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Visible-to-ultraviolet (<340 nm) photon upconversion by triplet-triplet annihilation in solvents[†]

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In this article, visible-to-ultraviolet photon upconversion (UV-UC) by triplet-triplet annihilation in the emission range shorter than 340 nm, which has not been explored well, is presented and the relevant physicochemical characteristics are elucidated. Investigations were carried out in several deaerated solvents using acridone and naphthalene derivatives as a sensitizer and emitter, respectively. Both upconversion guantum efficiency and sample photostability under continuous photoirradiation strongly depended on the solvent. The former dependence is governed by the solvent polarity, which affects the triplet energy level matching between the sensitizer and emitter because of the solvatochromism of the sensitizer. To elucidate the latter, first we investigated the photodegradation of samples without the emitter, which revealed that the sensitizer degradation rate is correlated with the difference between the frontier orbital energy levels of the sensitizer and solvent. Inclusion of the emitter effectively suppressed the degradation of the sensitizer, which is ascribed to fast quenching of the triplet sensitizer by the emitter and justifies the use of ketonic sensitizers for UV-UC in solvents. A theoretical model was developed to acquire insight into the observed temporal decays of the upconverted emission intensity under continuous photoirradiation. The theoretical curves generated by this model fitted the experimental decay curves well, which allowed the reaction rate between the emitter and solvent to be obtained. This rate was also correlated with the difference between the frontier orbital energy levels of the emitter and solvent. Finally, based on the acquired findings, general design guidelines for developing UV-UC samples were proposed.

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^c Nippon Kayaku Co., Ltd., 3-31-12 Shimo, Kita-ku, Tokyo 115-8588, Japan [†] Electronic supplementary information (ESI) available: Photostability of visibleto-visible UC in an ionic liquid, photostability of UV-UC using biacetyl and PPO in DMF, optical absorption spectra of the sensitizer and emitter used in this study, information about the solvents used in this study, experimental setup to controllably induce photodegradation, calculated dipole moments of the sensitizer and emitter, spectral overlap between emitter fluorescence and sensitizer absorption, calculated permittivity dependence of the sensitizer and emitter triplet energies, determination of $\Phi_{\rm UC}$, effect of solvent purity on temporal decay profiles of $\Phi_{\rm UC}$, temporal changes of fluorescence spectra of the sensitizer in the absence of the emitter during photoirradiation in different solvents, photoirradiation-induced changes of optical absorption spectra of samples containing only the sensitizer in different solvents, procedure to calculate $k_{\text{sen,degr}}$, plots of $k_{\text{sen,degr}}$ against ionization energy and electron affinity, procedure to calculate $\varPhi_{\rm sen,rxn},$ photoirradiation-induced changes of optical absorption spectra of samples containing both the sensitizer and emitter in different solvents, temporal changes of fluorescence spectra of the sensitizer in the presence of the emitter during photoirradiation in different solvents, effect of photoirradiation on the triplet lifetime of the emitter, calculation details of our theoretical model, and plots of $k_{\rm emi,rxn}$ against ionization energy and electron affinity. See DOI: 10.1039/d0cp04923a

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Introduction

Photon upconversion (UC) is a technology to convert presently wasted sub-bandgap photons into those with higher energies (*i.e.*, light of shorter wavelength), which are useful in many fields including photovoltaics and photocatalysis. To date, UC using triplet–triplet annihilation (TTA) between organic molecules has been widely explored because of its applicability to low-intensity and non-coherent light.^{1–5} Most of the previous studies focused on visible-to-visible UC.^{1–40} If TTA-UC technology can be reliably extended to the ultraviolet (UV) region (<400 nm), it will become suitable for a broader range of applications, such as for hydrogen generation by water splitting using anatase titanium dioxide (α -TiO₂), which has a band gap of 3.2 eV ($\lambda_{gap} \sim 385$ nm).⁴¹

Since the pioneering studies by Castellano and co-workers^{42,43} and Merkel and Dinnocenzo,⁴⁴ there have been multiple reports^{45–52} exploring UC of visible light to UV light (UV-UC). Here, the principle of TTA-UC is briefly described (Fig. 1a). First, a sensitizer molecule absorbs a low-energy photon (visible



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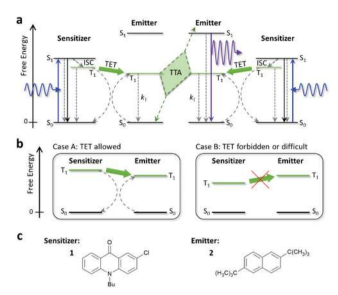


Fig. 1 (a) Schematic diagram of the process of TTA-UC. ISC, TET, and TTA mean intersystem crossing, triplet energy transfer, and triplet-triplet annihilation, respectively. Solid and dashed arrows represent radiative and non-radiative processes, respectively. (b) Schematic depictions of two cases where TET is allowed (Case A) and forbidden or difficult (Case B). (c) Molecular structures of the sensitizer **1** and emitter **2** used in this study.

photon in this context) and transforms to the excited singlet (S_1) state, which immediately converts to the triplet (T_1) state with a certain quantum yield through intersystem crossing. If the energy of the T_1 state of the emitter is similar to or lower than that of the sensitizer, the T_1 energy of the sensitizer can be transferred to the emitter (triplet energy transfer; TET), creating a T_1 emitter (Fig. 1b). When two T_1 emitters interact and undergo TTA, an S_1 emitter can be generated from which an upconverted photon (UV photon in this context) is emitted as delayed fluorescence.

Most previous UV-UC studies were carried out using pyrene or a derivative, whose UC emission maxima range between *ca.* 375 and 425 nm,^{42,45,50} or 2,5-diphenyloxazole (PPO), whose UC emission maxima range between 350 and 400 nm,^{43,44,46,48,49,51} as the emitter. For PPO, 2,3-butanedione (biacetyl) has often been used as the sensitizer.^{43,46,49} As far as we surveyed, except for our previously published open patent documents⁵³ on which this study is based and a recent report using iridium complex sensitizers in water,⁵⁴ both of which were UV-UC to wavelengths shorter than 340 nm, the shortest emission peak wavelength reported for UV-UC is 343–344 nm using terphenyl as the emitter.^{47,50} Therefore, UV-UC with emission maxima shorter than 340 nm has not been well explored thus far.

Shortening emission wavelengths further is meaningful for the following reasons. First, although λ_{gap} of α -TiO₂ is *ca.* 385 nm, which was determined by tangentially extrapolating its absorbance or reflectance spectrum to the horizontal axis,⁵⁵ a general characteristic of semiconductors is that their absorption coefficient is low near λ_{gap} .⁵⁶ For example, sufficient absorption is attained only below *ca.* 350 nm in the case of α -TiO₂ nanoparticles.^{55,57} Second, the quantum efficiency of watersplitting photocatalysts increases with the energy of incident photons.⁵⁸ This present article investigates UV-UC with emission maxima shorter than 340 nm and elucidates the relevant physicochemical characteristics.

However, we have noticed that such UV-UC, whether the samples used in this article or other samples such as those made using biacetyl and/or PPO, is accompanied by non-trivial or sometimes remarkable photodegradation, although such characteristics were not explicitly presented and discussed previously. Only recently, Lee *et al.*⁵⁰ showed fast photodegradation caused by continuous photoirradiation at 455 nm in deaerated tetrahydrofuran (THF) when PPO and terphenyl were used as emitters. They showed that, among the emitters tested, only pyrene exhibited satisfactory photostability in deaerated THF.⁵⁰

Previously, we reported visible-to-visible UC in systems using an ionic liquid as the solvent.^{16,21–23,28} These samples, when properly sealed, exhibited excellent photostability and their lifetime exceeded several years (Section S1 of the ESI†). However, when the same ionic liquid was combined with the sensitizer and emitter used in the present study for UV-UC (Fig. 1c), such photostability was not observed (Fig. S1, ESI† and also below). We also found that the combination of biacetyl and PPO in deaerated dimethylformamide (DMF), which were used previously,^{46,49} showed poor stability under continuous photoirradiation (Fig. S2, ESI†).

