2002a](#page-17-18), [2015;](#page-17-19) Santabarbara and Carbonera [2005\)](#page-17-20). In brief,

the |D|+|E| and |D|−|E| resonance lines of the FDMR spectra, recorded at multiple emission wavelengths, were simultaneously ftted using a linear combination of Gaussians as the model function. The microwave frequency of the Gaussian maxima and bandwidths were considered as global ft parameters and then constrained to assume the same value for the diferent spectral series (|D| +|E| or |D|−|E|). The amplitudes were instead treated as local parameters and hence allowed to change at the diferent emission wavelengths. Yet the ratio of the amplitudes, and of the widths of the Gaussian bands, in the |D|+|E| and |D|−|E| transitions were constrained to be the same. To take into account the diferences in the signal to noise levels between the microwave spectra recorded at diferent emission wavelengths, the root mean square noise over the average signal in microwave regions that did not induce triplet sublevel resonance (offresonance), served as an estimator of data point uncertainties. The search for the best ft parameters was performed with a non-linear least-square algorithm which minimises the global (sum over all measured transitions) χ^2 likelihood indicator, performing an initial search through the Simplex algorithm and a refnement using a Levenberg-Marquard algorithm (Santabarbara et al. [2007,](#page-17-21) [2015](#page-17-19)).

Time‑resolved electron magnetic resonance

TR-EPR experiments were performed on a Bruker ELEX-SYS E580 spectrometer operating at X-band (9.66 GHz), equipped with a dielectric cavity (Bruker ER 4117-DI5, $TE₀₁₁$ mode), an Oxford CF935 liquid helium flow cryostat and an Oxford ITC4 temperature controller. The microwave frequency was measured by a frequency counter (HP5342A). The temperature was controlled by an helium flow system, disabling the magnetic feld modulation and using pulsed sample photo-excitation from a Nd:YAG pulsed laser

Fig. 1 FDMR spectra of *A. marina* thylakoids recorded through a long-pass flter (*λ*>715 nm) in samples that were (**a**) dark-adapted for 15 min at RT or illuminated for 5 min at RT following incubation with (**b**) 10 mM Asc (**c**) 40 mM Asc (**d**) 10 mM Asc and 80 μM TMPD. Measurement conditions: T,1.8 K; mw-amplitude modulation, 33.3 Hz; phase, − 96°; scan rate, 2.5 MHz s^{-1}

(Quantel Brilliant) equipped with both second and third harmonic modules and an optical parametric oscillator (OPO-TECH) (λ =472 nm, pulse length=5 ns, E/pulse \approx 3 mJ, 10 Hz repetition time). The experiments were carried out by recording the time evolution of the EPR signal after the laser pulse with a LeCroy 9360 digital oscilloscope triggered by the laser pulse. At each magnetic feld position, 80 transient signals were usually averaged before transferring them to the PC controlling the instrument via the XEPR software; 250 points on the magnetic feld axis were recorded, with a sweep width of 100.0 mT. The microwave power for TR-EPR experiments was set to 0.47 mW (25 dB attenuation). The time vs. feld surfaces were processed using a homewritten MATLAB program that removes the background signal before the laser pulse (signal vs. magnetic feld) and the intrinsic response of the cavity to the laser pulse at an off-resonance field (signal vs. time). The TR-EPR spectra shown in the main text were extracted from the surfaces at the maximum of the transient (1500 ns after the laser fash) to avoid potential distortions arising from anisotropic relaxations.

Analysis of the TR-EPR spectra and their simulations were performed using Easyspin, a MATLAB toolbox (Stoll and Schweiger [2006\)](#page-18-22)

Results

Fluorescence detected magnetic resonance of chl triplet states

The photo-induced triplet states in thylakoid membranes of *A. marina* were initially investigated by FDMR employing broadband detection through a cut-on flter with half transmission at 715 nm. This corresponds, at the measurement

temperature of 1.8 K, to integrating over almost the entire emission bandwidth. Thylakoids were at frst studied in dark-adapted and redox ambient conditions (Fig. [1\)](#page-5-0), and employing amplitude modulation settings (33.3 Hz) optimised for lifetimes of \sim 1–3 ms, as expected for unquenched Chl triplet states (Krasnovsky [1982\)](#page-16-16). The presence of more than a single triplet population was observed both in the 510–670 MHz (|D|−|E|) and 810–970 MHz (|D|+|E|) transitions. Based on the previous analysis of Chls *a* and *d* in organic solvents (Di Valentin et al. [2007](#page-15-13)), the resonance frequency between 510 – 670 MHz is assigned to Chl *d* |D|−|E| transitions, whereas the Chl *d* |D|+|E| transition is more overlapped to that of Chl *a,* and hence less diagnostic. The |D|−|E| transition of Chl *a* is expected to fall in the 680–820 MHz window instead, where it was not possible to detect any signal under the conditions employed (Figure S1). This could be in part due to the fuorescence being substantially dominated by Chl *d* emission; yet Chl *a* triplet states could still be observed on Chl *d* emission because of singlet–singlet energy transfer, as observed for excited state coupling with long-wavelength emitting forms in other photosynthetic complexes **(**e.g. (Searle et al. [1981;](#page-18-23) Searle and Schaafsma [1992;](#page-18-24) Carbonera et al. [1997](#page-15-14); Santabarbara et al. [2002a\)](#page-17-18)).

Previous ODMR investigations of triplet states in thylakoids of organisms in which Chl *a* is the dominant chromophore involved in photochemical reactions under non-reducing conditions, showed the presence of a Chl *a* triplet population having an unusually short lifetime in the μs time scale (Santabarbara et al. [2002a\)](#page-17-18). Similarly fastrelaxing Chl *a* triplet states were also observed in isolated PSII either by transient spectroscopy (Hillmann et al. [1995](#page-16-21); van Mieghem et al. [1995\)](#page-18-25) or TR-EPR (Feikema et al. [2005](#page-16-22)). In ODMR investigations, this triplet state became dominant under phase sensitive detection at higher amplitude modulation frequencies of the resonant feld (333 Hz) than those employed for ms-lifetime populations (Santabarbara et al. [2002a](#page-17-18), [2003\)](#page-18-26). Measurements performed under these conditions, however, did not reveal either the presence of additional populations that might emerge because of phasesuppression, or any enhancement and/or changes in the FDMR spectral profle (Figure S1) indicating that the triplets observed under non-reducing conditions are all characterised by the typical lifetime of a few milliseconds.

The population of Chl triplet states is typically favoured under reducing conditions that lead to the accumulation of electron equivalents at the photosystems acceptor side, particularly the accumulation of the fully reduced form of the quinones, namely Q_A in PSII and A_1 in PSI. Under such conditions, the singlet radical pair becomes long-lived since electron transfer to the next acceptor in the chain is prevented, thereby promoting spin dephasing to the triplet state (Budil and Thurnauer [1991\)](#page-15-8). The triplet in the RC can

then be populated with high yield by the recombination mechanism.

An efficient method to obtain a complete reduction of the quinone acceptor is that of illuminating the sample in the presence of reducing agents, such as sodium dithionite (Carbonera et al. [1994](#page-15-15), [1997;](#page-15-14) Santabarbara et al. [2002a,](#page-17-18) [2007](#page-17-21)), or to expose the samples to long incubation in the dark in the presence of reducing agents (Frank et al. [1979](#page-16-23); Gast et al. [1983](#page-16-24); Rutherford and Sétif [1990](#page-17-22)). Yet, upon incubation of *A. marina* thylakoids in the presence of dithionite (in concentration of $1-40$ mM, and pH $7-10$) a very rapid change in colour of the suspension was observed, which was also accompanied by the pronounced bleaching of Chl absorption (Figure S2). Although the origin of the degradation is not clear at present, it may relate to some non-specifc redox reactivity between Chl *d* and dithionite that makes the conditions promoting the population of RC triplets in Chl *a*-binding photosystems not readily transferrable to *A. marina*.

In an attempt to overcome this limitation, the quinone photo-accumulation treatment was performed with a milder reducing agent, sodium ascorbate (Asc, 10–40 mM), alone or in the presence of the redox mediator N,N,N′,N′ tetramethyl-para-phenylene-diamine (TMPD, 80 μM). Based on the redox properties of Chl *a*-binding RCs, these sets of conditions shall be sufficient to elicit at least a substantial accumulation of fully reduced Q_A in PSII (Diner and Rappa-port [2002](#page-16-25) and references therein), but shall not be efficient in promoting the full reduction of A_1 in PSI since the acceptor side of PSI operates at much more negative redox potentials than that of PSII (Brettel [1997](#page-15-3) and references therein). Moreover, Asc can also act as an electron acceptor, thereby releasing reducing equivalent accumulation at PSI acceptor side (Trubitsin et al. [2014\)](#page-18-27).

Although illumination in the presence of 10 mM Asc alone was sufficient to induce a significant extent of recombination triplet in a PSII preparation isolated from land plants (Figure S3), the reducing agent alone, even at higher concentrations, did not appear to change by any signifcant extent the FDMR signals, neither in terms of band-shape or intensity, in the thylakoids of *A. marina* (Fig. [1](#page-5-0)). A signifcant change in the relative intensity of the diferent subpopulations was obtained upon illumination in the presence of Asc and TMPD acting as a redox mediator. In particular, there was a strong decrease in the signal of the triplets having a positive amplitude (under the phase convention used in the measurements) and a marked increase in signal intensity in triplets having a negative amplitude. Thus, FDMR spectra obtained in non-reduced/dark-adapted membranes and those pre-illuminated in the presence of Asc and TMPD were studied in further detail.

Fig. 2 Decomposition of FDMR spectra in *A. marina* thylakoids, dark-adapted at RT without the addition of reducing agents, at representative emission wavelengths. Black lines: experimental data; red lines: ftted spectra: dash-dotted lines: Gaussian sub-bands, orange (T_1) , dark yellow (T_2) , green (T_3) , blue (T₄). Triplets T₁–T₄ are defined as in Table [1](#page-9-0). Measurements conditions as in the legend of Fig. [1](#page-5-0)

Emission wavelength dependence of FDMR spectra

Figures [2](#page-7-0) and [3](#page-8-0) show the FDMR spectra recorded at a few selected emission wavelengths using interferential flters $(FWHM \sim 10 \text{ nm})$. Also shown in the figures is the Gaussian sub-bands decomposition of the FDMR spectra by a constrained global ftting approach (described in Material and Methods) in which the spectra acquired in the two diferent conditions discussed above are simultaneously ftted. This approach allowed the identifcation of a total of four triplet populations, characterised by the centre frequency, bandwidth, and associated zfs parameters reported in Table [1.](#page-9-0) The relative abundances of these populations were found to largely depend on the treatments performed at RT before freezing. In dark-adapted thylakoids under non-reducing conditions (Fig. [2](#page-7-0)), three triplets were mainly observed (labelled as a T_1 , T_2 and T_3 , zfs: |D|: 0.0252 cm⁻¹, |E|: 0.0049 cm−1; |D|: 0.0245 cm−1, |E|: 0.0049 cm−1; |D|: 0.0245 cm⁻¹, |E|: 0.0042 cm⁻¹, respectively), with the frst two having almost equal amplitude and being overall the most represented at all the emission wavelengths investigated, whereas the T_4 population was fundamentally negligible. In thylakoids illuminated at RT in the presence of Asc and TMPD the T₃ (|D|: 0.0245 cm⁻¹, |E|: 0.0042 cm⁻¹) and T₄ (|D|: 0.0248 cm⁻¹, |E|: 0.0040 cm⁻¹) populations became the most represented instead, particularly T_3 being the dominant one, whereas both T_1 and T_2 undergo an evident decrease in amplitude, with the latter being almost completely suppressed (Fig. [3](#page-8-0)).

The significant increase of the T_3 and T_4 populations under conditions expected to lead to the accumulation of reducing equivalents, at least at the PSII donor side, would seem to suggest that these Chl *d* triplets are associated with RC processes. It is yet worth nothing that the observed increase in the intensity of FDMR signals associated with T_3 (and T_4) under reducing conditions, although significant, is far less pronounced than that previously reported for Chl *a*-binding photosystems, either isolated (Carbonera et al. [1997](#page-15-14)) or embedded in the thylakoid membranes (Carbonera et al. [1997;](#page-15-14) Santabarbara et al. [2002a](#page-17-18), [2003](#page-18-26), [2007](#page-17-21)), albeit observed under stronger reducing conditions. This may be due to an only partial reduction of the photosystems'

Fig. 3 Decomposition of FDMR spectra in *A. marina* thylakoids, illuminated at RT in the presence of Asc (10 mM) and TMPD (80 μM), recorded at representative emission wavelengths. Black lines: experimental data; red lines: ftted spectra: dash-dotted lines: Gaussian sub-bands, orange (T_1) , dark yellow (T_2) , green (T_3) , blue (T_4) . Triplets $T_1 - T_4$ are defined as in Table [1](#page-9-0). Measurements conditions as in the legend of Fig. [1](#page-5-0)

acceptors. The poising conditions employed are expected to efficiently reduce the PSII acceptor, but not to be very efective on PSI terminal acceptors, as they operate at much more negative potential values $(< -500$ mV). Hence, to gain further insight into the nature of the triplet populations which displayed the largest sensitivity to the redox/illumination pre-treatment, the samples were also investigated by absorption-detected magnetic resonance.

Absorption‑detected magnetic resonance of chl d triplets

Figure [4](#page-10-0) shows the ADMR spectra recorded in the |D|−|E| transition in *A. marina* thylakoids pre-illuminated at RT in the presence of Asc and TMPD, at selected wavelengths in the Q_v transition. Also shown in Fig. [4](#page-10-0) is the decomposition in Gaussian sub-bands, which demonstrates that the ADMR signal was almost entirely due to the two already discussed **Table 1** Parameters of Gaussian decomposition of ODMR spectra of *A. marina* thylakoids

In Table [1](#page-9-0) are reported the values of the maxima and the widths of the Gaussian functions resulting from the global, joint, deconvolution analysis of the FDMR and ADMR spectra recorded in *A. marina* thylakoids dark-adapted under non-reducing and room temperature-illuminated under mild-reducing conditions (10 mM Asc+80 μm TMPD). Also indicated are the zero feld splitting (zfs) parameters |D| and |E|. Errors are in the zfs \pm 0.0001 cm⁻¹

populations T_3 and T_4 , with the former being the dominating species and T_4 accounting on average for $\sim 15-25\%$ of the overall ADMR signal depending on the detection wavelength.

Further details can be gained by recording the Tripletminus-Singlet (TmS) spectrum. Figure [5](#page-10-1) shows the TmS spectra obtained upon microwave excitation in the |D|−|E| transition at 610 MHz. The spectrum displays a broad and asymmetric maximal singlet bleaching centred at about 740 nm, accompanied by a complex set of satellite bands, the most clearly resolved being two sharp derivative-like features having maximal bleaching at 732 and 722 nm and positive companion bands at 728 and 717 nm, respectively. Other side bands are observed at 708 and 697 nm.

The TmS spectrum in Fig. [5](#page-10-1) has some significant similarities with the one recorded in highly PSI enriched particles from *A. marina* by transient optical spectroscopy at 5 K (Schenderlein et al. [2008](#page-18-10)). This signal became detectable upon pre-illumination of the sample at 230 K in the presence of dithionite and phenazine methosulfate (PMS) as a mediator and was attributed to the PSI-RC triplet, ${}^{3}P_{740}$. However, the TmS spectrum recoded here by ODMR in thylakoids at 1.8 K shows some additional spectral features. In particular, the sharp bleaching at 732 was not resolved in the previously reported spectrum and the exact position of the satellite bands, with the positive one at \sim 725 nm and also the side bands at \sim 705 nm, appearing more defined in the mw-induced TmS. The transient optical TmS showed a bleaching at ~ 685 nm, which is, however, not resolved here in the mw-induced one (Fig. [5\)](#page-10-1).

According to the decomposition of the ADMR spectra (Fig. [4](#page-10-0)), at the pump mw used to record the TmS spectrum (610 MHz) the T₃ ($|D|=0.0245$ cm⁻¹, $|E|=0.0042$ cm⁻¹) population should be by far the dominant one. Hence, although it is possible that some of the fne spectral structure may be the result of partial excitation of the T_4 population, this will require the TmS associated to each of these two states to be signifcantly diferent. The relative contribution of the T_3 and T_4 in ODMR spectra at different wavelengths appears to be almost the same (Figs. [3](#page-8-0) and [4](#page-10-0)) instead. It has not been possible, however, to record a TmS spectrum with preferential excitation of T_4 (mw > ~630 MHz) because of the signal weakness. Nonetheless, it appears unlikely, albeit not impossible, that the additional features in the ${}^{3}P_{740}$ TmS spectrum observed by ODMR with respect to the transient optical methods stem from lack of selective excitation, especially considering that optical excitation does not allow for triplet population selections, unless the populations have sufficiently different lifetimes.

The two dominating triplets, T_3 and T_4 , observed after pre-illumination of *A. marina* thylakoids under mildly reducing conditions, most probably represent two subpopulations of a triplet state located on the same chromophore, with the subtle variation in zfs and triplet sub-level resonance frequency resulting from microscopic heterogeneity in the protein-cofactor coordination. Similar heterogeneities have been previously observed in ODMR investigations of ${}^{3}P_{700}$ in Chl *a*-binding PSI, in both isolated photosystems (Carbonera et al. [1997\)](#page-15-14) and more intact environments (Santabarbara et al. [2002a](#page-17-18), [2007](#page-17-21)).

In Fig. [5,](#page-10-1) a decomposition of the TmS spectrum is also shown, with the ft parameters reported in Table [2.](#page-10-2) According to the Gaussian decomposition the main singlet bleaching bands are centred at 740 nm, 731 nm and 722 nm. As readily visible directly from the TmS spectrum inspection, the bleaching bands at 731 nm and 722 nm are relatively narrow (FWHM 4–4.5 nm), whereas the 740 nm is fairly broad (FWHM \sim 30 nm). The positive companion bands appear relatively narrow (3.5–7 nm, Table [2\)](#page-10-2) as well. The obtained sub-bands shall refect the shift and population redistribution of the singlet exciton (eigen) states upon population of the RC triplet state. Yet, the exact positions derived from ftting are subjected to some uncertainties, because of cross-compensation of positive/ negative amplitude sub-bands, therefore, the description shall be considered as a frst-order approximation only (an alternative solution is presented in the Supplementary Information, with slightly diferent band centres and widths, but displaying an overall consistency, Figure S4).

The broadness of the red-most transition, typically centred at 700–705 nm, is a characteristic also of Chl *a-*binding PSI (Carbonera et al. [1997\)](#page-15-14). Although the

Fig. 4 Decomposition of ADMR spectra in *A. marina* thylakoids, illuminated at RT in the presence of Asc (10 mM) and TMPD (80 μM), recorded at representative absorption wavelengths in the Chl *d* Qy band. Black lines: experimental data; red lines: ftted spectra: dash-dotted lines: Gaussian sub-bands, orange (T_1) , dark yellow (T_2) , green (T_3) , blue (T_4) . Triplets $T_1 - T_4$ $T_1 - T_4$ $T_1 - T_4$ are defined as in Table 1. Measurements conditions as in the legend of Fig. [1](#page-5-0), except that the phase was $+80^\circ$

exact bleaching positions and band widths are in part species dependent (Witt et al. [2003](#page-18-7); Nakamura et al. [2011](#page-17-7)), the width is most likely afected by a large contribution from homogenous broadening that depends on the electron–phonon coupling, rather than an unusually large heterogeneous broadening (site distribution). The large

Fig. 5 Microwave-induced TmS spectrum obtained upon excitation at 610 MHz, in *A. marina* thylakoids, illuminated at RT in the presence of Asc (10 mM) and TMPD (80 μM). Black lines and open symbols: experimental data; red lines: ftted spectra; dash-dotted lines: Gaussian sub-bands, with green shades representing singlet bleaching and yellow-orange shades describing the increase in absorption upon triplet population; the ft parameters are reported in Table [2.](#page-10-2) Measurements conditions as in the legend of Fig. [1,](#page-5-0) except that the phase was +80 $^{\circ}$ and the wavelength scan rate was 0.33 nm sec⁻¹

Table 2 Gaussian decomposition of mw-induced TmS spectrum in *A. marina* thylakoids

Band	Centre (nm)	FWHM (nm)	Amplitude
1	644.5	24.7	0.081
2	699.5	15.0	0.382
3	707.9	4.1	0.738
$\overline{4}$	717.4	7.2	0.643
5	722.1	4.2	-0.428
6	729.6	3.4	0.950
7	730.9	4.5	-0.784
8	740.1	29.7	-0.692
9	757.4	24.2	-0.067

In Table [2](#page-10-2) are reported the parameters employed in the description of the TmS spectrum obtained by microwave excitation at 610 MHz (|D|–|E|) and assigned to a triplet population residing in one or being shared between chromophores composing the PSI reaction centre () of *A. marina*

homogeneous broadening can in principle be due to coupling with a higher (mean) frequency mode with respect to Chls showing a narrow bleaching, to a larger Hung-Rhys, *S*, factor or both (e.g. Gillie et al. [1989](#page-16-26); Jankowiak and Small [1993;](#page-16-27) Jankowiak et al. [2011\)](#page-16-28). For Chl *a*-binding PSI a large value of S (\sim 2–4) was suggested to result from the

Fig. 6 X-band triplet TR-EPR spectra of *A. marina* thylakoids that were (**a**) dark-adapted for 15 min at RT or (b) illuminated at RT in the presence of Asc (10 mM) and TMPD (80 μM). The simulated (**c**) ISC and (d) recombination 3 Chl *d* spectra have been calculated using the following parameters: $D=25.8$ mT, $E=-5.35$ mT, D strain=2.5 mT, E strain=1.5 mT, isotropic linewidth=1.0 mT, microwave frequency 9.649 GHz. For the simulation of the (c) ISC triplet, a polarisation of $(P_x : P_y : P_z) = (0.33 : 0.43 : 0.24)$ has been employed. The spectra have been vertically shifted for a better comparison. *a* absorption, *e* emission

mixing with a charge-transfer state (Lathrop and Friesner [1994;](#page-17-23) Renger and Schlodder [2006](#page-17-24)) that in turn might be favouring a radical pair formation. If this were the case, the broadness of the main bleaching band at 740 nm in the PSI of *A. marina* seems to suggest a conservation of the partial charge-transfer character, as the main bleaching appears to be as broad, or broader, than in standard PSI.

Time‑resolved electron paramagnetic resonance

TR-EPR experiments were performed to further characterise the mechanism of triplet population in the thylakoids of *A. marina*. The two conditions in which the membranes were either dark-adapted at ambient conditions or pre-illuminated at RT in the presence of Asc and TMPD were tested by recording the TR-EPR spectra at 50 K, upon laser excitation at 472 nm. Similar results were obtained also performing the measurements at 10 and 100 K (not shown).

Figure [6](#page-11-0) shows the TR-EPR spectra obtained in the two investigated conditions. In both spectra, a narrow signal was observed in the centre of the spectrum $(g \sim 2.00)$, magnetic field \sim 344 mT), that can be attributed to spin-correlated radical pairs. Its characterization is beyond the scope of the present investigation.

Under non-reducing conditions, the TR-EPR spectrum was dominated by a triplet state contribution characterised by the polarisation pattern expected for a triplet state populated by the ISC mechanism (*eaeaea*), which would then be most likely populated in the photosystems' antenna. This triplet population(s) shall correspond, mainly, to the T_1 and $T₂$ retrieved by ODMR (Fig. [2,](#page-7-0) Table [1](#page-9-0)), also noticing that $T₃$ was already detectable under these conditions. However, the polarisation pattern of the triplet states observed in the membrane illuminated under mildly reducing conditions was still *eaeeaea* meaning that, also in this case, the triplets become populated by the ISC mechanism. On the contrary, the recombination mechanism should lead to an *aeeaae* pattern (see Fig. [6](#page-11-0)). This is a surprising result, since the conditions used should promote the recombination-induced RC triplet population. ODMR clearly detected a redistribution of the triplet populations after the pre-illumination treatment, with T_3 becoming the dominant one, followed by T_4 (Fig. [2,](#page-7-0) Table [2\)](#page-10-2), therefore, these are most likely the triplets also observed by TR-EPR. Yet, the treatments were not accompanied by a change in the population mechanism of the triplet states. The zfs parameters determined from the EPR spectra are compatible with those derived from the FDMR deconvolution (Table [2](#page-10-2)), but unfortunately, the intrinsically lower zfs resolution of EPR and the diference in temperature do not allow to discern between the four components identifed in the FDMR datasets. It is worth mentioning that a suppression of the recombination triplet signal like the one previously reported for the heliobacterial reaction centre, in which the recombination triplet $({}^{3}P_{800})$ was not spin polarised because of the D/E ratio being close to 3, with consequent cancellation of the absorptive and emissive contributions to the spectrum (Ferlez et al. [2017](#page-16-29)), can be excluded on the basis of the D and E values determined in ODMR (see Table [1\)](#page-9-0), that are characterised by a D/E ratio between 5 and 6 in all cases.

Discussion

The investigation of photo-induced chlorophyll triplet states in the isolated thylakoid membranes of *A. marina* revealed some interesting peculiarity of the mechanisms of population of these excited states in Chl *d*/*a* binding photosystems. The characterization remains, for the time being, limited by the difficulties in employing the reducing conditions commonly used for Chl *a*-binding photosystems. This is due to the rapid degradation of the *A. marina* photosystems, manifesting as a strong, non-specifc, absorption bleaching accompanied by an obvious change in colour of the thylakoid suspension and loss of both detectable ODMR and EPR signals. Hence, the investigation was restricted to the use of mildly reducing conditions, accompanied by illumination at room temperature.

Photo‑induced triplet states in A. marina thylakoids under non‑reducing and moderately reducing conditions: absence of signal attributable to 3 Chl a or 3 PSII‑RC

Under none of the conditions investigated in this study it was possible to observe photo-induced Chl *a* triplet states. Instead, all the ODMR and TR-EPR triplet signals were assigned to Chl *d*, based on their zfs parameters and reso-nance frequencies (Figs. [1,](#page-5-0) [2,](#page-7-0) [3](#page-8-0) and [6](#page-11-0)), Table [1\)](#page-9-0).

In Chl *a-*binding PSII-RC a fast-decaying (microsecond) triplet state was reported to be populated (Santabarbara et al. [2002a,](#page-17-18) [2003](#page-18-26)) even under non-reducing conditions and was assigned to the recombination triplet state. The decrease of the triplet state lifetime under these conditions was suggested to be due to the presence of a so-called "observer spin" (or radical) on Q_A^- in proximity of the Chl *a* cluster composing the RC (Hillmann et al. [1995;](#page-16-21) van Mieghem et al. [1995](#page-18-25)). Yet, in *A. marina* thylakoids it was not possible to detect any, neither Chl *a* nor *d*, fast-decaying triplet. Together with the detection of long-lived 3 Chl *d* only, this observation might be taken as a favourable evidence for Chl *d*, rather than Chl *a*, being the main, or even the only, photoactive pigment both in PSI and PSII of this organism. However, the absence of any signifcant long-lived ³ Chl *a/d* attributable to the PSII-RC after illumination under mildly reducing conditions is puzzling. This treatment was tested and proved to be efficient in promoting recombination triplet population in canonical Chl *a*-binding PSII (Figure S3) and was then expected to be efective also in *A. marina* since the redox properties of its PSII acceptor side were shown to be almost the same as those of canonical PSII (Cser and Vass [2007](#page-15-7); Cser et al. [2008;](#page-15-16) Viola et al. 2022). Moreover, the reported shift of ~ 60 mV in the redox potential of Pheo *a* in *A. marina* PSII should lead to a decrease in the energy gap with Q_A and hence potentially favour, with respect to canonical Chl *a* PSII, the repopulation of the [*P*⁺*Pheo*−] radical pair that, being the precursor for ³PSII-RC, is expected to increase the overall triplet yield in the Chl *d*-dominated RC. A higher triplet population was proposed as the cause of the increased sensitivity to lightinduced damage and propensity in ${}^{1}O_{2}$ production in this organism with respect to Chl *a-* and Chl *a*/*f*-binding PSII (Viola et al. [2022\)](#page-18-13). The results obtained here in thylakoids do

not appear, however, to indicate a higher yield of ³PSII-RC in *A. marina*. It is possible that the increased sensitivity to damage and the proposed higher PSII-RC triplet population, might be observable at RT but not at the cryogenic temperatures employed in ODMR (1.8 K) or TR-EPR (10–100 K). It is worth noting that these are however necessary conditions to observe 3 PSII-RC by EPR methods, that in Chl *a*-binding RCs are typically in excellent correlation with optical measurements that can be performed also at physiological or close-to-physiological conditions.

Although it is not possible, at present, to unambiguously determine the reasons leading to ³ Chl in PSII of *A. marina* escaping detection, it is nonetheless possible to advance some hypothesis. One possibility is that the PSII acceptor side in thylakoids is, to some extent, only partially accessible to the redox reagents and mediators; this, in turn, will make the pretreatment employed in this study inefficient. Another possibility, which is however not very likely especially at 1.8 K where ODMR experiments are performed, that the RC triplet is populated but escapes detection due to unusually fast spin–lattice relaxation. Finally, it may be considered the hypothesis that the recombination triplet is not efficiently populated because the other de-excitation routes of the precursor radical pair, such as recombination to either the excited or the ground state, dominate in the PSII of this organism.

Studies on PSII of *A. marina* on the isolated, purifed, supercomplex will be necessary to interpret the absence of 3 RC formation even under conditions that should, in principle, promote it.

Photo‑induced 3 Chl d triplet states in A. marina thylakoids under non‑reducing and moderately reducing conditions are dominated by PSI‑bound chromophores

The combination of FDMR and ADMR analyses, in particular the TmS spectrum recorded upon mw excitation at 610 MHz (|D|–|E|), seems to indicate that, at least under moderate reducing conditions, the detected ³ Chl *d* states (populations T_3 and T_4) in *A. marina* thylakoids belong to PSI chromophores. The assignment is based on specifc spectral features, like the maximum broad bleaching at 740 nm, and the intense flanking bands at \sim 730 and \sim 720 nm, being shared by the mw-induced TmS spectrum (Fig. [5\)](#page-10-1) and the spectrum previously obtained by transient optical spectroscopy in highly enriched PSI particles (Schenderlein et al. [2008](#page-18-10)). The sensitivity to illumination under mild-reducing conditions and the complex spectral shape would suggest that T_3 (and likely also T_4 that might be only a subtly different conformer) sits on a Chl *d* that is part of PSI-RC. However, diferently from what may be expected in view of the response to the (mild) redox poise, this triplet state does not

show the characteristic *aeeaae* polarisation pattern resulting from the almost pure population of the T_0 level when the triplet stems from a singlet radical pair state precursor, that is, by the recombination mechanism. It rather displays the *eaeaea* polarisation pattern that is instead compatible with the ISC mechanism (Di Valentin et al. [2007](#page-15-13)). This is actually the same electron spin polarisation observed under non-reducing conditions (Fig. [6\)](#page-11-0) when the dominating triplets $(T_1$ and T_2) are most likely populated in the antenna.

The triplet assigned to PSI-RC (T_3) is already detected, although not being the dominant species, under non-reducing conditions (Fig. [2\)](#page-7-0). This is similar to what previously reported for both isolated Chl *a*-binding PSI (Carbonera et al. [1997](#page-15-14)) and thylakoids embedded PSI (Santabarbara et al. [2002a,](#page-17-18) [2007](#page-17-21)) that, moreover, also showed a heterogeneity of resonance transitions, distinguishable by FDMR because of the very high sensitivity of the technique to even very subtle variations in the zfs parameters.

The significant damping of the populations T_1 and T_2 that, under the mildly reducing conditions employed, correlates with the increase of the populations T_3 and T_4 , suggests that they originate from the same photosystem, being in competition for the localisation of the singlet excited state on the Chl *d* molecule(s) from which the triplet is then formed. Similar behaviours have been previously observed in Chl *a-*binding PSI (McLean and Sauer [1982;](#page-17-25) Carbonera et al. [1997\)](#page-15-14). However, it shall be noted that the fuorescence emission spectra of PSI and PSII of *A. marina* are not quite as distinguished as those of Chl *a-*binding super-complexes, where PSI shows a red-shift of at least 20 nm, and commonly as large as 30–40 nm, with respect to PSII in most model organisms (e.g. for reviews Croce and van Amerongen [2013;](#page-15-17) Santabarbara et al. [2020](#page-18-28)). In this respect, the emission wavelength selectivity ofered by FDMR is less unambiguous for the Chl *d/(a)*-binding photosystem. It is, therefore, not possible to exclude completely that one of the T_1 and T_2 population, or both, might stem, at least in part, from PSII antenn. Yet, the observed decrease in population after illumination under mildly reducing conditions still seem to indicate a dominance of PSI-bound chromophores to the triplet populations detected.

Triplet states in Chl d‑binding PSI‑RC of A. marina: an alternative population mechanism?

The surprising observation that the ³Chl *d* population that increases upon illumination under mildly reducing conditions, namely T_3 , shows a TmS spectrum that closely resembles the one previously reported and assigned to PSI-RC (Schenderlein et al. [2008\)](#page-18-10), ${}^{3}P_{740}$, but manifests an electron spin polarisation resulting from ISC, deserves further attention and discussion.

One hypothesis to reconcile this apparently contradicting evidence is that the observed bleaching in the TmS spectrum at 740 nm does not correspond to the same molecule, or cluster of excitonically coupled molecules, giving rise to the bleaching at 735–740 nm upon light-induced cation formation (P_{740}^+) . The TmS and the P⁺-minus-P spectra are not expected to be exactly the same because, whereas ${}^{3}P$ is electrically neutral and the spectral feature shall originate from excitonic interaction only, the charged nature of P^+ (and often the concomitant presence of a reduced acceptors, *A*[−]) gives rise to local electric feld manifesting a Stark efect (also called electrochromic) on pigments that also contribute to the diference spectrum. Thus, the diferences in fne structure of the previously reported P^+ -minus-P and ${}^{3}P$ -minus-P spectra (Schenderlein et al. [2008](#page-18-10)), with the main bleaching being located almost at identical positions, might just be due to the contribution from local Stark band-shifts. It can not however be completely excluded, instead, that the triplet sits on a diferent (cluster of) chromophore, having the lowest singlet state at almost the same energy as P_{740} . This would, in other terms, represent a low-energy Chl *d* spectral form (red form) in *A. marina* PSI antenna, being almost iso-energetic with the lowest RC excited state. Red forms are characteristics of many Chl *a*-binding PSI (Croce and van Amerongen [2013](#page-15-17); Santabarbara et al. [2020\)](#page-18-28). Their presence has been reported in the isolated PSI of *A. marina* both by low temperature optical spectroscopy and the associated structure-based spectral-kinetic modelling (Kimura et al. [2022\)](#page-16-30). On the other hand, the response of the T_3 population to illumination in the presence of mediators does not corroborate this hypothesis, because the lowest energy state shall remain, at low temperature, a (local) singlet exciton trap irrespectively of the reduction state of the photosystem acceptor or donor side.

The second hypothesis, that we tend to favour, is that the $T₃$ population does actually represent a triplet sitting on one of the Chl *d* present in the PSI-RC, but being populated with a diferent mechanism than in Chl *a-*binding PSI, *i.e.* not through the recombination mechanism. The recombination mechanism can be expressed, in simplifed kinetic terms, by the following generalised scheme:

$$
RC^* \xrightarrow{k_{\infty}} \left({}^{1}[P^{\dagger}A^{-}] \xleftarrow{\circ} \begin{pmatrix} {}^{3}[P^{\dagger}A^{-}] \\ \downarrow \\ {}^{3}P \end{pmatrix} \right)
$$

Where ${}^{3}[P^{+}A^{-}]$ and ${}^{1}[P^{+}A^{-}]$ are the triplet and singlet state of the spin-correlated radical pair, respectively, formed with an effective rate constant k_{css} from the reaction centre excited state RC^* . ω is the singlet–triplet mixing frequency of the radical pair, and the triplet becomes fnally localised

on ³*P*, with *P* being one (or more) of the chromophores composing the RC, having the lowest triplet energy. This mechanism is kinetically favoured when ω is much faster than the rate of direct recombination from $1[P^+A^-]$ to the reaction centre excited state *RC*[∗], which is shown, but unlabelled, in scheme 1, to stress that it represents a "slow" kinetic process.

If the recombination rate to the singlet state (k_{cr}) becomes signifcant or even dominant, the kinetic scheme can be modifed as:

$$
\begin{pmatrix}\nRC^* \frac{k_{\text{cos}}}{k_x} \left[P^* A^- \right] \\
\downarrow \\
{}^{3}P\n\end{pmatrix}\n\longleftrightarrow {}^{3}[P^* A^-]\n\begin{pmatrix}\n\downarrow & \downarrow & \downarrow \\
\downarrow & \downarrow & \downarrow \\
{}^{3}P\n\end{pmatrix}
$$
\nScheme 2

In this case, repopulation of *RC*[∗] would kinetically outcompete the population of the radical pair triplet state. Then, under this scenario, the triplet would be populated by ISC, analogously to the mechanism observed for Chl molecules in the antenna.

An increase in the rate of RC singlet state repopulation can be caused by a relatively small free energy diference, making the population of the radical pair rapidly reversible. Photochemical mechanisms involving a substantial equilibration between the primary radical pair and the reaction centre (or antenna bed) excited state have been extensively discussed, for instance by Holtzwart and cowokers (Schatz et al. [1988\)](#page-18-29), initially in Chl *a*-binding PSII and have been successively extended to PSI (Müller et al. [2003](#page-17-26), [2010](#page-17-27)). Yet, following primary photochemistry, the charge separated state was discussed as being stabilised through the almost irreversible formation of a secondary radical pair, either by electron or hole transfer. Recombination to the triplet state in the RC takes place from the latter "stabilised", and hence relatively long-lived, radical pair. In Chl *a*-binding PSI this state corresponds to $[P_{700}^+ A_0^-]$, where A_0 is the Chl *a* at the structural eC_3 position, acting as the electron acceptor. The proposed RC triplet population in the PSI of *A. marina* by ISC would require that the energy gap between *RC*[∗] and the corresponding radical pair to be much smaller than in canonical PSI, particularly considering that at the ODMR measurement condition of 1.8 K the thermal energy for uphill transfer is only 0.15 meV, and hence the radical pair would have to be almost iso-energetic with RC excited state.

A decrease in the driving force for primary photochemistry and successive charge stabilisation reactions might be expected considering the lower photon energy of P_{740}^* with respect to P_{700}^* , which is in the order of ~95–100 meV. On the other hand, the structural model of *A. marina* PSI showed the presence of Pheo a at the eC_3 position. The triplet precursor radical pair in *A. marina* PSI should then be

[*P*⁺ ⁷⁴⁰*Pheo*−]. The reduction midpoint potential of Pheo *a* in organic solvents is 100–200 mV more positive than that of Chl *a* (Kobayashi et al. [2013](#page-16-13)), which in principle can even over-compensate the photon energy loss in P_{740}^* . Hence, for the proposed mechanism for PSI-RC triplet proceeding through ISC to hold true, and the energy gap between *RC*[∗] and $[P_{740}^{+}Pheo^{-}]$ to be particularly small, the potential of Pheo *a* at the eC_3 site shall be tuned by the photosystem subunits to signifcantly more reducing values than expected based on the comparisons with Chl *a* in organic solvents. A small energy gap cannot instead be ascribed to diferences in the oxidation potential of P_{740}^+ and P_{700}^+ , that differ only by about 10 mV as estimated by direct titration (Schenderlein et al. [2008](#page-18-10)).

It shall also be worth considering that, most studies that discussed a reversible radical pair mechanism in canonical PSI, postulated the population of a primary radical pair, acting as precursor to $[P_{700}^+ A_0^-]$ (Müller et al. [2003](#page-17-26), [2010](#page-17-27); Holzwarth et al [2005](#page-16-31); Giera et al. [2010](#page-16-32); Molotokaite et al. [2017;](#page-17-28) Russo et al. [2020](#page-17-29)). The energy gap in between *RC*[∗] and the primary radical pair, comprising the cofactors at the $eC₂$ position is therefore smaller than between the further stabilised pair. Thus, the decrease in the driving force for the primary charge separation could favour in *A. marina* recombination to RC^* starting from the primary charge separated state rather than from the secondary $[P_{740}^{+}Pheo^{-}]$.

Although the proposed mechanism provides a rationale for the observation of the *eaeaea* instead of the *aeeaae* polarisation pattern for a triplet state residing on the RC, it remains to be validated by experimental approaches through which the free energy difference between RC^* and [*P*⁺ 740*Pheo*−] could be assessed (*i.e.* ultrafast transient absorption, delayed luminescence).

Conclusions

The investigation of triplet states by ODMR and TR-EPR in the thylakoid membranes of *A. marina* allowed to identify the presence of four diferent triplet populations, which can all be assigned to Chl *d* based on their microwave resonance frequencies and associated zfs parameters. All ³Chl *d* populations appear to be long-lived, having a ms-lifetime. On the other hand, neither short- or long-lived Chl *a* triplet states could be detected under the conditions employed.

The ³Chl *d* populations display different behaviour in response to the treatment in the presence of mild redox poise and pre-illumination at RT. In particular, a population characterised by zfs $|D|=0.0245$ cm⁻¹ and $|E|=0.0042$ cm⁻¹ increases and becomes dominant following sample illumination in the presence of a weak reductant. Its associated TmS spectrum, detected at a resonant frequency of 610 MHz

at 1.8 K, resembles the one previously attributed to ${}^{3}P_{740}$, the PSI reaction centre triplet, by optical spectroscopy at 5 K (Schenderlein et al. [2008](#page-18-10)). Moreover, the microwaveinduced TmS spectrum presents additional fne spectral features. Surprisingly, TR-EPR spectroscopy indicates that this triplet is populated through ISC rather than the recombination mechanism. A possible explanation for this apparent contradiction is proposed, based on kinetic competition between the singlet–triplet mixing and the repopulation of the RC singlet excited state, from the precursor radical pair $[P_{740}^{+}Pheo^{-}]$. Population of ³ P_{740} by ISC can become efficient if recombination to *RC*[∗] is the dominant kinetic process. This would in turn imply a signifcantly lower energy gap between *RC*[∗] and the precursor radical pair state than in canonical, Chl *a*-binding, PSI.

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Data Availability Data sets generated during the current study are available from the corresponding author on reasonable request.

Declarations

Competing interests The authors declare no competing interests.

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