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Introduction

Photophysical properties of betaxanthins: miraxanthin V – insight into the excited-state deactivation mechanism from experiment and computations†

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Miraxanthin V is a betaxanthin dye occurring in Caryophyllales plants. This work describes its photophysical properties in aqueous and alcoholic solutions. After excitation at $\lambda_{\text{exc}} = 480$ nm (2.6 eV), transient absorption spectrum of the excited S₁ state was observed (S₁ \rightarrow S_n absorption band at $\lambda_{\text{probe}} = 394$ nm). The S_1 state population decays with two time constants (4.2 ps and 24.2 ps in water) corresponding to the photoexcited miraxanthin V in its two major stereoisomeric forms. The $S₁$ state decay is mainly radiationless, since the fluorescence quantum yield is low ($\Phi_F = 0.003$ in water, the smallest value among all of the studied betaxanthins). Strong correlations were obtained between the solvent viscosity and the S₁ state lifetime in linear alcohols, as well as in methanolic solutions at low temperatures. These correlations suggest that intramolecular rotation or conformational relaxation precedes the S_1 state decay. Full recovery of the electronic ground state S₀ was observed after excitations at $\lambda_{\text{exc}} = 480$ nm and 298 nm (4.2 eV), without forming the triplet T_1 state or any photoproducts. Experimental results are complemented by the ab initio S_0 state calculations using MP2 and DFT methods, and also by the excited-state calculations at the ADC(2) and TD-DFT levels of theory. The ab initio ADC(2) results give insight into the mechanism of the excited-state deactivation through the conical intersection region. We conclude that most probably the excited-state deactivation process is initiated by the rotation about the C=N double bond, present in the chain linking the electron-donor catechol to the electron-acceptor dihydropyridine group, and it is followed by collective geometry changes involving a large rotation about the C=C double bond. This mechanism seems to be universal for different betaxanthin dyes present in plants, and plays a photoprotective role. PAPER

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Betaxanthins are yellow dyes present in Caryophyllales plants.¹ Their photophysical properties, such as efficient light to heat conversion, indicate that betaxanthins play a photoprotective role in biological systems.^{2,3} Recent studies for betaxanthins in solutions show that the major S_1 state deactivation path is due to the $S_1 \rightarrow S_0$ internal conversion, while fluorescence contributes only

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weakly (Φ_F < 0.01 in H₂O), with neither the S₁ \rightarrow T₁ intersystem crossing nor the product formation ever observed.^{2,3} The internal conversion is likely enhanced by the intramolecular rotations, since an increase in the solvent viscosity causes an increase of the S_1 state lifetime and fluorescence quantum yield. Advanced quantum chemical calculations are required for a better understanding of the mechanisms responsible for the efficient internal conversion. Presently, we use miraxanthin V (MIR, Fig. 1) as a model betaxanthin dye, which probably has the highest rate of the internal conversion among betaxanthins, as follows from its lowest fluorescence intensity.⁴

Stintzing et al. have characterized **MIR** structure using ¹³C-NMR spectroscopy.⁵ Four stereoisomeric forms have been identified, existing in a $49:36:8:7$ equilibrium mixture in slightly acidified aqueous solutions. Gandía-Herrero et al. have developed a protocol for semi-synthesis of MIR by condensation of betalamic acid with dopamine, followed by HPLC purication.⁶ MIR undergoes thermal degradation in solutions, being sufficiently stable for spectroscopic studies at room temperature.

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Fig. 1 The two dominant stereoisomers of miraxanthin V in aqueous solutions.⁵

It is more stable at low temperatures in the pH range between 4 and $7^{7,8}$ The absorption maximum of **MIR** in water is at about 472 nm (2.63 eV) ,^{9,10} while its fluorescence spectrum peaks at 512 nm (2.42 eV) .⁴ The **MIR** molar absorption coefficient is expected to be similar to those of other betaxanthins, $\varepsilon_{\text{max}} = 48000$ M^{-1} cm⁻¹.^{2,3} MIR is the dominant dye in yellow Mirabilis jalapa L., Portulaca oleracea L. and Celosia argentea L. inflorescences,¹¹⁻¹³ with flower fluorescence patterns presumably attracting insects pollinators.13,14 MIR has a potential to be used as food colorant.12,15 It might also be used in DSSC (Dye Sensitized Solar Cells),¹⁶ where we need to understand the S_1 state intramolecular deactivation (the undesirable phenomenon) competing with the electron injection (the desirable phenomenon).

The aim of this work is characterization of the MIR photophysical properties using stationary and time-resolved optical spectroscopies. The solvent viscosity is probably an important factor controlling the S_1 state deactivation dynamics that will be explored. Theoretical studies will elucidate the mechanistic reasons behind the efficient radiationless $S_1 \rightarrow S_0$ transition, by calculating the excited-state minimum-energy path for the isomerization reaction. The new insights are expected to be important for betaxanthins, and also for betacyanins in general.

Results and discussion

Stationary absorption and fluorescence spectra

The recorded absorption and fluorescence spectra ($\lambda_{\rm exc}$ = 470 nm) of MIR in aqueous and alcoholic solutions are presented in Fig. 2. The absorption band maximum is weakly affected by the solvent, being located around $\lambda_{\text{abs}}^{\text{max}} = 475 \text{ nm}.$

These spectra correspond to the $S_0 \rightarrow S_1$ transition, predicted using TD-DFT calculations (see Table S1 in ESI†). The shapes of MIR absorption spectra are all similar in alcoholic solutions, differing slightly in water. This is also confirmed by calculations that predict no signicant difference between the absorption spectra of **MIR** in H_2O *vs.* MeOH (see Table S1[†]). However, the fluorescence maxima λ_f^{max} show a stronger dependence on solvent. The emission spectra in ethylene glycol and water are red-shifted $(\lambda_f^{\text{max}} \approx 510 \text{ nm}, \text{ Table 1})$ in

Fig. 2 Normalized stationary absorption and fluorescence spectra of miraxanthin V in selected solvents.

comparison to linear alcoholic and TFE solutions ($\lambda_f^{\text{max}} \approx 498$ nm). The experimental Stokes shift ($\Delta v_s = \tilde{v}_{\text{abs}}^{\text{max}} - \tilde{v}_{\text{f}}^{\text{max}}$) reaches the highest value (\approx 1530 cm⁻¹) in water.

The MIR fluorescence excitation spectrum agrees with the absorption spectrum (Fig. S1B†) implying a similarity of the shape of the absorption spectra of two major **MIR** stereoisomers in H_2O , in line with calculations (Table S1†). The contour plot of 3D fluorescence of MIR in water is shown in Fig. S2.† Fluorescence is placed in one zone at 480–640 nm in the emission scale. The peak of the contour plot shows a maximum at $\lambda_f^{\text{max}} \approx 510$ nm upon the excitation wavelength $\lambda_{\text{exc}} \approx 470$ nm. The shapes of the emission spectra weakly depend on the excitation wavelength. The fluorescence quantum yields Φ_{F} , determined with rhodamine 6G as a reference with quantum yield $\Phi_{\rm F}^{\rm ref} = 0.9,$ ¹⁷ are in the range of $3-22 \times 10^{-3}$ in the solvents tested (Table 1). The clear Φ_F dependence on the solvent viscosity in linear alcohols (Fig. S3†) agrees with the excited state behavior of other betaxanthins.^{2,3} The smallest $\Phi_{\rm F}$ value is observed in water $(\Phi_{\rm F}=3\times 10^{-3})$ which is the lowest value among the betaxanthins studied so far ($\Phi_F = 5.3$ and 7.3×10^{-3} for indicaxanthin and vulgaxanthin I in H₂O, respectively,^{2,3} see Table S2[†]). It also agrees with Gandía-Herrero et al. reporting that the smallest fluorescence intensity is observed for MIR in relation to other betaxanthins.⁴ The weak fluorescence of MIR has been explained by the lack of electron withdrawing group (such as carboxyl) in the amine and the electron-donating character of the catechol moiety.⁴

On the basis of the stationary measurements, we determined the radiative rate constant for MIR in water using the Strickler– Berg relation:¹⁸

$$
k_{\rm r} = 2.88 \frac{10^{-9} n^2}{\left\langle \tilde{\nu}_{\rm f}^{-3} \right\rangle_{\rm Av}} \int \varepsilon(\tilde{\nu}) \frac{d\tilde{\nu}}{\tilde{\nu}} \tag{1}
$$

$$
\left\langle \tilde{\nu}_{\rm f} \right\rangle_{\rm Av} = \frac{\displaystyle \int \tilde{\nu}^{-3} F(\tilde{\nu}) \mathrm{d} \tilde{\nu}}{\displaystyle \int F(\tilde{\nu}) \mathrm{d} \tilde{\nu}}
$$

where

Solvent	Viscosity ^a [cP]	$\varPhi_\mathrm{F}\left[\times 10^{-3}\right]^b$	$\lambda_{\text{abs}}^{\text{max}}$ (S ₀ \rightarrow S ₁) [nm]	λ_f^{max} [nm]	Stokes shift \lceil cm ⁻¹ \rceil
Water	0.89	3	474	511	1528
Methanol	0.54	4.7	474	500	1097
Ethanol	1.07	7.6	475	497	932
Trifluoroethanol	1.73	5.8	475	498	972
1-Propanol	1.945	14	476	498	928
1-Butanol	2.54	20	475	497	932
1-Pentanol	3.619	22	477	497	844
Ethylene glycol	16.1	15	478	508	1235
	here, $\varepsilon(\tilde{\nu})$ is the molar absorption coefficient, $F(\tilde{\nu})$ the fluores- cence intensity, $\tilde{\nu}$ the wavenumber, <i>n</i> the refractive index. ¹⁸ Note, that $F(\tilde{\nu}) = \lambda^2 F(\lambda)$. The resulting calculated value is $k_r = 2.2$ \times 10 ⁸ s ⁻¹ for MIR in water. Note that the S ₀ \rightarrow S ₁ absorption		Satisfactory fits were obtained using a double exponential function (Fig. S5†). All of the analyzed band integrals (EBI, PBI and NBI) produce quite similar time constants (Table 2), although the NBI data present a better signal-to-noise ratio.		
	band is well-defined, with the molar extinction coefficient ε_{max}		water) were tentatively assigned to the S_1 state lifetimes of the two major stereoisomers. The ratio of the partial amplitudes		
$\approx 48000 \text{ M}^{-1} \text{ cm}^{-1}$. Transient absorption spectroscopy			with the respective time-constants (0.4 and 0.6, NBI in water) reflects the relative populations of the stereoisomers in the S_1		
	Time-resolved absorption measurements were performed in the		state, which likely coincide with the S_0 state equilibrium exist-		The two time constants ($\tau_1 = 4.2$ ps, $\tau_2 = 24.2$ ps, NBI in
	spectral range of 330-660 nm with excitation at $\lambda_{\text{exc}} = 480$ nm.		ing before photoexcitation. Indeed, the ratio 0.42 : 0.58 between		
	Fig. 3 shows typical data obtained for MIR in water. Note that		the two dominant stereoisomers was estimated for MIR in		
	two separate contributions are responsible for the negative		aqueous solutions using the NMR data (Fig. 1). ⁵ Based on their		
	band: (i) the bleaching at \approx 470 nm caused by the reduced MIR		respective abundances, we infer that the EE stereoisomer is		
	concentration in the ground state S_0 upon the sample photo-		associated with the longer time constant, while the ZE stereo-		
	excitation (ii) the $S_1 \rightarrow S_0$ stimulated emission peaking at		isomer with a shorter time constant. Inspection of Table 2 also		
	\approx 530 nm. The shape of the latter band may be well approxi-		shows that the relative amplitudes of the two exponential		
	mated by the spectrum of the stimulated emission cross-section		components, describing the band integrals kinetics, depend on		
the factor of λ^4 (Fig. 3B).	obtained from the stationary fluorescence spectrum scaled by		solvent. Fig. $S6†$ shows an increase of the partial amplitude $A1$		

Table 1 Fluorescence quantum yields Φ_F ($\lambda_{\rm exc} = 470$ nm), maxima of the S₀ \rightarrow S₁ absorption bands ($\lambda_{\rm abs}^{\rm max}$), S₁ fluorescence bands ($\lambda_{\rm f}^{\rm max}$) and Stokes shift values for miraxanthin V in aqueous and alcoholic solutions

Transient absorption spectroscopy

The positive band with the maximum at 394 nm (3.14 eV) is mainly assigned to the $S_1 \rightarrow S_n$ absorption, with quite small S_0 -bleaching contribution at 394 nm. Normalization of the $S_1 \rightarrow S_n$ absorption band shows no spectral shift or band narrowing (Fig. S4†) within our signal-to-noise ratio, thus the contributions from solvation and vibrational cooling are not noticeable. A weak contribution of the solvation was to be expected, based on small Stokes shift values (below 1600 $\text{cm}^{-1})$ obtained in the stationary measurements, and a weak dependence of the Stokes shift on the solvent polarity (compare water to the less polar methanol, Table 1).

In order to retrieve the S_1 state lifetime, band integral kinetics were computed using

$$
BI = \int_{330}^{670} \Delta A(t, \lambda) \frac{d\lambda}{\lambda},
$$
 (2)

where the integration is done over the entire probe spectral range (EBI, 330–670 nm). Note that the usage of the band integral minimizes the potential contributions of the vibrational cooling and solvation effects.¹⁹ Band integral kinetics were also calculated separately for positive (PBI, 330–430 nm) and negative (NBI, 430–670 nm) transient absorption bands.

The two time constants ($\tau_1 = 4.2$ ps, $\tau_2 = 24.2$ ps, NBI in water) were tentatively assigned to the S_1 state lifetimes of the two major stereoisomers. The ratio of the partial amplitudes with the respective time-constants (0.4 and 0.6, NBI in water) reflects the relative populations of the stereoisomers in the S_1 state, which likely coincide with the S_0 state equilibrium existing before photoexcitation. Indeed, the ratio 0.42 : 0.58 between the two dominant stereoisomers was estimated for MIR in aqueous solutions using the NMR data (Fig. 1).⁵ Based on their respective abundances, we infer that the EE stereoisomer is associated with the longer time constant, while the ZE stereoisomer with a shorter time constant. Inspection of Table 2 also shows that the relative amplitudes of the two exponential components, describing the band integrals kinetics, depend on solvent. Fig. S6† shows an increase of the partial amplitude A_1 with solvent polarity π^* . We can expect, that a more polar solvent stabilizes a more polar stereoisomer. Indeed, theoretical predictions show that the ZE form has a larger dipole moment than EE form (Tables S1 and S3†).

The global analysis of the transient absorption data results in time constants ($\tau_1 = 2.2$ ps, $\tau_2 = 20.8$ ps, in water) similar to those deduced from the band integral kinetics (Table 2); note similar shapes of the two respective decay-associated spectra (Fig. 3C).

The **MIR** transient absorption data in H_2O (Fig. 3A) were also analyzed by independent fitting of the individual kinetic traces (Fig. S7†). The obtained time constants approximately reproduce the values retrieved in the global analysis (Fig. 3C). The exception is at probe $\lambda = 493$ nm where the individual timeconstants are too short. This can be explained by contribution from an intermediate absorption,²⁰ indeed species nascent upon $S_1 \rightarrow S_0$ deactivation can be structurally unrelaxed and possess a red-shifted absorption band in relation to the relaxed **MIR** in the S_0 state.

Time-resolved absorption measurements were also performed for MIR in the range of 825–1380 nm (the NIR range) with the excitation at $\lambda_{\text{exc}} = 485$ nm in water (Fig. S8A†). The

Fig. 3 (A) Transient UV-vis absorption spectra obtained miraxanthin V in water ($\lambda_{\rm exc}$ = 480 nm). (B) Reversed UV-vis absorption and cross-section σ_E spectrum for stimulated emission. (C) Decay associated spectra obtained by the two-exponential fit.

recorded $S_1 \rightarrow S_m$ (*m* > 1) absorption band is located around \approx 920 nm. The band integral over the entire spectral range (825– 1380 nm) was calculated. The two time constants obtained from the double exponential fit: 2.7 ps (0.41) and 21.9 ps (0.59) , are in a reasonable agreement with those derived from the global analysis (Fig. S8B†). Thus, the determined amplitude associated spectra (Fig. $S8B\dagger$) likely reflect respective stereoisomers $S₁$ state absorption bands. Similar transient absorption data are produced for MIR in MeOH (Fig. S9A†), with the amplitude associated spectra resembling those in H_2O .

The long-lived stereoisomer in the S_1 state mainly contributes to the stationary fluorescence spectra and to the fluorescence quantum yield Φ_F . This also agrees with inspection of the strength of the stimulated emission transient bands at longer delay times exceeding 6 ps, Fig. 3A. The radiative rate constant for this stereoisomer was calculated using the standard expression:

$$
k_{\rm r}\tau_2 = \Phi_{\rm F} \tag{3}
$$

where Φ_F is fluorescence quantum yield, and τ_2 is the lifetime of the long-lived stereoisomer. Linear regression was performed (Fig. S10†) and the radiative rate constant of MIR k_r was determined at $k_r = 2 \times 10^8 \text{ s}^{-1}$. Note that k_r agrees with value 2.2 \times 10^8 s⁻¹ calculated from the Strickler-Berg relation (1) for **MIR** in water.

Very similar spectral shape of transient absorption data (Fig. S11A†) at delays longer than 3 ps was obtained for MIR in water using excitation at $\lambda_{\rm exc}$ = 298 nm instead of 480 nm (Fig. 3A). The decay time-constants retrieved from band integral analysis (NBI, $\tau_1 = 3.8$ ps, $\tau_2 = 23.6$ ps) are comparable to those obtained with $\lambda_{\rm exc} = 480$ nm. Inspection of transient absorption kinetics at probe wavelengths 460 nm and 520 nm (Fig. S11E†) shows an extra decay (460 nm) and rise (520 nm) with a timeconstant of 0.6 ps. This time-constant likely reflects vibrationally hot population of the S_1 state generated through the $S_n \rightarrow S_1$ internal conversion process, following the initial $S_0 \rightarrow S_n$ (n > 1) state photoexcitation (with $\lambda_{\text{exc}} = 298$ nm). Thus, the vibrationally hot S_1 state contributes to the fast decay of the stimulated emission at probe 460 nm. In contrast, probe 520 nm shows the initial growth (0.6 ps) of the stimulated emission signal because the vibrationally relaxed $S₁$ state population increases. The early changes caused by the S_1 state vibrational relaxation are virtually absent with a long excitation wavelength ($\lambda_{\rm exc}$ = 480 nm, Fig. S11E[†]), since under these conditions a small excess of vibrational energy is produced in the S_1 state. A very fast heat dissipation capability of MIR can be explained by strong solute–solvent coupling through hydrogen bonding. Note that the vibrational cooling in case of betanin has been also described by a short time-constant (≈ 0.9 ps in H_2O).²¹

The transient absorption measurements were also performed for **MIR** in alcoholic solutions with the excitation at $\lambda =$ 480 nm (Fig. S12†). The time constants deduced from both global and band integral analysis all depend on the solvent viscosity in linear alcohols (Table 2). Note that while the dynamics is affected by the solvent, the shape of the transient absorption spectra remains unchanged (Fig. S12†). Additional measurements were performed at temperatures between 180 and 300 K for MIR methanolic solutions, and again a correlation of the S_1 lifetime vs. viscosity η was observed, although the influence of temperature should also be considered (Fig. S13, data in Table S4†). These viscosity-dependent data may be described by a commonly used empirical model.²² Thus, the

Table 2 Miraxanthin V S₁ state decay time constants retrieved from the band integrals and the global analysis of the transient UV-vis data^a

		π^*	Positive BI		Negative BI		
Solvent	Viscosity [cP]		A_1, A_2	$\tau_1, \tau_2^{\ b}$ [ps]	A_1, A_2	τ_1, τ_2^c [ps]	Global analysis τ_1 , τ_2^c [ps]
Water	0.89	1.09	0.42, 0.58	3.7, 22.7	0.4, 0.6	4.2, 24.2	2.2, 20.8
Methanol	0.54	0.60	0.34, 0.66	6.1, 33.3	0.35, 0.65	6.9, 36.1	5.6, 35.9
Ethanol	1.07	0.54	0.23, 0.77	5.6, 56.7	0.24, 0.76	6, 58.4	5.7, 58.9
Trifluoroethanol	1.73	0.73	0.32, 0.68	5.3, 40.1	0.34, 0.66	6.1, 43.4	4.5, 40.7
1-Propanol	1.945	0.52	0.22, 0.78	8,79.9	0.23, 0.77	10.7, 84.2	6.2, 80.8
1-Butanol	2.54	0.47	0.23, 0.77	7.5, 103.5	0.22, 0.78	9.3, 106.8	5.8, 102.7
1-Pentanol	3.619	0.40	0.27, 0.73	8.1, 121.4	0.27, 0.73	10.4, 127.8	7.7, 126.7
Ethylene glycol	16.1	0.92	0.36, 0.64	9.6, 82.2	0.37, 0.63	9.5, 83.2	7.6, 79
	rates of the S_1 decay in methanol as a function of temperature, and also in linear alcohols at $T = 22$ °C, may all be fitted using						
the following equation: ²³⁻²⁶							
	$k = \frac{\nu}{\eta^{\alpha}} \exp\left(-\frac{E_{A}}{RT}\right)$			$\left(4\right)$			the plane that is approximately parallel to the plane containing the electron-accepting moiety and the C_{11} , C_{10} , N_9 and C_8 atoms. The two moieties contained in the above-mentioned planes are linked by the $C_7 - C_8$ single bond bridge - a linkage of a geometry typical for saturated hydrocarbons. Such molec- ular arrangement makes the potential energy profile asym-
	where η is the solvent viscosity, T temperature, R the ideal gas						metric with respect to the rotations described in this paragraph.
	constant, k the reciprocal of the S_1 state lifetime. Here α indi-						
	cates the influence of viscosity (dynamical molecule-solvent						
	friction) upon the overall S ₁ state relaxation, with $0 \le \alpha \le 1$, E_A						Both the B3LYP/cc-pVDZ and MP2/cc-pVDZ calculations in gas phase confirm that the ZE isomer is by ≈ 0.16 eV higher in energy than the most stable EE form. The calculated excitation
	is the activation energy of the barrier between the locally excited						
	state and the twisted state, ν describes the pressure exerted by						energies along with the oscillator strengths for different rota- meric forms of MIR (computed with the aid of the TD-DFT
	MIR at unit viscosity. After applying the logarithm, we obtain:						method for different solvents and in vacuum) are presented in Tables S1 and S3, \dagger with ADC(2) results given for comparison.

$$
k = \frac{\nu}{\eta^{\alpha}} \exp\left(-\frac{E_{\rm A}}{RT}\right) \tag{4}
$$

$$
\ln(k) = \ln(v) - \alpha \ln(\eta) - \frac{E_{A}}{RT}
$$
 (5)

The fitted surface is presented in Fig. S14.[†] The parameter values $\alpha_2 = 0.74 \pm 0.04, E_{A2} = 57 \pm 33$ cm⁻¹, and $\alpha_1 = 0.25 \pm 10^{-1}$ $0.10, E_{A1} = 350 \pm 80 \text{ cm}^{-1}$ were obtained for the long- and shortlived stereoisomers, respectively. As expected,²⁶ the higher barrier $(E_{A1} > E_{A2})$ for the internal rotations or conformational relaxation in the singlet excited state should be accompanied by a weaker dependence of the rate k on viscosity ($\alpha_1 < \alpha_2$). The activation energy for both stereoisomers is small and comparable to kT value in room temperature (\approx 210 cm⁻¹), thus the solvent viscosity is the main factor controlling the S_1 state lifetime. Comparing the α values ($\alpha_2 = 0.74$ vs. $\alpha_1 = 0.25$), we deduce that the stereoisomer 2 (EE) experiences a stronger friction against the solvent molecules. Indeed, a more hampered rotation around the central bridge bonds is expected for the EE stereoisomer (Fig. 1) due to the increased friction of the dopamine moiety against the solvent molecules.

Quantum chemical calculations

The NMR data indicate that MIR exists in solution in an equilibrium between the two main stereoisomers: EE (9E,11E) and ZE (9Z,11E), which are the most abundant (see Fig. 1).⁵ The quantum chemical calculations produced the optimized structures of the most abundant stereoisomers shown in Fig. S15.† In both the forms the electron-donating catechol moiety lies in

the plane that is approximately parallel to the plane containing the electron-accepting moiety and the C_{11} , C_{10} , N_9 and C_8 atoms. The two moieties contained in the above-mentioned planes are linked by the C_7-C_8 single bond bridge – a linkage of a geometry typical for saturated hydrocarbons. Such molecular arrangement makes the potential energy profile asymmetric with respect to the rotations described in this paragraph. Both the B3LYP/cc-pVDZ and MP2/cc-pVDZ calculations in gas phase confirm that the ZE isomer is by ≈ 0.16 eV higher in energy than the most stable EE form. The calculated excitation energies along with the oscillator strengths for different rotameric forms of MIR (computed with the aid of the TD-DFT method for different solvents and in vacuum) are presented in Tables S1 and S3,† with ADC(2) results given for comparison. The $S_0 \rightarrow S_1$ transition was computed at 401 nm (3.09 eV) with oscillator strength $f = 0.20$ for *EE*, and at 422 nm (2.94 eV) with lower intensity for ZE in vacuum (Table S1†). This band gets significantly stronger for both forms in polar solvents (water and alcohols), as its oscillator strength f increases to ≈ 0.6 (see Table S1†). Moreover, the calculations show that the two absorption peaks for both the EE and ZE forms are even closer in a given solvent (alcohol or water) than in vacuum. This computational result suggests that the photoexcitation at $\lambda_{\rm exc} \approx$ 480 nm in a given solvent results in an initial population of the isomers in the S_1 state that reproduces the equilibrium existing in the S₀ state. The molecular orbitals involved in the $S_0 \rightarrow S_1$ transition were plotted with isosurfaces at 0.03 value (Table S5†). The major contribution to this transition is provided by HOMO $(\pi) \rightarrow$ LUMO (π^*) component, while the second most relevant HOMO -1 $(\pi) \rightarrow$ LUMO (π^*) contribution is a little smaller for both the forms. The coefficients of electronic configurations contributing to the ADC(2) S_1 state wavefunction are given in Table S5.† Note that the $S_0 \rightarrow S_1$ transition has a π,π^* character for both the *EE* and *ZE* forms. The shape of orbitals indicates that after excitation within the Franck-Condon area in the S_1 state, a part of the electron density moves from the electron-donating catechol moiety to the one of the electron-accepting carboxyl groups,²⁷ suggesting that MIR may act as a sensitizer in DSSC. The presence of COOH groups in MIR facilitates anchoring of this dye to the TiO₂ layer.²⁸ The difference between the experimental excitation wavelength (474

Table 3 Comparison of the geometric parameters (bond lengths, in Å), adiabatic energies (E_a , in eV) and dipole moments (μ , in debye) for the equilibrium ground state (S_0) and the excited state (S_1) geometries optimized at the MP2/cc-pVDZ (for S_0) and the ADC2/cc-pVDZ (for S_1) levels of theory. The fluorescence energy (E_f , in eV) and oscillator strength $f(S_1 \rightarrow S_0)$ are given for the S₁-state

	C_8-N_9	$C_{10} = N_9$	C_{10} – C_{11}	$C_{11} = C_{12}$	$E_{\rm a}$ [eV]	E_f [eV]	μ [D]	$f(S_1 \rightarrow S_0)$
S_0 EE	1.455	1.299	1.454	1.374	0.00		2.8	
S_1 EE $\pi\pi^*$	1.447	1.318	1.427	1.415	3.32	2.98	5.1	0.541
$S_1 \theta = 90^\circ$	1.455	1.434	1.3334	1.486	1.83	-1.55	8.3	0.000
S_1 ZE $\pi\pi^*$	1.455	1.321	1.429	1.420	3.38	2.96	5.3	0.512
S_0 ZE	1.457	1.302	1.460	1.377	0.16		3.4	

nm) and its predicted theoretical value (401 nm) is quite significant (3837 $\text{cm}^{-1} = 0.48 \text{ eV}$), while still at the level of the mean absolute error for the B3LYP functional (\approx 0.34 eV for excited singlet states).²⁹ Note that our attempt to use either a CAM-B3LYP functional or a larger aug-cc-pVTZ basis set with B3LYP functional produced no improvements to the TD-DFT results.

The strongly absorbing $\pi\pi^*$ state is always the lowest excited state in the spectrum. Thus, we presume that this is the initially photoexcited state, active in the photophysics of MIR, however, the second lowest excited state has $n\pi^*$ character and lies approx. 0.5 eV above the $S_1(\pi \pi^*)$ state, according to the ADC(2)/ cc-pVDZ method (see Table S3†).

As already mentioned, we expect that upon UV excitation of the $S_0 \rightarrow S_1(\pi \pi^*)$ transition, the initially formed $S_1(\pi \pi^*)$ state in solution exists in two major forms: $S_1(EE)$ and $S_1(ZE)$, with the population ratio similar to that of the S_0 state. Moreover, the transient absorption data show two characteristic time constants assigned to the S₁ state lifetimes. Those two $S_1(\pi \pi^*)$ state forms are almost isoenergetic in the gas phase, according to the ADC(2)/cc-pVDZ results (compare the adiabatic energies of the $\frac{1}{2}\pi\pi$ *-excited states in Table 3). The electronic structure analysis indicates locally-excited character of the respective minima (see Table S5†). Both the electron-donating and accepting orbitals are localized on the dihydropyridine moiety of MIR. However, the experimentally determined S_1 state lifetimes (4.2 ps and 24.2 ps in water) are relatively short. This suggests existence of an efficient molecular mechanism for the radiationless decay of the electronic excitation. A possible explanation for the fast excited-state deactivation is given by the existence of a low-lying S_1-S_0 conical intersection (CI) seam (see the colored beige rectangles in Fig. 4 and bottom-right circle in Fig. 5), where non-adiabatic transition takes place from the excited state (S_1) to the (S_0) electronic ground state.^{30,31}

Such transitions are the most effective in the geometries where the excited and the ground states become degenerate. In our experiments the dependence of the $MIR-S₁$ state lifetimes on the solvent viscosity was determined (Fig. S14†). This implies that the driving coordinate for the excited-state decay should be a large-amplitude motion described as a rotation about one of the torsional angles. Studying the MIR chemical structure, we may consider several potential reaction coordinates that may drive the system towards the S_1-S_0 CI region.

There are six torsional angles which should be considered as a driving coordinate. These are two double bonds: $C_{11} = C_{12}$ and C_{10} =N₉, shortly referred to as C=C and C=N, and four single bonds: C_1 – C_7 , C_7 – C_8 , C_8 – N_9 , and C_{10} – C_{11} . Single bonds can be immediately excluded from consideration for a following reason; the shape of the minimum energy profile (MEP) of the excited state calculated along the coordinate described as a rotation about a single bond is known to be parallel to the MEP along the same coordinate for the S_0 state, $30,32-34$ thus such rotation would demand overpassing the excited-state energy barrier as it also exists in the S_0 state. This implies that single bond rotations cannot lead to the fast deactivation pathway through the CI regions. We are pointing to the conclusion that rotations about double bonds are good candidates for the driving coordinates for fast deactivation process ongoing in the excited state. Why is that? **BSC Arbaness**

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There are several known classes of molecules where such excited-state rotations around different type of double bonds

Fig. 4 Potential-energy profiles of miraxanthin V in the excited states: $\pi\pi^*$ state (blue triangles), 1 n π^* state (red squares) determined at the ADC(2)/def-SV(P) level (MP2/def-SV(P) for the ground state) along the minimum energy path (solid line) for the isomerization between the EE and ZE stereoisomers. $S_0^{(\pi \pi^*)}$ (empty black triangles) and $S_0^{(\pi \pi^*)}$ (empty black squares) denote the energy of the ground state calculated along the minimum-energy path of the appropriate excited state (dashed curves). Shaded rectangles denote areas where the ADC(2) iteration cycle ceases to converge due to degeneracy of the S_1 and the S_0 states. Empty blue triangles and empty red squares denote the points corresponding to the given ${}^1\!\pi\pi^*$ or ${}^1\!n\pi^*$ excited state energy profile for which the ADC(2) method ceased to converge.

Fig. 5 Minimum-energy profile of the $\frac{1}{n}\pi^*$ (blue circles) and $\frac{1}{n}\pi^*$ (red circles) excited state of miraxanthin plotted as a function of θ (C₈–N₉=C₁₀–C₁₁) and θ (C₁₃–C₁₂=C₁₁–C₁₀) coordinates illustrating the deactivation pathway of the ZE form according to two alternative scenarios: (A) the initial rotation about the C=N bond (as in Fig. 4) is followed by strong twisting of the C=C bond, (B) the initial rotation about the C=C bond (as in Fig. S16†) followed by large twisting of the C=N bond to enter the CI region. The red circle in the bottom right corner of the surfaces represents the conical intersection region: $Cl(S_0 - S_1)$ of a so-called "collapsed" structure. The results were obtained with the aid of the ADC(2)/def-SV(P) method.

are observed, such as azobenzenes where rotation around the N=N double bond leads to profound conformational changes.³⁵⁻³⁸ The excited-state rotation around the C=C double bond occurs in the extensively studied stilbenes (Z/E) isomerization), $39,40$ cyanines, $41-43$ the 3-hydroxy-picolinic acid, $44,45$ and the $7HQ^{46,47}$ molecule. Quite recently, a new photoswitching molecule N-(3-pyridinyl)-2-pyridinecarboxamide (NPPCA) has been investigated, where excited state rotation occurs about the $C=N$ double bond,⁴⁸ present also in the **MIR** chromophore; the same effect was observed in light-switchable peptides.^{33,39}

Upon UV excitation, a double bond nominally short in the S_0 state, may get weaker and longer in the excited state. This effect was detected in a series of derivatives of 7-hydroxy-quinoline (7HQ)⁴⁷ and salicylidene methylamine.⁴⁹ Thus, a hindered rotation around a nominally double bond in the S_0 state may become allowed in the S_1 -excited state due to lowering of the barrier, explained by this bond becoming a longer and weaker (single) bond in the S_1 state.

Indeed, according to the calculations, comparing the ground- and excited-state equilibrium geometries in MIR, both the $C=$ C and $C=N$ double bonds undergo elongation upon excitation, with the single C–C bond simultaneously getting shorter. This is observed for both the EE and ZE rotamers. The respective geometric parameters are compared in Table 3.

Our calculations indicate an elongation of the $C=C$ bond by 0.039 and 0.043 Å for the EE and ZE forms, respectively, comparing their equilibrium ground-state geometry to their excited-state equilibrium geometry; similarly, we obtained elongation of the C=N bond by 0.019 \AA for both forms. To get insight into the mechanism of deactivation process of MIR the excited-state MEPs were calculated along two driving coordinates, separately: rotations about the C_{10} =N₉ and C_{11} =C₁₂ double bonds. Until the system is in the $^1\pi\pi^*$ -state minimum, both planes including two ring moieties are almost parallel to

one another. Comparison of the estimated in calculations excited-state barriers, indicates that in the initially photoexcited $S_1(\pi \pi^*)$ state, the barrier for the rotation about the $C_{10} = N_9$ bond is lower than analogous barrier for the $C_{11} = C_{12}$ and $C_{10}-C_{11}$ rotations, for a given stereoisomer, especially for EE (cf. 4, S16 and S17†). That is why we may assume that it is a rotation about the $C=N$ bond that initiates the deactivation process of overpassing the $^1\pi\pi^*$ -state barrier. Such evolution of the system toward the $\frac{1}{2}\pi\pi^*$ -state barrier one may observe in Fig. 5A for which the $C=C$ bond does not change much.

The initial C=C rotation as $^1\!\pi\pi^*$ -state deactivation path can be considered for ZE stereoisomer (Fig. 5B), since the barrier for the C=C rotation is only slightly higher than that for $C=N$ rotation. However in both scenarios (Fig. 5A and B) the initial bond rotation along one coordinate is followed by twisting of the second coordinate before reaching the conical intersection point, where MIR deactivates to the ground state.

The conical intersection region is reached upon the initial rotation about the $C=N$ double bond of the ZE form by the dihedral angle of $\approx 30^\circ$. The most relevant orbitals that contribute to the nonadiabatic transition between the excited and the ground state in the S_1/S_0 conical intersection are shown in Table S7.† The most important in this arbitrarily chosen geometry is the $n\pi^*$ contribution, while the $\pi\pi^*$ contribution is also present. However, based on this contribution set we can't state that the conical intersection is purely of the $n\pi^*/S_0$ nature. Its character is thus a mixture of the two contributions.

The photophysical deactivation of MIR proceeds as follows. After photoexcitation of the EE form (as well as ZE), the $S_1(\pi \pi^*)$ state is populated. As shown in Fig. 4, the potential-energy profile of the $S_1(\pi \pi^*)$ excited state has a small barrier of 0.09 eV for the rotation about the $C=N$ bond (see blue curve in the upper-left corner of Fig. 4). The analogous $S_1(\pi \pi^*)$ -state barrier for the ZE form is lower, of 0.04 eV (see the blue curve in the upper-right

corner of Fig. 4). Such a small barrier may be easily overcome in experiments undertaken at room temperature. It is also evident that a rotation by just about $25-30^\circ$, either from the EE or from the *ZE* $S_1(\pi \pi^*)$ -state minimum form, may put the system into a close proximity of the CI region (beige areas). The process is also shown in Fig. 5A which illustrates the deactivation pathway eye-guided along the points presented in the two-coordination space. In this region $(\theta(C=N) \approx 30^{\circ})$, the S₁($\pi \pi^*$) state meets the descending $-\int n\pi^*$ state – which was found to be the second excited state in the absorption spectrum both for the EE and ZE forms (Table S3†), and $S_1(\pi \pi^*)$ populates this ${}^1n\pi^*$ state. This intersection of the $\pi \pi^*$ and $n \pi^*$ states is illustrated in Fig. 5A, where at the intersection point the two states have the same driving coordinate values what helps the transition from one state into another because the barrier between the states disappears. In general, however, one may note that the intersection point presented in Fig. 5A is still just an apparent crossing, since the remaining geometry parameters in the two states can differ. As soon as the rotation about the $C=C$ bond is allowed, the optimization procedure immediately heads to the "collapsed" structure with the C₁₃-C₁₂=C₁₁-C₁₀ dihedral $\approx 80^\circ$ (shown in the bottom right corner of Fig. 5A), which is a structure with $\approx 60^{\circ}$ angle between the two planes of the ring moieties. Upon large amplitude motion, including other degrees of freedom such as rotation about single bonds in the bridge, the energy of the excited state rapidly decreases and collective relaxation immediately drives the system toward the $CI(S_1-S_0)$ region where the nonadiabatic transition to the S_0 state takes place. Unfortunately this leads the ADC(2) method to cease in convergence. BSC Advances

corner of Fig. 4). Such a small barrier may be easily oscenent in the strong time serient barrier corner in the strong corner in the strong corner in the strong common common barrier correl in the strong com

Exactly the same mechanism of excited-state deactivation was made for the EE and ZE tautomers (compare the second left and the second right structures in Fig. 4). Moreover, since after overpassing the $\frac{1}{2}\pi\pi^*$ -state barrier, the system rotates $\approx 80^\circ$ about the $C=C$ bond, it can be assumed that the system will come back to the same S_0 state stereoisomer which was photoexcited (aborted isomerization path), instead of proceeding through the higher-in-energy central path connected with lonely rotation about the $C=N$ double bond, in which the $C=$ C rotation is not present.

The involvement of the $n\pi^*$ state in the overall excited-state deactivation process through the conical intersection region(s) in MIR is an interesting idea. Additional evidence in support is given by the fact that the $n\pi^*$ -state is found driving the same processes in several other molecules, e.g. for the hydrogenbonded azo aromatic molecules,³⁷ dyes with the molecular scaffold of 2-hydroxyazobenzene (Sudan I and Orange II),³⁸ and for the picolinic acid derivatives.^{44,45} In the latter case the 1 n π^* state lies below the $^1\pi\pi^*$ excited state in all of four tautomeric forms studied. Apparently the electronically excited picolinic acid decays non-radiatively through the conical intersections of the $1n\pi^*$ state with the ground state, quenching the fluorescence of all of the tautomeric forms.⁴⁴

Conclusions

Experimental data show an increase of the $MIR S₁$ state lifetimes with an increase of solvent viscosity in linear alcohols. We

postulate that torsional relaxation of the $S_1(\pi,\pi^*)$ state around the central bonds (N=C and/or C=C) leads to $S_1 \rightarrow S_0$ radiationless relaxation. This rotational motion is slower in a more viscous solvent, therefore the internal conversion rate is limited by the time required to reach the necessary geometry. The $S_1 \rightarrow$ $S₀$ radiationless relaxation is the main deactivation channel for the S₁ state. Fluorescence is a minor relaxation pathway (Φ _F < 0.03 in all of the solvents tested), with a secondary role in the overall S_1 -population deactivation. We obtained no indications of photoproduct or T_1 state formation from the S_1 state, with the full S_0 state recovery observed. The S_1 state population exhibit biexponential decays in solution attributed to the presence of two major stereoisomers in the ground state of MIR. While numerous photophysical properties, including stationary UVvis absorption, fluorescence, spectral locations of the $S₁$ state absorption band and the S_1 deactivation paths make MIR similar to other betaxanthins studied so far (see Table S2† for the comparison), $2,3$ MIR exhibits exceptionally low fluorescence quantum yield Φ_F , and relatively short S_1 state lifetimes. These features are controlled by structural factors, such as the electron-donating character of the catechol moiety, while the more fluorescent betaxanthins (vulgaxanthin I, indicaxanthin) typically possess an electron-withdrawing group (e.g. carboxyl) in the amine, instead of catechol.

Ab initio calculations give insight into the mechanism of fast $S_1 \rightarrow S_0$ internal conversion in MIR. Easy access to the CI region through the rotation about the C $=N$ and C $=C$ bonds explains fast radiationless deactivation of the singlet excited state and its short lifetime. The rotation about the C $=N$ double bond seems to initiate the excited-state deactivation process, with an energy barrier lower than that for the rotation about the $C=C$ bond, and much lower than that for the rotation about the C–C single bond in the central bridge of the molecule. However, as soon as the system leaves the initially photoexcited $\frac{1}{l}\pi\pi^*$ -state minimum, collective geometry changes, foremost the rotation about the $C=C$ double bond, drives the system toward the CI region and next to the ground state repopulation, closing the photocycle through the aborted isomerization process. The twisting angle of $\approx 30^\circ$ is required for the system to reach the CI region, thus it explains found influence of solvent viscosity on $S₁$ lifetimes and fluorescence quantum yields in solution. Since the other betaxanthins studied so far^{2,3} also have two distinct values of the S_1 state lifetime, we expect them to have a similar S_1 state deactivation mechanism, with the S_1 state decay rate controlled by structural factors, location of the CI region, and solvent viscosity.

Experimental section

Pure MIR was obtained as proposed by Gandía-Herrero et al.⁶ Before the semi-synthesis of MIR, a separation of the synthetic substrate, indicaxanthin (IND), from the orange-yellow fruits of Opuntia ficus-indica L. Mill, was performed as described previously.² During the semi-synthesis, dopamine was added in a 25 fold molar excess to an alkaline solution of betalamic acid obtained directly after IND hydrolysis at pH 11.4 through addition of 25% NH₄OH. After 20 min since the addition of dopamine,

the solution was acidified to pH 6.5 by formic acid and chromatographically purified by open column chromatography on C18 sorbent (Merck, Darmstadt, Germany).² Typically, the C18 sorbent was activated with 100% methanol and then rinsed with 1% aqueous formic acid. The samples were applied to a column rinsed again with 1% formic acid and betaxanthin was eluted with methanol. The eluates were pooled and preconcentrated under reduced pressure at 25 °C. Concentrated extract was submitted to semipreparative flash chromatography (preparative HPLC system with LC-20AP pumps, UV-VIS SPD-20AV detector and LabSolutions 5.51 operating software, Shimadzu Corp., Japan) on a C18 flash column (Interchim, France) for the separation of MIR.²¹ Further purification of the pigment was performed on an semipreparative HPLC column Luna C18(2) 250×30 mm i.d., 10 µm (Phenomenex, Torrance, CA, USA).⁵⁰

The details of femtosecond UV-vis-NIR transient absorption spectra have been described elsewhere.² The ultrafast laser system consists of a short-pulse titanium-sapphire oscillator (Mai-Tai, Spectra Physics, 70 fs) followed by a high-energy titanium-sapphire regenerative amplifier (Spitfire Ace, Spectra Physics, 100 fs). The 800 nm beam was split into two beams to generate: (1) a pump ($\lambda_{\rm exc}$ = 480, 485 or 298 nm) in the optical parametric amplifier (Topas Prime with a NirUVis frequency mixer) and (2) probe pulses – white light continuum in the UVvis range by using a CaF₂ plate (330-660 nm, Ultrafast Systems, Helios) or in the NIR range by using a YAG crystal (830– 1390 nm, Ultrafast Systems, Helios). The remaining 800 nm photons in the probe pulse were filtered out before the sample. The pump pulse energy was typically about $0.5-0.8$ μ J. In most of the transient absorption experiments the absorbance was about 0.5 at the excitation wavelength in a 2 mm optical path quartz cell. Sample solution was stirred by a Teflon-coated bar. The entire set of pump–probe delay positions was repeated at least four times to ensure data reproducibility. The transient absorption data were corrected for chirp of the white light continuum. The instrument response function (IRF) was about 200 fs (FWHM) in the experiments with 480 nm excitations, and 350 fs (FWHM) with the excitation at 298 nm. The data were analyzed by determination of the band integral kinetics and fitted with a double-exponential function. The convolution with a Gaussian response function was included into the global fitting procedure (ASUFIT program), and satisfactory fits were obtained. Temperature-dependent experiments between 180 and 300 K were performed in a liquid nitrogen cryostat (Janis VPF-100). The solutions were placed into a cuvette consisting of two parallel plastic windows, separated by 1.5 mm distance. The temperature was measured by two thermocouples, located over and under the sample. Paper

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Fluorescence emission and excitation spectra were recorded on a Horiba Jobin-Yvon-Spex Fluorolog 3-22 spectrofluorometer. Right-angles geometry was used in all of the steady-state emission measurements. The sample emission and excitation spectra were recorded with three replicates at a reduced scanning rate. The experiments were performed on 3.5 mL of sample solutions contained in a quartz cell (1 cm \times 1 cm) with the MIR absorbance not exceeding 0.1 at the excitation wavelength (2 \times 10^{-6} M in water). The excitation wavelength was set at $\lambda_{\rm exc} =$

470 nm for the fluorescence measurements. Fluorescence spectra were collected with 1 nm excitation and emission slits, using 0.5 s integration time. Fluorescence spectra typically were not corrected for contribution of the solvent Raman scattering (Fig. S1A†).

For the 3D fluorescence spectra, excitation and emission slits were 2 nm; the acquisition interval was 2 nm and the integration time was 0.2 s. 3D emission spectra were collected between 480 to 675 nm in the 355–550 nm range of the excitation wavelengths, spaced by 5 nm intervals in the excitation domain. Stationary UV-vis absorption spectra were recorded using a Jasco V-550 spectrophotometer. Alcohols and rhodamine 6G were purchased from Aldrich, water was purchased from Fluka. Unless otherwise stated, experiments were performed at room temperature (22 \degree C).

Computational methods

To simulate the UV-absorption spectra of MIR, the equilibrium geometries of the molecular system in its closed-shell ground singlet state (S_0) were determined at the level of density functional theory (DFT) with the use of the of the B3LYP⁵¹ hybrid correlation–exchange functional in conjunction with the ccpVDZ⁵² correlation-consistent basis set as implemented in TURBOMOLE software package.⁵³ In the calculation of the excitation energies, mimicking the absorption spectra, the basis set was augmented with diffuse functions (aug-cc-pVD Z^{52}) and single-point TD-DFT calculations were done on the top of the geometries optimized with the cc-pVDZ basis set. Such calculations are supposed to be in good agreement with the experimental UV-absorption spectra. In the next step, the geometry was optimized in the Conductor-like Screening Model⁵⁴ (Cosmo) implemented in TURBOMOLE, to have comparison of the UV-absorption spectra obtained in the solvent and vacuum. Such satisfactory comparison was done in our report on betanin,²¹ based on TD-DFT calculations with the B3LYP functional⁵¹ with 6-311++G(3df,3pd) basis set⁵⁵ (the largest Poplestyle basis set) and polarizable continuum model (PCM) for the solvent,^{56,57} as implemented in Gaussian software package.⁵⁸ Thus, here we use the same approach for MIR.

However, the TD-DFT method is also known to fail in prediction of the excited states with strong charge-transfer,^{59,60} with small overlap of the two orbitals contributing to the electron excitation (e.g. HOMO and LUMO). To bypass this problem in predicting the excited-state potential energy profile (PE), additional calculations were performed, where the equilibrium geometry of the molecular system in its closed-shell singlet ground state (S_0) was determined with the MP2 method.⁶¹ Excitation energies, equilibrium geometries, and response properties of the lowest singlet excited states were calculated using the algebraic diagrammatic construction method of the second order, $ADC(2)$.^{62,63} To get insight into the mechanism of excited-state deactivation process, the minimum-energy reaction path (MEP) along the photophysically relevant reaction coordinates in the lowest excited singlet state were also determined with the ADC(2) method. For a suitably chosen driving coordinate, all other nuclear degrees of freedom were optimized for a given value of the driving coordinate. To allow costeffective exploration of the high-dimensional potential-energy surfaces, the standard split-valence double-zeta basis set with polarization functions on the heavy atoms (def-SV(P)) was employed in these geometry optimizations.

Ab initio calculations for all of the model systems were performed with the TURBOMOLE program package,⁵³ making use of the Resolution-of-the-Identity (RI) approximation for the evaluation of the electron-repulsion integrals.⁶⁴

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