Based on these observations, we consider that UV-UC at wavelengths shorter than *ca.* 370 nm tends to suffer from low photostability, presumably because the use of high-energy triplet states may induce photochemical reactions, such as hydrogen abstraction from the solvent. This is an unaddressed issue that should be investigated before UV-UC technology is used in applications. Therefore, it is important to obtain understanding of the governing factors and/or mechanism of such photodegradation in UV-UC.

In this study, based on our previous technological findings regarding UV-UC,⁵³ we develop UV-UC samples that exhibit photoemission peaks in the 320–340 nm range. We find that both the UC quantum efficiency ($\Phi_{\rm UC}$) and photostability of these samples depend on the solvent. To understand this phenomenon, we conduct a systematic investigation by performing both experiments and theoretical analysis. The aim of this article is to elucidate the factors governing such solvent dependence and obtain general guidelines for designing UV-UC systems with high UC efficiency and photostability.

Experimental

We used 10-butyl-2-chloro-9(10*H*)-acridinone (1) and 2,6-di-*tert*butylnaphthalene (2) as the sensitizer and emitter, respectively (Fig. 1c). Both 1 and 2 (purity: >98%) were purchased from TCI; 1 was recrystallized before use and 2 was used as received. We chose 1, in which the photoexcitation is the $n \rightarrow \pi^*$ transition, because the small overlap between the n and π^* orbitals around its carbonyl group leads to a small S₁-T₁ energy gap and the n, π^* state has a high quantum yield of S₁-to-T₁ intersystem crossing ($\Phi_{T,sen}$),⁵⁹ both of which are desirable for sensitizers for TTA-UC.

Table 1 List of samples and selected results

Solvent $\langle E_{\rm T}(30) \rangle^a$ $\Phi_{\rm UC} / \%$ (Exct.							
$(P')^b$	$A_{405\mathrm{nm}}{}^f$	$\Phi_{ m F,sen}$	${\Phi_{\mathrm{T,sen}}}^g$	$\Phi_{ m F,emi}$	intensity/W cm ^{-2})	$k_{ m sen,degr}^{j}/M \ m s^{-1}$	$k_{\rm emi,rxn}^{k}/{\rm s}^{-1}$
Hexane (30.9)	0.067	0.006	0.994	0.33	$4.5(0.40)^h$	7.35×10^{-7}	$5.73 imes10^{-3}$
(0.1) Ethyl acetate $\langle 38.0 \rangle$	0.20	0.274	0.726	0.39	$\begin{array}{c} 8.2 \ (1.75) \\ 1.9 \ (0.20)^h \end{array}$	$\frac{-}{2.35\times10^{-6}}$	$1.92 imes 10^4$
(4.4)	0.20	0.274	0.720	0.39	4.9 (1.74)	2.55 × 10	1.92 × 10
Toluene (33.9)	0.23	0.191	0.809	0.57	$2.2 (0.17)^h$	1.56×10^{-5}	4.47×10^{-1}
(2.4)						_	
Acetonitrile $\langle 45.6 \rangle$	0.15	0.598	0.402	0.43	$0.38 \ (0.50)^h$	4.09×10^{-7}	$3.29 imes10^4$
(5.8) DMF ^c $\langle 43.2 \rangle$	0.17	0.593	0.407	0.49	$0.015 (0.44)^h$	$1.48 imes 10^{-5}$	_
(6.4)							
D-Limonene $\langle N/A \rangle^d$ (N/A)	0.20	0.030	0.970	0.16	$\cong 0 \ (0.15)^h$	1.95×10^{-5}	—
$\left[C_4 \text{dmim}\right]$ [NTf ₂] $\langle 40.9 \rangle^e$ (N/A)	0.15	0.575	0.425	0.58	$0.25 \ (0.44)^i$	1.15×10^{-6}	—
Methanol $\langle 55.4 \rangle$ (5.1)	0.16	0.657	0.343	0.37	$\cong 0 \ (0.52)^h$	—	—

^a From ref. 62 unless otherwise specified. ^b From ref. 63. ^c N,N-Dimethylformamide. ^d Measured using Reichardt's dye but no absorption peak was found in the visible to near-infrared region. ^e Measured using Reichardt's dye. ^f Absorbance of 1 at 405 nm with an optical path length of 1 mm. ^g Determined by assuming the Ermolev's rule $\Phi_{T,sen} = 1 - \Phi_{F,sen}$. ^h Excitation intensity where the T_1 state of 1 is generated at 1.9×10^{-3} M s⁻¹. ⁱ Excitation intensity where the T_1 state of 1 is formed at 1.77×10^{-3} M s⁻¹. ^j Obtained by the procedure described in Section S13 of the ESI. ^k Obtained from the fit to the experimental decay curves of the UC emission intensity using our model; see Fig. 4c.

After testing several acridones, we found that 1 was preferable over the other candidates because of its visible absorption in the 400-425 nm range (Fig. S3, ESI⁺) and ability to undergo TET with naphthalenes. We chose 2 because of its relatively high fluorescence quantum yield and suitable fluorescence spectrum for the purpose of this study.

Samples were prepared using the solvents listed in Table 1. Details of the solvents are given in Table S1 in the ESI.[†] We included p-limonene because it has been reported to prevent degradation of solutes in visible-to-visible UC by functioning as a strong antioxidant that quickly scavenges residual oxygen.⁶⁰ Additionally, in the former half of this study, we included the ionic liquid [C₄dmim][NTf₂] as a reference solvent because it enables highly stable red-to-blue UC16,21-23 (Fig. S1, ESI†). Throughout this report, the concentrations of 1 and 2 were 2×10^{-4} and 2×10^{-3} M, respectively. A continuous-wave laser with an emission wavelength of 405 nm and spot diameter of 0.8 mm was used as the excitation source unless otherwise specified. The absorption spectra of 1 and 2 are depicted in Fig. S3 in the ESI[†] and their fluorescence spectra are shown in Fig. 2a.

Except for the sample with [C₄dmim][NTf₂], all samples, which contained both 1 and 2 or only 1, underwent at least seven (typically eight or nine) freeze-pump-thaw (FPT) cycles using our FPT system to carefully remove dissolved air. Our FPT system consisted of a small glass jar (#33.010007.11A.710, EVAC) with a flange, an O-ring-sealed stainless-steel (SUS) flange coupling to it, and all-SUS vacuum line consisting of a flexible metal hose and Swagelok valves and tube fittings. The vacuum line was connected to an oil-free dry scroll pump (nXDS15i, Edwards) able to attain a vacuum of ca. 1 Pa. To efficiently remove dissolved gas, the volume of liquid in the jar was small (ca. 2 mL). The increase of solute concentration induced by the FPT cycles was negligible for all samples,

as confirmed by their unchanged absorbance in UV-vis measurements. Typically, the emergence of bubbles in the liquid ended within three or four FPT cycles, and thus the aforementioned number of FPT cycles was believed to be sufficient. After the FPT cycles, the glass jar coupled with a closed SUS valve was detached from the vacuum line and transferred into a vacuum-type SUS glovebox containing freshly replaced nitrogen gas (purity: >99.998%). In the glovebox, the liquid sample was injected into a square glass capillary (inner dimensions: 1×1 mm, outer dimensions: 2×2 mm, length: 27 mm) with a closed end. The open top of the capillary was immediately closed with a low-melting-point solder as previously described.^{16,21-23} The seals were checked by placing the capillary under vacuum for a long period (hours or days); an effective seal was confirmed by the sample volume remaining constant. This sealing method works for at least several years (e.g., the sample in Fig. S1, ESI^{\dagger}). For the sample with [C₄dmim][NTf₂], oxygen and moisture were removed by stirring the sample, which had a small volume ($<600 \mu$ L), in an open vial at 60 °C and 200 rpm for 2 h inside a nitrogen-filled glovebox equipped with a gas-purification system (OMNI-LAB, VAC; oxygen and moisture: <1 ppm) before it was sealed in a glass capillary inside the glovebox.

Time-resolved measurements of the UC emission intensity were carried out using nanosecond light pulses generated from an optical parametric oscillator (OPO; NT-242, Ekspla) at 410 nm and 20 Hz. The fluorescence quantum yields of 1 and 2 ($\Phi_{\rm F,sen}$ and $\Phi_{\rm F,emi}$, respectively) and their fluorescence spectra were acquired by an absolute quantum yield spectrometer (Quantaurus-QY, Hamamatsu) using a quartz cell $(1 \times 1 \text{ cm})$; the solute concentration for these measurements (of the order of 10^{-5} to 10^{-4} M) was chosen so that the absorptance of each sample in the integrating sphere was between 0.35 and 0.55 at the excitation wavelength.

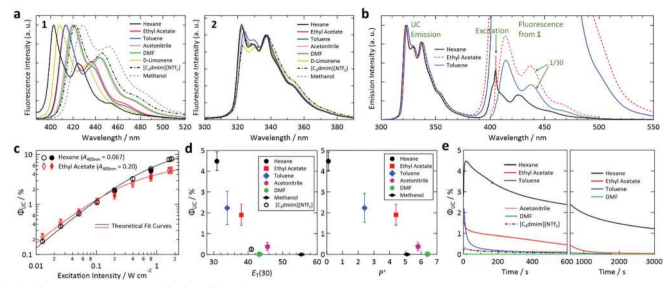


Fig. 2 (a) Fluorescence spectra of **1** (left) and **2** (right) in the solvents used in this study, acquired using an absolute quantum yield spectrometer and degassed dilute solutions (concentrations of the order of $10^{-4}-10^{-5}$ M). (b) Photoemission spectra of the samples prepared with hexane, ethyl acetate, and toluene acquired upon excitation at 405 nm. (c) Dependence of the upconversion (UC) quantum efficiency of the samples prepared using hexane and ethyl acetate on excitation intensity. Open symbols denote data acquired while increasing excitation intensity and filled symbols represent data acquired with decreasing excitation intensity. (d) UC quantum efficiencies plotted against $E_{T}(30)$ (left) and solvent polarity scale P' (right). (e) Temporal profiles of UC quantum efficiency, which is proportional to UC emission intensity, measured for samples prepared using different solvents under continuous photoirradiation at 405 nm. In (b, d and e), the excitation light intensity was chosen such that the irradiation generated the T₁ state of **1** at a rate of 1.9×10^{-3} M s⁻¹ (1.77×10^{-3} M s⁻¹ for [C₄dmim][NTf₂]). See the main text for the details.

In reference experiments, photodegradation was controllably induced in a sample using a setup where the excitation laser beam was expanded to irradiate almost the entire volume of a sample liquid (*ca.* 2 mL) in a hermetically sealed glass vial from below (see Fig. S4 in the ESI† for details). In these experiments, the photoirradiation was continued until each molecule of **1** in the sample turned to the T₁ state 85 times on average. The duration of photoirradiation was set by assuming that the initial absorbance of **1** at 405 nm did not change during the course of the irradiation. All photoemission spectra in this report were corrected by the wavelength-dependent sensitivities of the grating in a monochromator and CCD array detector as described in our previous reports.^{16,21–23}

All quantum-chemical simulations were carried out using Gaussian $16^{(R)}$ at the B3LYP/6-31G++(d,p) level.

Results and discussion

The fluorescence and absorption spectra of sensitizer **1** exhibited large solvatochromic shifts whereas those of emitter **2** showed small solvatochromic shifts (see Fig. 2a for the fluorescence spectra and Fig. S3 in the ESI† for the optical absorption spectra). This behavior is ascribed to the large (negligible) permanent dipole moment of **1** (**2**) (Fig. S5, ESI†). Fig. 2b shows photoemission spectra of samples prepared using hexane, ethyl acetate, and toluene upon excitation at 405 nm. The UC emission spectra were structured with the emission maximum at 322 nm and other peaks in the range of 320–340 nm. The photoemission spectra also contained peaks originating from fluorescence from the S₁ state of **1** in the 400–500 nm range. The intensity of this fluorescence relative to that of the UC emission varied considerably between samples, which is partially attributed to the difference of $\Phi_{\rm F,sen}$ in these solvents ($\Phi_{\rm F,sen}$ = 0.006, 0.274, and 0.191 in hexane, ethyl acetate, and toluene, respectively; *cf.* Table 1). Re-absorption of UC fluorescence from **2** by **1** is weak because of the transmission window of **1** from 310–370 nm (Fig. S6, ESI⁺).

The dependence of $\Phi_{\rm UC}$ of the samples with hexane and ethyl acetate on excitation intensity was determined (Fig. 2c). For $\Phi_{\rm UC}$ in this article, we customarily describe efficiency in percent and thus the maximum is 100%, which is twice the maximum UC quantum yield of 0.5. The emission intensity between 310 and 380 nm was used to calculate $\Phi_{\rm UC}$; *i.e.*, the emission between 380 and 405 nm was not used to exclude the tail of the fluorescence and thermally induced UC emission. The procedure used to determine $\Phi_{\rm UC}$ is described in Section S9 of the ESI.† As shown in Fig. 2c, the samples with hexane and ethyl acetate attained high $\Phi_{\rm UC}$ of 8.2% and 4.9%, respectively, at an excitation intensity of *ca.* 1.75 W cm^{-2} . The data points in Fig. 2c were acquired while first increasing the excitation intensity and then while decreasing the excitation intensity to confirm the reproducibility of the $\Phi_{\rm UC}$ values. Although $\Phi_{\rm UC}$ measured while decreasing the excitation intensity were slightly lower than those obtained with increasing excitation intensity for both samples, the differences were smaller than the related error bars and thus $\Phi_{\rm UC}$ values were considered reproducible.

In Fig. 2c, theoretical curves³¹ were fitted to the data points. From the fitting, excitation threshold intensities (I_{th}) for the samples with hexane and ethyl acetate of *ca.* 1.3 and 0.43 W cm⁻²,

respectively, were obtained. These high $I_{\rm th}$ values and the solvent dependence of $I_{\rm th}$ are explained as follows. First, $I_{\rm th}$ strongly depends on the absorbance of the sample at the excitation wavelength; $I_{\rm th}$ decreases as the absorbance increases and *vice versa*.⁶¹ The samples with hexane and ethyl acetate exhibited low absorbances of 0.067 and 0.20 (corresponding optical absorption coefficients of 0.154 and 0.461 mm⁻¹), respectively, at the fixed excitation wavelength of 405 nm used in this study. Therefore, the high $I_{\rm th}$ here mainly arose from the low absorbance of 1 at the excitation wavelength and the solvent dependence mainly originated from the large solvatochromic shift of 1 (*cf.* Fig. 2a). Besides this factor, the variation of the triplet lifetime ($\tau_{\rm T}$) of 2 in different solvents (*ca.* 114 µs in hexane and 47 µs in ethyl acetate) should also contribute to the solvent dependence of $I_{\rm th}$.

We found that $\Phi_{\rm UC}$ and photostability strongly depended on the solvent. To systematically compare $\Phi_{\rm UC}$ and the rates of photoinduced changes of the samples prepared using different solvents, in the following experiments we set the laser power irradiated onto the sample sealed in a glass capillary (see the Experimental section for details) such that the irradiation generated the T₁ state of 1 at a rate of 1.9×10^{-3} M s⁻¹ in the photoirradiation volume (a cylinder with a diameter of 0.8 mm and length of 1 mm). At this rate, each 1 molecule in the volume transitions to the T₁ state 9.5 times per second. Note that the rate was 1.77×10^{-3} M s⁻¹ in the sample with [C₄dmim][NTf₂]. The actual laser power irradiated onto each sample, which was in the range of 0.73-2.6 mW or 0.15- 0.52 W cm^{-2} , was determined using the absorbance at 405 nm (A_{405nm}) and $\Phi_{T,sen}$ listed in Table 1. The $\Phi_{T,sen}$ values were estimated assuming Ermolev's rule of $\Phi_{\rm T} pprox$ 1 – $\Phi_{\rm F}$.⁵⁹

The determined $\Phi_{\rm UC}$ values are plotted against the polarity scales $E_{\rm T}(30)^{62}$ and P'^{63} in Fig. 2d. We were unable to determine the $E_{\rm T}(30)$ value for D-limonene because it did not exhibit an absorption peak in the visible to near-infrared range in a solution of Reichardt's dye. No P' values for D-limonene or $[C_4 \text{dmim}][\text{NTf}_2]$ were found in the literature. From these plots, we found that $\Phi_{\rm UC}$ is correlated with the solvent polarity and decreases as the polarity increases.

As mentioned above, **1** has a large dipole moment (Fig. S5, ESI†) and thus exhibits a large bathochromic shift as the solvent polarity increases, whereas 2 does not (Fig. 2a). Therefore, as the solvent polarity increases, the T₁ level of **1** is considered to be lowered relative to that of **2**, making TET thermodynamically unfavorable (*i.e.*, Case B in Fig. 1b). The solvent dependence of $\Phi_{\rm UC}$ of our samples is mainly attributed to this mechanism, which is supported by our calculation of the dependence of the triplet energies of **1** and **2** on solvent permittivity (Fig. S7, ESI†). In addition, the solvent dependences of $\Phi_{\rm UC}$.

The stability of the samples under continuous photoirradiation strongly depended on the solvent (Fig. 2e). For example, UC emission in toluene decayed rapidly whereas that in hexane lasted much longer; the reason for this behavior is investigated below. It is noted that no UC emission was observed when p-limonene and methanol were used (Table 1). While the lack of UC emission in methanol can be explained by the above discussion regarding Fig. 2d, the reason for the absence of UC emission in D-limonene is unclear. It may be caused by the high reactivity of D-limonene, which has a reactive unsaturated C==C bond, with high-energy triplet states of 1 and 2.

Here, we note the following three points. First, although the use of solvents with different certified purities resulted in a minor but recognizable effect on the intensity of UC emission, this difference did not alter the qualitative profile of the temporal UC emission intensity change (Fig. S8, ESI[†]). Second, the temporal decays of the UC emission intensity observed in Fig. 2e were not considered to be governed by residual oxygen in the solvents, which was the case in previous visible-to-visible UC studies. $^{64-68}$ This is partly because the use of D-limonene, which scavenged residual oxygen efficiently and helped to attain stable visible-to-visible UC,⁶⁰ completely suppressed the UC emission in the present study. That the UC emission decays observed in Fig. 2e were not caused by residual oxygen was also supported by the thorough FPT treatment and tightly sealed samples used here. Third, the decay rate of the UC emission in hexane in the present study is much slower than that of a previously reported biacetyl/PPO/DMF system46,49 when compared using the similar triplet generation rate on 1 (Fig. S2, ESI†).

In the following investigations, we excluded the sample with methanol because it did not realize UC and the sample with $[C_4 dmim][NT_2]$ because its UC efficiency was low and the photochemical reaction with a molten salt is complex.

To understand the solvent-dependent photostability of our samples, first, we investigated samples containing only **1**. When each sample in a glass capillary was excited at a triplet generation rate of 1.9×10^{-3} M s⁻¹, the decay rate of the fluorescence intensity of **1** depended on the type of solvent (Fig. 3a and Fig. S9 in the ESI† for the fluorescence intensities and spectra, respectively). We confirmed that the photoirradiation induced a decrease of the absorbance of **1** (Fig. 3b and Fig. S10 in the ESI†), indicating that photoirradiation induced degradation of **1**. To compare the degradation rates of **1** in different solvents, we determined the rate of sensitizer degradation $k_{\text{sen,degr}}$ [M s⁻¹] from the decay curves in Fig. 3a by applying a double-exponential fit (see Section S13 of the ESI† for the procedure used to calculate $k_{\text{sen,degr}}$).

Here we use the frontier orbital theory to discuss the observed photoinduced degradation of **1** in the solvents. