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Singlet Fission in Naturally-Organized Carotenoid Molecules

Annamaria Quaranta[†], Anja Krieger-Liszak[†], Andrew A. Pascal[†], François Perreau[‡], Bruno Robert[†], Mikas Vengris^{§*} and Manuel J. Llansola-Portoles^{†*}

[†] Université Paris-Saclay, CEA, CNRS, Institute for Integrative Biology of the Cell (I2BC), 91198, Gif-sur-Yvette, France

[‡] Institut Jean-Pierre Bourgin, INRAE, AgroParisTech, Université Paris-Saclay, 78000, Versailles, France

[§] Laser Research Center, Faculty of Physics, Vilnius University, Saulėtekio Ave. 10, LT-10223, Vilnius, Lithuania

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ABSTRACT: We have investigated the photophysics of aggregated lutein/violaxanthin in daffodil chromoplasts. We reveal the presence of three carotenoid aggregate species, the main one composed of a mixture of lutein/violaxanthin absorbing at 481 nm, and two secondary populations of aggregated carotenoids absorbing circa 500 and 402 nm. The major population exhibits an efficient singlet fission process, generating μ s-lived triplet states on an ultrafast timescale. The structural organization of aggregated lutein/violaxanthin in daffodil chromoplasts produces well-defined electronic levels that permit the energetic pathways to be disentangled unequivocally, allowing us to propose a consistent mechanism for singlet fission in carotenoid aggregates. Transient absorption measurements on this system reveal for the first time an entangled triplet signature for carotenoid aggregates, and its evolution into dissociated triplet states. A clear picture of the carotenoid singlet fission pathway is obtained, which is usually blurred due to the intrinsic disorder of carotenoid aggregates.

INTRODUCTION

The bright colours of many flowers and fruits originate from the carotenoids accumulated in specialized plastids in the plant cell, termed chromoplasts. Chromoplasts are generally produced in flower petals and ripe fruit through a remodeling of the photosynthetic chloroplast¹, involving chlorophyll degradation, down-regulation of genes encoding for photosynthetic proteins, and up-regulation of stress-response and carotenoid-synthesis genes (although differentiation may also occur from other plastid types²). In mature chromoplasts, the carotenoid pigments are sequestered in the form of crystal-like structures (crystalloids), granules, or stromal plastoglobules³. The main function of the chromoplast is thought to be secondary signaling, attracting animals to flowers and fruits to aid pollination and seed dispersal, respectively. They are also the site of protein, lipid and carbohydrate biosynthesis⁴, and play an important role in fruit and flower scent (for instance, one of the main components of the tomato scent is a degradation product of chromoplastic lycopene)⁵. It is interesting to note that the chloroplast/chromoplast transition is in some cases a reversible process, such that chromoplasts can re-acquire photosynthetic activity and differentiate back into chloroplasts^{1, 2}.

Carotenoid molecules are tetraterpenoid derivatives formed by eight isoprene units. They display a wide variety

of functions in photosynthetic tissues, some of which are also observed throughout nature. They play an important role in harvesting photons in the blue-green range in photosynthesis⁶, and ensure efficient photoprotection of the photosynthetic apparatus - through the quenching of chlorophyll triplet⁷ and singlet⁸ excited states, and by short-circuiting lipid peroxidation⁹. Carotenoids exhibit strong electronic absorption corresponding to a transition from the ground state to the second excited state ($S_0 \rightarrow S_2$), while transition to the first excited state (which has the same symmetry as the ground state) is one-photon forbidden¹⁰. After photoexcitation, isolated carotenoids decay rapidly to the ground state in a few picoseconds^{11, 12}. Although chromoplasts are not expected to display any specific functional photoactivity, we recently discovered that the lycopene crystalloids in tomato chromoplasts do indeed exhibit photophysics that are not observed in isolated carotenoids¹³. This self-associated lycopene dissipates excitation energy via singlet fission, a mechanism we tentatively proposed to be photoprotective. In the singlet exciton fission process, one singlet excited state splits into two excited triplets ($T_1 + T_1$), fulfilling the energetic condition $E(S_1) \geq 2 \bullet E(T_1)$ ¹⁴⁻¹⁶. Carotenoids in tightly-interacting assemblies *in vitro* have been shown to generate triplets by singlet fission - such as aggregated astaxanthin^{17, 18}, β -carotene^{19, 20}, and

zeaxanthin²¹⁻²³ - but up to now, tomato chromoplasts represent the only known example of this process in a biological structure¹³.

We investigated the lutein/violaxanthin photochemistry in daffodil (<*Narcissus pseudonarcissus* L.> chromoplasts^{24, 25}, to determine whether singlet fission is unique to the lycopene in tomato crystalloids, or is a more general chromoplast phenomenon. Daffodil chromoplasts accumulate these carotenoids in a concentric membrane system, forming condensed structures involving carotenoids and acyl lipid molecules (they belong to the class of so-called membranous chromoplasts)^{24, 25}. These structures allow close interactions between carotenoids, which perturb the absorption properties of lutein and violaxanthin. We show that singlet fission indeed occurs in these plastids, suggesting that this process may be common to all carotenoid-containing chromoplasts. Our experiments reveal the singlet fission mechanism in unprecedented detail. The information on specific energy levels of intermediate species allows us to identify the entangled triplet pair, and to pinpoint the spectral signatures of the disentangled triplet excited state(s) of the excitonic lutein/violaxanthin system.

MATERIALS AND METHODS

Chromoplasts of the inner coronae of fully-opened daffodil flowers (*Narcissus pseudonarcissus* L. cv Dutch Master), at developmental stage IV²⁶, were isolated according to the method of Liedvogel *et. al.*²⁵ using a step gradient in which chromoplasts accumulate at the 15 % / 30 % sucrose interphase²⁷. The isolated chromoplasts were suspended in 100 mM Tris/HCl pH 7.4, 10 mM MgCl₂, 1 mM 1,4-dithioerythritol. The preparations were stored at -80 °C.

For pigment determination, the preparation was centrifuged, the supernatant discarded and the pelleted chromoplasts were resuspended in 2 mL acetone. 5 µL was then subjected to liquid chromatography at a flow rate of 0.5 mL min⁻¹, using an HPLC column (Uptisphere Strategy C18-HQ 250 × 3 mm 3 µm, Interchim, Montluçon, France) with a liquid chromatography chain HPLC/UV (Shimadzu, Tokyo, Japan) which included two pumps (LC-20AD), a sample manager (SIL-20AC HT), a column oven (CTO-20A) and a UV diode-array detector (UVSPD-M20A) monitoring the absorbance at 450 nm. Mobile phase A was composed of 10 % water and 90 % acetonitrile with 0.5 % acetic acid, mobile phase B of ethyl acetate containing 0.5 % acetic acid, and the elution gradient was as follows: 1 min 90 % A : 10 % B, then a linear gradient to 5 % A : 95 % B after 25 min. Concentrations of each pigment were calculated based on their peak areas, calibrated using injected standards or (for some minor pigments) with known absorption factors for each compound at 450 nm.

Absorption spectra were measured using a Varian Cary E5 scanning spectrophotometer. Low temperature measurements were performed in a helium bath cryostat (Maico Metriks, Tartu, Estonia); 60 % glycerol (v/v) was added to the sample prior to freezing.

Circular dichroism spectra were recorded on an Aviv 215 spectropolarimeter (Aviv Biomedical). Near-UV CD spectra were recorded between 300 and 600 nm using a square

cell with a 1 cm path length, and an averaging time of 1 s per step. Three consecutive scans from each sample were merged to produce an averaged spectrum, which was then corrected against buffer baselines measured under the same conditions. Dichroism value is measured in mdeg.

Resonance Raman spectra were recorded at 77 K with laser excitations obtained from Coherent Ar⁺ (Sabre) and Kr⁺ (Innova 90) lasers. Output laser powers of 10–100 mW were attenuated to < 5 mW at the sample. Scattered light was collected at 90° to the incident light, and focused into a Jobin-Yvon U1000 double-grating spectrometer (1800 grooves/mm) equipped with a red-sensitive, back-illuminated, LN₂-cooled CCD camera. The sample stability and integrity were assessed based on the stability of the Raman signal.

Nano-to-millisecond transient absorption experiments were performed on an Edinburgh Instruments LP920 Flash Photolysis Spectrometer system, incorporating a Continuum Surelite OPO. The OPO was pumped by a Q-switched Nd:YAG laser operating at 355 nm. The LP920 system is equipped with a 450 W Xenon arc lamp as the probe for the transient absorption measurements; this lamp was pulsed in the time range 10 ns to 100 µs. Detection in the LP920 system was performed either via a 500-nm-blazed Czerny-Turner monochromator (bandwidth 1-5 nm) coupled with a Hamamatsu R928 photomultiplier tube (kinetics mode), or via a 500-nm-blazed spectrograph (bandwidth 5 nm) coupled with a water-cooled ICCD nanosecond Andor DH720 camera (spectral mode). Excitation energy was approximately 15 mJ / pulse.

Femto-to-nanosecond time-resolved transient absorption spectra were recorded on a setup described in detail elsewhere²⁸. Pump and probe pulses were derived from an amplified Ti:Sapphire laser (Libra, Coherent). Tunable pump pulses with center wavelength at 490 nm were obtained from an optical parametric amplifier (Topas, Light Conversion). White light continuum generated in CaF₂ from the fundamental output of the laser was used to probe the absorption changes in the 350-750 nm range. The time resolution of the instrument is ca. 120 fs. Excitation energies were set to approximately 200 nJ / pulse.

RESULTS AND DISCUSSION

The carotenoid content of daffodil chromoplasts was determined by HPLC (Supporting information, Figure S1). The carotenoids with significant concentrations are lutein and violaxanthin, with a ratio circa 5:4. Lutein and violaxanthin in cyclohexane both present the typical absorption spectrum for carotenoids, with four bands corresponding to the vibronic sublevels 0-3, 0-2, 0-1, and 0-0 of the S₀→S₂ electronic transition - at 395, 416, 440 & 474 nm and 393, 422, 452 & 471 nm, respectively (Fig. 1a)²⁹. Figure 1b shows the absorption spectrum of broken daffodil chromoplasts in 99 % ethanol, which disrupts the chromoplast and solubilizes the carotenoids. This spectrum, with maxima at 394, 418, 442 & 471 nm, is essentially identical to a linear combination of isolated lutein and violaxanthin using the proportion obtained from the HPLC analysis (supporting information, Figure S2). The absorption spectrum of intact

chromoplasts at room temperature exhibits broad vibronic transitions peaking at 393, 422, 447 and 481 nm (Fig. 1c). The o-o band at 481 nm is redshifted relative to the spectrum of solubilized chromoplasts (note that the absorption of lutein may be observed as high as 485 nm in high polarizability solvents³⁰). The intensity of the o-3 and o-2 electronic transitions is significantly higher in chromoplasts than in monomeric carotenoids (cf. Fig. 1a,b,c), which may indicate some perturbation of the electronic transition caused by packing interactions³¹. The absorption spectrum of daffodil chromoplasts recorded at cryogenic temperature (4.2 K) reveals a complex picture, indicating the presence of several carotenoid populations (Fig. 1d) which can be resolved by second derivative analysis (Fig. 1e). The red part of the absorption spectrum shows the presence of two species with close electronic transitions: a dominant species (species 1) absorbing at 422, 451 & 487 nm, and a minor one (species 2) with absorption at 430, 463 & 500 nm (o-2, o-1 & o-o, respectively). An additional band is present at 402 nm whose second derivative shows no clear structure. This suggests that this 402-nm band is not a o-3 vibronic transition of species 1 & 2, but could rather arise from a third, blue-shifted carotenoid population (species 3). Overall, the absorption at 4.2 K presents a characteristic bathochromic shift relative to the spectrum at room temperature^{32, 33} (note red-shift of the major o-o band from 481 to 487 nm). The circular dichroism of daffodil chromoplasts

at room temperature (Fig 1f) exhibits chirality across the whole spectral range, with a dominant S-shaped feature on the blue side of the envelope (maximum 397 nm, minimum 375 nm). The CD spectrum of solubilized chromoplasts displays no chirality (data not shown). The observed chirality for intact chromoplasts indicates that all three species are constituted from aggregated (interacting) carotenoid molecules. Species 3 has a blue shifted o-o transition, around 402 nm at 4.2 K, and displays a large CD signature. This is representative of H-aggregates, where exciton coupling is strong and extends over multiple chromophores. The absorption of such strongly-coupled H-aggregates collapses into a single electronic transition, with the large hypsochromic shift observed here (circa 5000 cm^{-1} relative to monomeric violaxanthin and lutein)³⁴. Species 1 & 2 conserve the vibronic peaks of monomeric carotenoids, and exhibit only a small bathochromic shift relative to them. Carotenoid aggregates that have a predominant bathochromic shift have been traditionally interpreted as J-aggregates^{35, 36}. However, species 1 & 2 do not resemble the typical narrow exciton-coupled bands expected for strongly-coupled J-aggregates^{17, 18, 37}. Their absorption and CD spectra more closely resemble model spectra of weakly-coupled H-aggregates of lutein³¹. Further studies are necessary to determine the precise molecular interaction(s) involved in species 1 & 2.

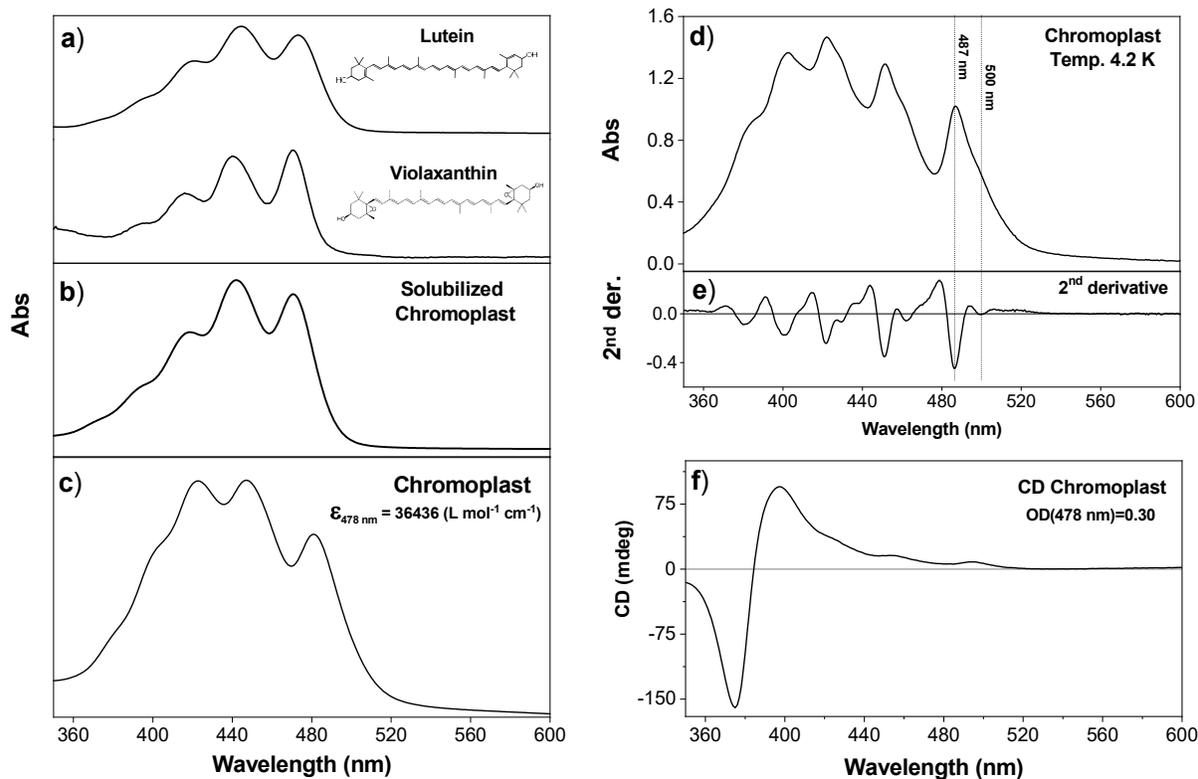


Figure 1: Room temperature absorption spectra of (a) monomeric lutein and violaxanthin in n-hexane; (b) solubilized chromoplasts; and (c) intact daffodil chromoplasts in buffer solution. (d) 4.2 K absorption spectrum of daffodil chromoplasts, and (e) its second derivative. (f) CD spectrum of daffodil chromoplasts in buffer solution, obtained with 0.3 OD at 478 nm.

We used resonance Raman spectroscopy³⁸ at three different excitation wavelengths to characterize the three carotenoid species in daffodil chromoplasts. The high selectivity of this technique provides information for pigment populations individually, by exciting close to their respective absorption maxima. Figure 2 shows the resonance Raman spectra at 77 K for isolated *all-trans* lutein and violaxanthin in cyclohexane, measured at 488.0 nm, and for intact chromoplasts in buffer excited at 488.0, 501.7 and 406.7 nm. The position of the ν_1 vibrational mode depends on the length of the π -electron conjugated chain in isolated carotenoids^{39, 39}. This band occurs at 1530.2 and 1529.1 cm^{-1} for violaxanthin and lutein in cyclohexane, respectively. The ν_1 frequency in daffodil chromoplasts exhibits a clear dependence on the excitation wavelength, peaking at 1530.4, 1526.7 & 1528.0 cm^{-1} for excitation at 488.0, 501.7 & 406.7 nm, respectively (see figure 2d). These different frequencies support the presence of three non-identical carotenoid species – note that the three excitation wavelengths correspond to the absorption peaks of species 1-3 (487, 500 & 402 nm, respectively). Due to the excitonic nature of aggregates, it is not possible to relate the ν_1 frequency to that of monomeric violaxanthin or lutein. However, the position of the ν_1 mode permits us to confirm the presence of more than one aggregate species with very close absorption.

Around 1160 cm^{-1} , a group of bands termed ν_2 constitutes a fingerprint for the isomerization state (*cis/trans*) of the carotenoid carbon backbone⁴⁰⁻⁴². Chromoplasts display the same modes at all wavelengths used, with a dominant band at 1058 cm^{-1} and two satellites circa 1188 & 1214 cm^{-1} . The absence of a vibrational mode at 1130 cm^{-1} in daffodil chromoplasts, as for the isolated standards, confirms that both lutein and violaxanthin are in the *all-trans* configuration⁴⁰⁻⁴². Observed around 1000 cm^{-1} , the ν_3 band arises from in-plane rocking vibrations of the methyl groups attached to the conjugated chain, coupled with in-plane bending modes of the adjacent C-H's⁴³. The ν_3 mode is a fingerprint of the conformation of carotenoid conjugated end-cycles⁴⁴. This band can thus be used to track the presence of lutein and/or violaxanthin in the chromoplast spectra, as a result of differences in their end-rings. Lutein in cyclohexane shows a sharp asymmetric band peaking at 1003.1 cm^{-1} with a small shoulder ca. 1007.9 cm^{-1} , while violaxanthin presents a broad band peaking at 1006.5 cm^{-1} with a clear shoulder ca. 1002.9 cm^{-1} . Chromoplasts excited at 501.7 nm exhibit a sharp vibrational mode at 1004.4 cm^{-1} , resembling the spectrum of monomeric lutein, thus indicating that this carotenoid is dominant in species 2. At 488.0 & 406.7 nm, the ν_3 band is wider with a prominent shoulder extending up to 1007.2 cm^{-1} - indicating the presence of both violaxanthin and lutein in species 1 & 3 (see figure 2c). The ν_4 mode around 960 cm^{-1} arises from C-H out-of-plane wagging motions coupled with C=C torsional modes,³⁹ and this region is an indicator of out-of-the-plane distortions of the carotenoid conjugated chain^{39, 45}. In chromoplasts, this region contains small but clear modes at 951.8 and 963.8 cm^{-1} , which are also present for isolated lutein and

violaxanthin. Thus, all three species appear to experience no significant distortions from their relaxed conformation.

