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Twisted Carotenoids Do Not Support Efficient Intramolecular Singlet Fission in the Orange Carotenoid Protein

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individual carotenoid molecules within protein complexes provided that the conjugated backbone is twisted out of plane. However, this hypothesis has been forwarded only in protein complexes containing multiple carotenoids and bacteriochlorophylls in close contact. To test the hypothesis on twisted carotenoids in a "minimal" one-carotenoid system, we study the orange carotenoid protein (OCP). OCP exists in two forms: in its orange form (OCPo), the single bound carotenoid is twisted, whereas in its red form (OCPr), the carotenoid is planar. To enable room-temperature spectroscopy on canthaxanthin-binding OCPo and OCPr without laser-induced photoconversion, we trap them in a trehalose glass. Using transient absorption spectroscopy, we show that there is no evidence of long-lived triplet generation through intramolecular singlet fission despite the canthaxanthin twist in OCPo.



C inglet exciton fission (SF) is the conversion of a spin-0 \bigcirc singlet exciton¹ (or excited singlet state) into a pair of spin-1 triplet excitons.²⁻⁵ This multiexciton generation process has been studied over the past decade primarily because of its promise to improve solar cell efficiency;⁶⁻¹⁰ one high-energy photon creates two low-energy excited states, which could be harvested by conventional photovoltaic devices in a process minimizing energetic losses due to thermalization. SF has other potential applications for nonlinear optics,¹¹⁻¹³ organic lightemitting diodes,¹⁴ or even quantum technologies^{15–18} by taking advantage of the virtue that a single photon creates a pair of spin-entangled quantum states. However, despite promising results,^{19,20} practical applications have yet to be realized, in part due to the limited library of materials that undergo SF, none of which is yet ideal.^{4,7}

In the search for other SF materials, the polyenes, "class III" SF materials according to Smith and Michl's categorization, form an intriguing materials class. In these materials, the lowest-lying singlet excited state (S_1) has dominant triplet-pair character, denoted $^{1}(TT)$ (see refs 3, 21, and 22), and thus demonstrates negligible one-photon absorption from the ground state. S1 is instead accessed by internal conversion following excitation to the strongly absorbing S₂ state.

This SF class includes conjugated polymers such as polydiacetylene,^{23–25} poly(alkylthienylenevinylene),^{26–28} a new generation of donor–acceptor singlet fission polymers,^{29–32} quinoidal thiophenes,^{33–37} carbene-based diradica-loids,³⁸ and antiaromatic core-structured molecules.^{39,40} The polyene family also includes the carotenoids, a large class of over 1000 naturally occurring molecules,^{41,42} represented here by canthaxanthin (CAN), which forms the subject of this work (see structure in Figure 1a).

In comparison with better-studied "class I" SF materials, $^{2,46-49}$ mostly based on molecules such as pentacene 50,51 or tetracene, $^{52-55}$ SF in polyenes is less well understood. This is attributable in part to their complex manifold of low-lying triplet-pair states⁵⁶⁻⁵⁸ and strong vibronic coupling^{59,60} and also partly due to the sensitivity of the photophysics to conjugation length and molecular geometry. In polyenes, the lowest-lying ${}^{1}(TT)$ state that makes up the dominant contribution of $S_{1}^{21,22,61,62}$ contains tightly bound triplets that are unlikely to easily separate into free triplets⁶³ without additional energy.⁵⁰

Indeed, while intramolecular singlet fission (iSF) has been observed in a variety of long-chain polyenes in solution,²³⁻³² unlike the recently designed "class \hat{I} " iSF systems, $^{47-49,64}$ the triplet pairs in polyenes decay rapidly (ps-ns) to the S₀ ground state.³ Even in carotenoid aggregates, where intermolecular SF occurs between neighboring chromophores,^{65–70} the majority of triplet excited states decay to S₀ surprisingly quickly (within

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Figure 1. The orange carotenoid protein (OCP) photoswitches from orange (OCPo) to red (OCPr) forms with different carotenoid conformations. The OCP studied here binds a single CAN carotenoid whose skeletal structure is shown in (a). The bound CAN conformation depends on the OCP form, as shown in (b): when CAN is bound in OCPo (green, data from X-ray diffraction structure, PDB entry 4XB5^{43,44}), it has a twisted conformation; when bound in an OCPr N-terminal homologue (blue, data from X-ray diffraction structure of red carotenoid protein (RCP), PDB entry 4XB4^{43,45}), it is planar. (c) In solution, OCPo converts to OCPr under white-light illumination (1600 μ mol of photons m⁻² s⁻¹), resulting in a change in color (see the inset) and absorbance spectrum (main panel). The spectra were taken in 1 min intervals under constant white-light illumination. In the dark, OCPr converts back to OCPo (see Figure S1). The optical path length for solution measurements was 1 mm.

a nanosecond).^{3,65,70} In isolated carotenoids in solution, the dominant deactivation channel from the photoexcited S_2 state is internal conversion to S_1 . To our knowledge, there is no evidence that isolated carotenoids in solution demonstrate iSF.

Nevertheless, similarly to recent reports that torsion or twisting along a molecular backbone can allow both rapid iSF and formation of long-lived triplets in "class I" SF materials, ^{48,64} iSF along a single *twisted* carotenoid chain to produce long-lived (μ s) triplets has been suggested to occur in some photosynthetic light-harvesting complexes (LHCs).^{71–75} In these systems, the protein binds the carotenoid, so that it is constrained in a twisted geometry. This twist reportedly stabilizes a triplet at either end of the molecule.^{71,73}

This hypothesis was initially proposed to explain the presence of SF in the light-harvesting antenna (LH1) from *Rhodospirillum rubrum* because of the large intermolecular distances between neighboring carotenoids (>10 Å).⁷¹ More recently, Yu et al.⁷³ observed a correlation between the presence of SF and the so-called ν_4 resonance Raman peak (~980 cm⁻¹) in LHCs (LH1-RC and LH2) from *Thermochromatium tepidum* and *Rhodobacter sphaeroides* 2.4.1. The intensity of ν_4 is related to carotenoid backbone twisting,^{73,76} so this finding led to the conclusion that backbone twisting of the carotenoid is the "structural determinant" that enables iSF.⁷³

To test the hypothesis that SF can occur along a single twisted carotenoid chain, we examined a protein that binds a single carotenoid: the orange carotenoid protein (OCP). In OCP the protein exists in two forms, orange (OCPo) and red (OCPr), with the carotenoid in either a twisted or planar conformation, respectively (see Figure 1). By studying both forms with the protein fixed in a trehalose–sucrose glass, we demonstrate that a twisted backbone is not sufficient to enable iSF in a protein-bound carotenoid. In light of recent work understanding magnetic field effects (MFEs) in SF systems,^{77,78} we also discuss published reports of MFEs in

LHCs from purple bacteria^{79–84} and find that the reported MFEs are also inconsistent with iSF. Overall, we conclude that iSF is not supported on carotenoids bound to the OCP and is unlikely to occur in LHCs.

In this study, the OCP was produced in *Escherichia coli* by virtue of a dual plasmid system comprising pET28a with the *Synechocystis* sp. PCC 6803 OCP gene (slr1963) and pAC-CANTHipi, which provides near-100% accumulation of CAN.⁸⁵ Carotenoid-containing protein was isolated according to the method described in Supporting Information (SI) section S1.1.

In solution, upon illumination with white light, the darkadapted OCPo form undergoes a conformational switch to the OCPr form, with a concomitant red shift of its steady-state absorbance spectrum due to the effective conjugation length extension of the bound carotenoid^{76,86–88} (see Figure 1c). The change is reversible, with back-conversion from OCPr to OCPo occurring in the dark (see Figure S1).

Previously published X-ray diffraction structures by Leverenz, Sutter, and co-workers⁴³ show that the conjugated backbone of the bound carotenoid is twisted out of the plane of conjugation in OCPo (PDB entry 4XB5⁴⁴), while in OCPr N-terminal domain homologues such as red carotenoid protein (RCP) it is relatively planar (PDB entry 4XB4⁴⁵). The difference between the two conformations of CAN is depicted in Figure 1b using data from X-ray diffraction structures.⁴³ The different protein conformations containing a twisted and nontwisted form of CAN provide an uncomplicated model system to study the role of carotenoid geometry on iSF.

To avoid the problems associated with using spectroscopy to probe a light-activated conformational switch, we prevent the conformational change by trapping the protein in either its OCPo or OCPr conformation in a trehalose–sucrose glass as previously described.⁷⁰ This glass matrix prevents OCPo \rightleftharpoons OCPr conversion, as demonstrated in Figure 2a,b, and allows us to probe each conformation in isolation at room temperature without altering its conformation or photophysics.⁸⁹

To confirm the twisted/planar conformations of CAN in the OCPo/OCPr glass films, we turn to resonance Raman spectroscopy. As described above,^{73,76} the presence of a so-called ν_4 peak at ~980 cm⁻¹ in the resonance Raman spectrum of carotenoids (due to out-of-plane C–H wagging modes⁹⁰) is generally associated with a backbone twist of the carotenoid.^{73,76} Figure 2c shows the resonance Raman spectra of OCPo (blue) and OCPr (green). Consistent with previous measurements on an echinenone-binding OCP,⁷⁶ we observe a larger twist-induced ν_4 peak in OCPo than in OCPr, confirming that the native geometry is maintained in trehalose-encapsulated OCPo and OCPr.

Having established that the CAN backbone is more twisted in OCPo than in OCPr, we test the suggestion that such a twist is the determinant for iSF reactivity.^{72,73,75} Picosecond transient absorption spectra and dynamics are shown in Figures 3 and 4, respectively. Global lifetime analysis of the data is shown in Figures S5 and S6, but simply from inspection of the raw data in Figure 3 we see that all spectral features in both OCPo (green) and OCPr (blue) decay to <1% of the initial population within 30 ps. Importantly, we observe no obvious formation of SF-generated triplets. Instead, both OCPo and OCPr broadly demonstrate the expected isolated carotenoid behavior characterized by rapid internal conversion from S₂ to S₁ (as evidenced by the instrument-limited decay of



Figure 2. OCPo and OCPr trapped in trehalose glass films. The steady-state absorbance spectra of the OCPo film (a) were taken in 1 min intervals under constant white-light illumination (1600 μ mol of photons m⁻² s⁻²), and the spectra of the OCPr film (b) were taken in 1 min intervals at 22 °C in darkness. No changes in spectra were observed over 900 min. (c) The resonance Raman spectra of OCPo (green) and OCPr (blue) films in trehalose glass show vibrational peaks typical of carotenoids, labeled following convention. The spectra show a significant difference in the intensity of the ν_1 vibrational peak between OCPo and OCPr and a shift of the ν_1 peak. The Raman measurements were performed by using a 532 nm laser. Data are averages of two successive scans and normalized to the peak ν_1 intensity.



Figure 3. Transient absorption spectra of CAN-binding OCPo (a) and OCPr (b) trapped in trehalose films. The spectral time slices have been averaged between the times indicated in the figure and are consistent with an $S_2 \rightarrow S_1 \rightarrow S_0$ decay scheme in both cases, with no discernible long-lived features (see also global lifetime analysis of the data in SI section S4.2). The films were excited with 532 nm, 5 kHz, ~100 fs, and 200 μ J cm⁻² pump pulses.

an excited-state absorption (ESA) in the near-infrared region) and subsequent decay of S_1 -like states to the ground state.



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Figure 4. Transient absorption dynamics of CAN-binding OCPo (a) and OCPr (b) in trehalose glass. The dynamics have been averaged between the wavelengths indicated in the legend and demonstrate that no discernible long-lived features are seen. The films were excited with 532 nm, 5 kHz, ~100 fs, and 200 μ J cm⁻² pulses. Note that the plots have a linear time axis up to 1 ps and subsequently logarithmic up to 40 ps.

Therefore, a twist along the carotenoid backbone is not sufficient to enable iSF.

The lack of iSF in the protein-twisted carotenoid in OCPo appears to question the currently accepted hypothesis that the determinant for iSF in carotenoids is a twist along its backbone.⁷³ Indeed, while the carotenoid environments in OCP and photosynthetic complexes are very different, the carotenoids in OCPo and LHCs seem to demonstrate similar backbone twists (see Figure S3 and the SI text for further comparison).⁷³ Therefore, the lack of iSF in OCPo suggests that it would be worth revisiting the mechanism of SF in purple bacterial LHCs, particularly considering recent work on the nature of intermediate triplet-pair states involved in SF, as probed by magnetic-field-dependent measurements.^{77,78} We therefore return to the original studies of SF in these LHC systems and discuss them in light of this recent work.^{77,78}

Singlet fission in LHCs was first observed in a series of experiments that probed their magnetic-field-dependent fluorescence.^{79–81} Representative data for oxidized cells from *Rhodobacter sphaeroides* 2.4.1 from ref 79 are reproduced in Figure 5; similar behavior has been reported for whole cells and isolated LHCs from several strains of purple bacte-



Figure 5. Magnetic field effect (MFE) of *Rhodobacter sphaeroides.* Data were taken from ref 79. The data are plotted as the normalized change in fluorescence, $\Delta F/F$ (detected at 890 nm), as a function of the magnetic field strength upon 515 nm excitation in oxidized cells of *Rhodobacter sphaeroides* 2.4.1, with 1 mM K₃Fe(CN)₆ added. $A_{850-960}$ = 0.35; optical path length = 2 mm.

ria.^{79–81,84} The shape of the magnetic field effect (MFE) in Figure 5, with an initial dip in fluorescence as the field increases from 0 to 40 mT and then a rise in fluorescence to saturation beyond 100 mT, is a characteristic signature of SF.

This behavior is very well described by the kinetic model of SF by Johnson and Merrifield,^{51,91-94} published in the 1960s and 1970s. Recent work^{77,78} shows that this low-field Merrifield-type MFE behavior can only be observed when the intertriplet exchange interaction *J* is negligible or, more precisely, when $J \ll D$,⁹⁵ where *D* is the intratriplet dipolar zero-field splitting parameter. In carotenoids, and indeed most organic chromophores, *D* is relatively small, on the order of 4–10 μ eV.^{96,97} If *J* increases beyond *D*, the MFE has a different behavior, showing dips in fluorescence at much higher field strengths.^{77,78,98,99} Therefore, to determine whether SF along a single carotenoid chain is capable of producing the measured MFEs in LHCs, we must estimate the values of *J* and *D*.

Before doing so, we make several observations about the carotenoids involved in SF in LHCs of purple bacteria: (1) the $S_0 \rightarrow S_2$ absorbance spectra of the carotenoids in lightharvesting antenna are similar to those of their all-trans forms in organic solvent and depend sensitively on the carotenoid conjugation length.^{74,100} A full break in conjugation along the chain would lead to a dramatic blue shift of the carotenoid absorption feature that is not observed. (2) The carotenoid T_1 \rightarrow T_n excited-state absorption feature seen in transient absorption of LHCs^{71,72,74,75} is very similar to that seen in aggregated carotenoids of comparable conjugation lengths forming triplets by intermolecular SF.^{65–67,70} The $T_1 \rightarrow T_n$ feature is also sensitive to the carotenoid conjugation length,^{74,101,102} and a conjugation break along the chain would similarly lead to a blue shift that is not observed. (3) The dipolar *D* and *E* parameters of the SF-generated triplets in LHCs from transient electron parametric resonance (EPR) spectroscopy are similar to full-chain triplet D and Eparameters rather than to their half-chain alternatives.⁸⁴ These observations suggest that the conjugation along the chain is not broken, even in the LHC antenna protein, and therefore that the triplets at either end of the chain maintain orbital overlap and, presumably, non-negligible J.

The exchange interaction, *J*, between triplets within a pair is equal to one-sixth of the energy difference between the pure singlet triplet pair, denoted ¹(TT), and the pure quintet, ⁵(TT).³ In addition, to first approximation, the energy of ⁵(TT) is equal to twice the free triplet energy.^{3,56,103} In carotenoids, as described above, the lowest-energy singlet state (S₁) is predominantly a pure singlet ¹(TT) state. Therefore, a comparison between twice the energy of a triplet on half a chain against the energy of S₁ on a full chain provides an indication of the exchange interaction.

Recent high-level density matrix renormalization group (DMRG) calculations of the Pariser–Parr–Pople–Peierls Hamiltonian⁵⁶ show that $2 \times E(T_1)$ for a half chain is higher in energy than $S_1 (\approx^1(TT))$ for a full chain at all conjugation lengths. This is supported by experimentally determined energies: for diphenylexatriene with N = 5 conjugated double bonds, $2 \times E(T_1) = 3.02 \pm 0.1 \text{ eV}$,^{101,104} while for spheroidene with twice the number of double bonds (N = 10), $E(S_1) = 1.77$ eV.¹⁰⁵ This would indicate an exchange interaction of J = 0.2 eV, which is orders of magnitude larger than the dipolar parameter $D \sim 4-10 \ \mu \text{eV}$.^{96,97} These energies indicate that the triplets within ¹(TT) should be strongly exchange-coupled.

The triplets within a single carotenoid chain are therefore exchange-coupled $(J \gg D)$ even in a protein that twists the carotenoid backbone,⁷³ as no breaks in conjugation along the carotenoid chain have been observed (i.e., no observable shifts in absorption spectra⁷⁴ or changes in dipolar D and E parameters⁸⁴). Therefore, MFEs such as those reproduced in Figure 5, that were the initial proof of SF in purple bacteria, cannot be explained with an intramolecular model of SF. We speculate that the mechanism for singlet fission in light-harvesting complexes must be intermolecular, possibly occurring via the neighboring bacteriochlorophyll a molecules in the protein complex.

We conclude that singlet fission (SF) to produce long-lived triplets does not occur along a single twisted carotenoid chain in the OCP and is unlikely to occur on a single carotenoid chain in purple bacterial light-harvesting complexes (LHCs), contrary to the current notion.^{71–75} We conclude this because (1) immobilized OCPo—an uncomplicated, minimal carotenoprotein—shows similar twisted carotenoid geometry to LHCs but shows no evidence of SF and (2) the MFEs that identified SF in purple bacteria are irreconcilable with iSF without a significant break in conjugation (which is not observed in LHCs). These findings therefore call into question the mechanism of SF observed in LHCs.

ASSOCIATED CONTENT

Data Availability Statement

Data are available on the University of Sheffield's repository, ORDA.

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.jpclett.3c01139.

Experimental methods, including canthaxanthin-binding OCP production, OCPo and OCPr in sugar glasses preparation, spectroscopic setups, and data analysis procedures; back-conversion of solution OCPr to OCPo in the dark; comparison of the resonance Raman spectra of OCPo and OCPr of this work against previously published spectra of purple bacterial LHCs; resonance Raman spectrum of blank trehalose; transient absorption pump spectrum; results of global lifetime analysis on the transient absorption data (PDF)

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Author Contributions

¹G.A.S. and J.P.P. contributed equally. G.A.S. conceived the study. G.A.S., J.P.P., S.W. and J.C. designed the experiments. G.A.S., M.S.P. and A.H. prepared protein samples under the supervision of M.P.J. and C.N.H. Absorbance measurements were performed by G.A.S. Resonance Raman measurements were performed by H.K.H.L. under the supervision of W.C.T. Transient absorption was conducted by J.P.P. and S.W. within the Lord Porter Laser Facility, with D.C. providing facility management. J.P.P. and G.A.S. analyzed the data. J.P.P., G.A.S. and J.C. wrote the manuscript and prepared the figures with input from all authors.

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Notes

The authors declare no competing financial interest.

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