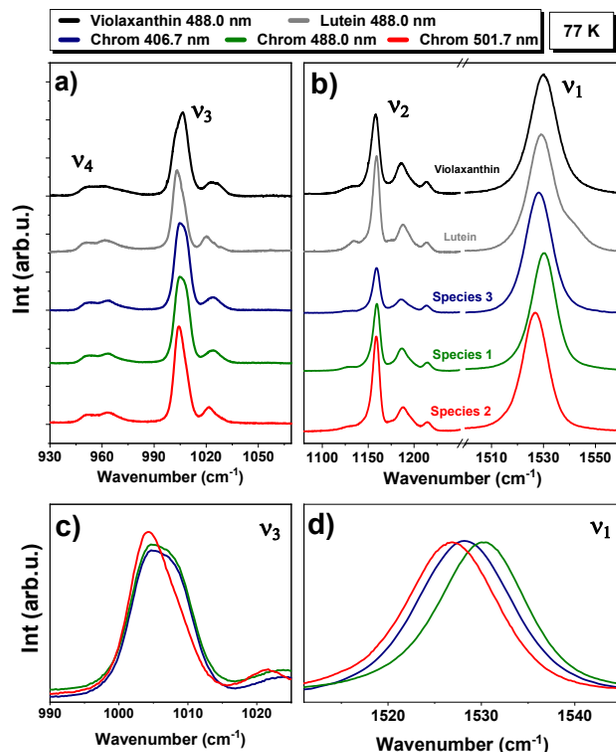


Figure 2: 77 K resonance Raman spectra of violaxanthin (black) and lutein (grey) in cyclohexane, excited at 488.0 nm, and of intact chromoplasts excited at 406.7, 488.0 & 501.7 nm (blue, green, red respectively), showing the 930-1050 and 1100-1600 cm^{-1} regions (a, b respectively). The ν_3 and ν_1 regions are expanded in panels c and d, respectively (normalized to peak height).

Figure 3 shows the time-resolved absorption data obtained for chromoplasts in the 100 fs - 5 ns time window, measured with an excitation pulse centered at 488 nm to investigate the photochemistry of the major population of carotenoid aggregates (species 1). The dataset overview (Figure 3a) shows that different parts of the ground-state bleaching (GSB) region in the 350-475 nm range decay with markedly different kinetics, and the excited state absorption (ESA) band in the 500-600 nm region exhibits two prominent peaks that reach their maxima at different times. Both these trends can be observed in more detail in the kinetic traces, presented in Figure 3b. The two ESA bands evolve at different rates, with the 552-nm peak appearing and then decaying slightly before the one at 528 nm. The complex evolution of these ESA bands is clearly observed in time-gated spectra (Figure 3c). The initial S_2 excited state, observed by its negative stimulated emission (SE) band visible above 505 nm in the 120-fs spectrum (black trace), relaxes very quickly - evolving into a broad, almost featureless band peaking around 560 nm (red trace, 300 fs). This broad feature can be attributed to a vibrationally-hot S_1 state. The evolution of this state is different from that observed for isolated carotenoids - instead of

narrowing to a well-defined S₁ ESA peak, a nearly-symmetrical doublet appears with maxima at 528 and 552 nm (green trace, 2 ps). These peaks decay at different rates, so that the 528-nm maximum dominates the 15-ps spectrum (blue trace). At later times the ESA band shifts further to the blue, where it partly decays within 100 ps (cyan trace).

However, a small fraction of this signal remains beyond the 5-ns time window of the experiment (eg. pink trace, 2-5 ns). The evolution at longer timescales is again different from that seen for monomeric carotenoids in solution – including violaxanthin and lutein – which return to the ground state in tens of ps¹².

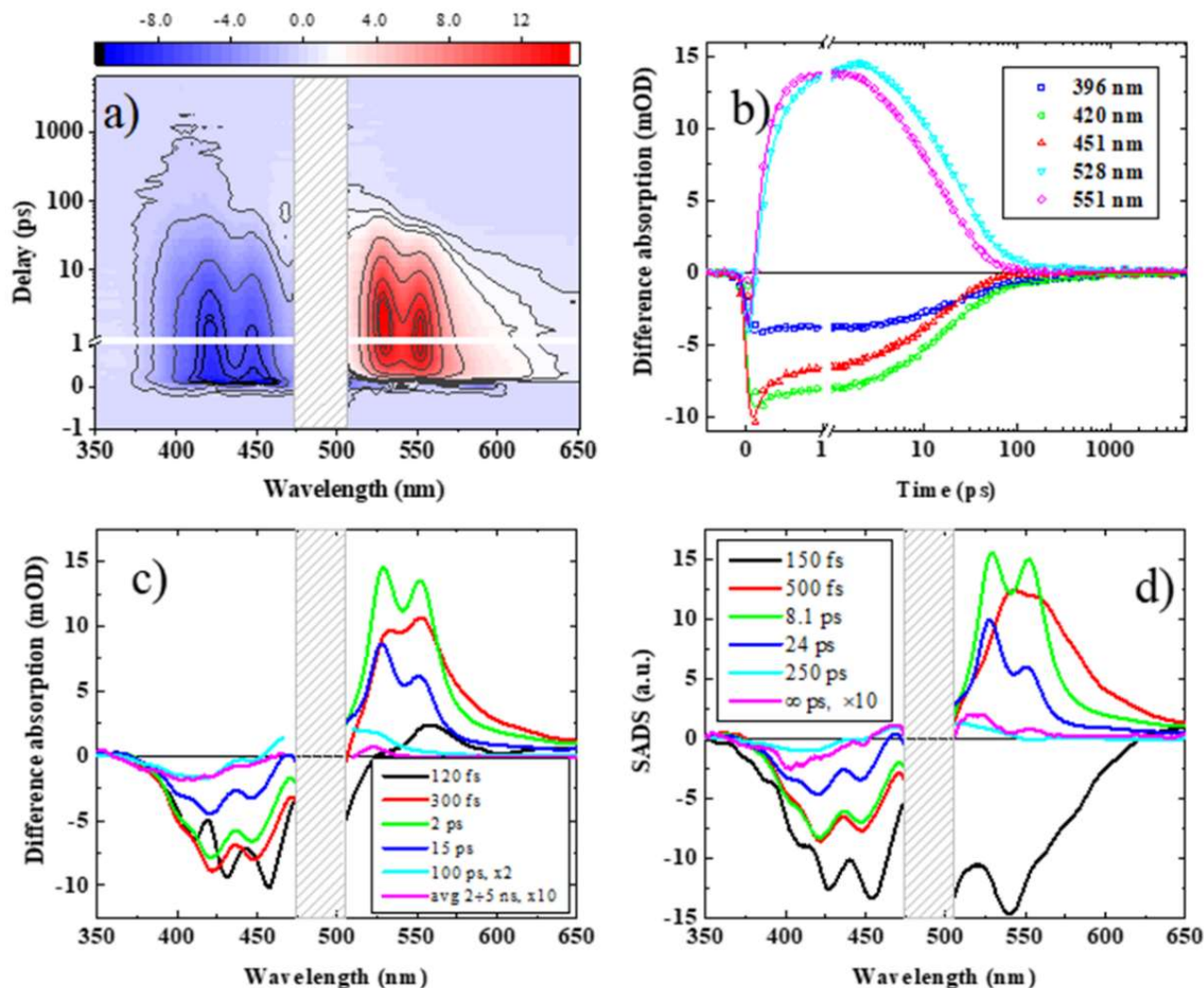


Figure 3: Femtosecond-to-nanosecond transient absorption data of daffodil chromoplasts obtained with excitation at 488 nm: (a) dataset overview; (b) selected absorption-difference kinetic traces (symbols - experimental data; solid lines - results of global fit, as per d); (c) time-gated spectra at selected delay times, as indicated; (d) EADS estimated by global fitting, using 6-component evolutionary model (decay times indicated in the legend). Greyed regions indicate spectral range corrupted by scattering of the pump pulses.

Global analysis using an evolutionary model yields the Evolution-Associated Difference Spectra (EADS) shown in Figure 3d, reflecting a cascade of events with the following time constants: 150 fs → 500 fs → 8.1 ps → 24 ps → 250 ps → non-decaying. These first two EADS at 150 and 500 fs (black, red spectra) are similar to those observed for monomeric carotenoids, reflecting analogous evolution over the same timescales¹². The first step corresponds to the decay of the S₂ state (150 fs, black spectrum), somewhat obscured by cross-phase modulation between pump and

probe. The second step corresponds to cooling of the resulting hot S₁ state (500 fs, red spectrum), whereby ESA loses its red shoulder at 550-650 nm. The third EADS (8.1 ps, green spectrum) presents two peaks of similar intensity at 528 and 552 nm, while the GSB in the blue part of the spectrum remains unchanged relative to the 500-fs spectrum. In the fourth EADS (24 ps, blue spectrum) a substantial decrease of relative intensity of the 552-nm peak is observed. The fifth EADS (250 ps, cyan spectrum) presents a distorted shape of the GSB on its red wing, being attenuated by about 90 %, possibly indicating the presence of a

transient species with ESA overlapping in this region and pushing the signal up. The peaks at 528 and 552 nm disappear completely, while a broad ESA band peaks at ca. 507 nm. This sixth EADS (∞ , pink spectrum) is similar to that at 250 ps and persists beyond 5 ns (time window of the experiment). It should be noted that the two excited species with ESA peaking at 528 and 552 nm decay in parallel, and they cannot be separated in this sequential model. This model is therefore unable to describe the dynamics after photoexcitation completely, while it is adequate to perform an initial assessment of the excited species that will lead to the design of a representative model (see target analysis, below and Fig 5). In monomeric carotenoids in solution, S_1 decays to the ground state in a few picoseconds, with no long-living species. The ESA position and decay rate for S_1 in monomeric violaxanthin and lutein are 532 nm & 24 ps, and 558 nm & 14 ps, respectively¹². In addition, the GSB intensity for isolated carotenoids decays in proportion to the S_1 decay. In daffodil chromoplasts, the unchanged intensity of GSB between 500 fs and 8.1 ps indicates that the recombination of excited states is somewhat compensated by the formation of additional excited species depleting the ground state. It suggests that the two ESA cannot be assigned to independent S_1 states decaying to the ground state. In the literature, similar ESA doublets have been reported for aggregates of β -carotene²⁰ and zeaxanthin^{22, 23}. Hence, the red ESA may be assigned to S_1 (generated by excitonic coupling) decaying in a few ps, while the blue ESA can be assigned to a triplet state decaying over a wide range of timescales, from 10s to 100s of ps. Hence, the ESA observed for daffodil chromoplasts at 528 nm may indicate the presence of triplet states.

In order to analyze the fate of the excitation at longer timescales, transient absorption spectra of daffodil chromoplasts in the ns-to- μ s range were measured in buffer solutions saturated with oxygen, air and argon (Fig. 4). All transient absorption kinetics can be fitted to a mono-exponential decay, and the gated spectra for chromoplasts at three different oxygen concentrations are nearly identical. They all show a wide negative region around 350-440 nm, attributable to GSB. In the 440-500 nm region, the ground state bleaching is mixed up with ESA, with negative contributions at 480 and 444 nm corresponding to the o-0 and o-1 vibronic peaks of the $S_0 \rightarrow S_2$ electronic transitions, respectively. The ESA in the region 500-530 nm exhibits a peak at 505 nm, accompanied by a signal at 520 nm appearing as a peak in argon or a shoulder in air and oxygen. These signals decay with lifetimes of 180 ns, 700 ns, and 3.5 μ s in oxygen, air and argon, respectively. The dependence of this lifetime on oxygen concentration (Fig. 4), a molecule known to enhance inter-system crossing in carotenoid triplet states^{46, 47}, is consistent with the attribution of these ESA to $T_1 \rightarrow T_n$ transitions. In monomers of the plant antenna protein LHCI, no significant difference is observed between lutein and violaxanthin triplets, in the native protein and the *npq1utz* mutant, respectively - $T_1 \rightarrow T_n$ peaks at 508 nm in both cases⁴⁸. The weak excitonic coupling of species 1 and 2 may introduce changes in their $T_1 \rightarrow T_n$ ESA position. It therefore seems reasonable to assign the

higher-energy $T_1 \rightarrow T_n$ transition at 505 nm to associated lutein/violaxanthin triplets in species 1 (o-o at 481 nm a room temperature), while associated lutein (species 2; 500 nm) contributes to the lower-energy triplet signal at 520 nm.

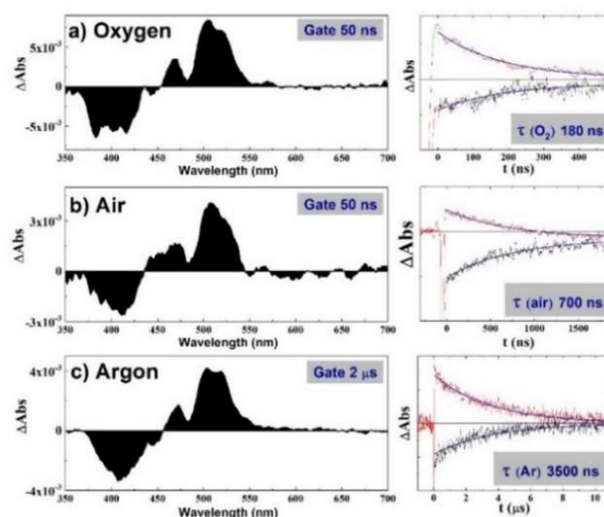


Figure 4: Transient absorption in the ns-to- μ s window for daffodil chromoplasts upon 488-nm excitation: gated spectra and time evolution for 405 and 500 nm in (a) oxygen, (b) air, and (c) argon.

The only known process capable of forming triplet states in carotenoids in the absence of a photo-sensitizer is singlet exciton fission. This process has been observed in several cases where intermolecular interactions occur between carotenoids^{13, 17-23}. In acenes, the mechanism of triplet formation is fairly well understood. A singlet-excited state (S_1) on one pigment shares its energy with a neighboring pigment in the ground state (S_0), resulting in a correlated pair of triplets ($^1(TT)$) on these two molecules, entangled with spin coherence and overall zero spin. This strongly-entangled state has a singlet character, so that its decay to the singlet ground state is allowed with rates reminiscent of singlet excited states (ps-ns)^{14, 15}. Conversely, the intermediates involved in the evolution of this $^1(TT)$ state remain controversial. A weakly-entangled triplet ($T_1 \dots T_1$) is necessary as an intermediate step, which may either yield two independent triplet states ($T_1 + T_1$), or recombine by triplet-triplet annihilation. The spectral position of the $^1(TT)$ signature is not well established - for acenes it appears blue-shifted with respect to an independent triplet, while it has been proposed as red-shifted for terrylene-diimide dimers⁴⁹ and β -carotene aggregates^{14, 15}.

In carotenoids, photoexcitation leads to the formation of the second excited state (S_2), with an energy of the lowest vibrational level at 2.57 eV. The (S_1) singlet state consists of two coupled triplets residing on the same isolated molecule¹⁰. Consequently, the mechanism of singlet fission in daffodil chromoplasts must take into account the contribution of the second excited state, while also allowing for the excitonic character of S_2 , S_1 and T_1 . Excitation energy is transferred from S_2 to the higher vibrational levels of the first excited state (hot S_1) in 150 femtoseconds, whereupon two pathways compete in ca. 500 fs: i) decay to the lowest

vibrational states of S_1 , and ii) formation of an entangled triplet state $^1(TT)$ between two neighboring carotenoid molecules, with both triplets residing on the same excitonic manifold. The relaxed S_1 state decays in 11 ps to the ground state (S_0), whereas the entangled triplets $^1(TT)$ either decay in 36 ps to the ground state (transition allowed due to singlet character), or dissociate into weakly-entangled and then independent triplets ($T_1\dots T_1$ & $T_1 + T_1$, respectively). Target analysis based on this model, presented in Figure 5a, produces a remarkably good fit to the femtosecond data (see solid lines in Fig. 5b), and the resulting spectra exhibit the exact features expected from the model. This is clear from the hot S_1 , the relaxed S_1 and the $^1(TT)$ states, which exhibit very well-separated ESA. Hot S_1 has a broad and featureless induced absorption with a pronounced red shoulder, the relaxed S_1 state exhibits a narrow and well-defined peak at 553 nm, and the $^1(TT)$ state peaks at 528 nm. This $^1(TT)$ decays following two competitive channels, with a 36-ps decay to the ground state, or towards ($T_1\dots T_1$) in 240 ps. The weakly-entangled triplets ($T_1\dots T_1$) and independent triplets (T_1+T_1) are modeled as two exponentially-decaying states with identical spectra. They show a broad induced absorption band peaking around 510 nm, with no ESA above 550 nm at long time-scales.

This set of experiments additionally allows a fairly precise estimation of the energy of the T_1 state of lutein/violaxanthin aggregates, a parameter which has never been measured directly. The energy of the S_2 state is 2.57 eV, as obtained from the absorption spectrum, and that of the relaxed S_1 state has been estimated at 1.55 eV⁵⁰. The energy level of $^1(TT)$ can be assumed to be equal to S_1 within a difference depending on the entanglement energy (0.15 eV)^{14, 15}, while the independent triplets must obey the energetic condition $E(S_1) \geq 2 \bullet E(T_1)$. Hence the $^1(TT)$ energy is 1.55 ± 0.20 eV, while the T_1 energy level for aggregated carotenoids in daffodils is 0.78 ± 0.10 eV (Scheme 1).

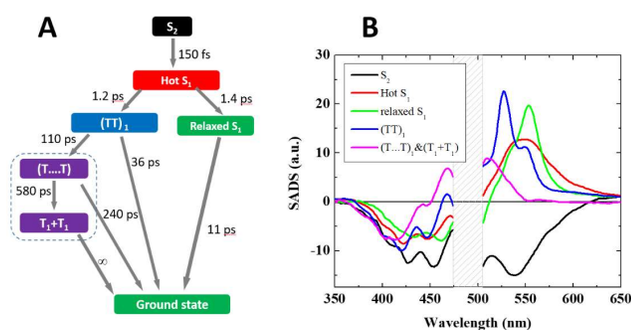
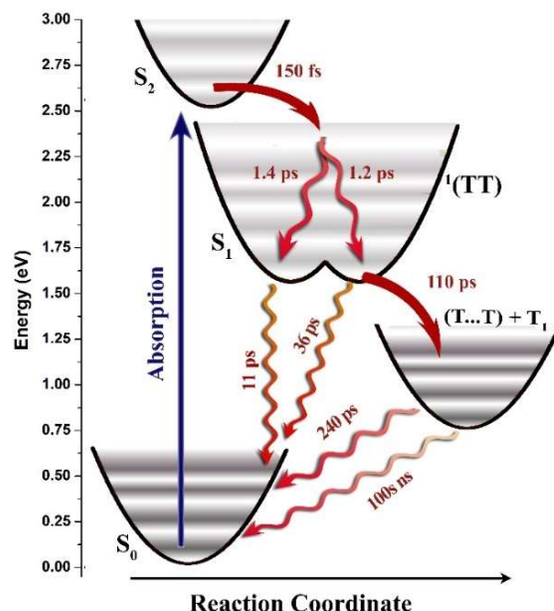


Figure 5: Target model (a) and species-associated difference spectra (b) resulting from target analysis of femtosecond pump-probe data on daffodil chromoplasts. The estimated time constants are shown next to the corresponding arrows in panel a.



Scheme 1: Energetic pathways in aggregated lutein & violaxanthin after absorption of one photon (lifetimes as obtained from target analysis).

CONCLUSIONS

In daffodil chromoplasts, the interactions of lutein and violaxanthin lead to the existence of three different species of carotenoids aggregates, the most abundant being a mixture of lutein/violaxanthin with o-o transition at 481 nm. The photodynamics of this species induces the emergence of μ s-lived triplet states produced via singlet fission, as already observed for self-associated lycopene in ripe tomato fruit. This first report of singlet fission occurring in mature flower plastids suggests that triplet fission is a phenomenon general to carotenoid aggregates in chromoplasts. While in tomatoes, lycopene interactions induce both a large redshift and strong distortions of the lycopene molecule¹³, in daffodil chromoplasts singlet fission is observed from lutein/violaxanthin aggregates which exhibit only a moderate red-shift in absorption transitions, and no molecular distortion of the interacting carotenoids. The fact that we observe singlet fission in both fruits and, for the first time here, flowers, occurring in carotenoid assemblies displaying different organizations and levels of interaction (strong and medium-to-weak, respectively), suggests a specific role of singlet fission in plastids. Finally, the structural organization of lutein/violaxanthin in daffodil chromoplasts produces well-defined electronic levels, allowing the energetic pathways following carotenoid singlet fission to be disentangled unequivocally for the first time.

ASSOCIATED CONTENT

Supporting Information. HPLC determination of carotenoid content in daffodil chromoplasts (Fig. S1); linear combination of *in vitro* absorption spectra of isolated carotenoids, to reproduce the spectrum of solubilized chromoplasts (Fig. S2). This material is available free of charge via the Internet at <http://pubs.acs.org>.

AUTHOR INFORMATION

Corresponding Author

* Mikas Vengris, Laser Research Center, Faculty of Physics, Vilnius University, Saulėtekio Ave. 10, LT-10223, Vilnius, Lithuania.

e-mail: mikas.vengris@ff.vu.lt

* Manuel J Llansola-Portoles, Université Paris-Saclay, CEA, CNRS, Institute for Integrative Biology of the Cell (I2BC), 91198, Gif-sur-Yvette, France.

e-mail: manuel.llansola@izbc.paris-saclay.fr

Author Contributions

The manuscript was written through contributions of all authors.

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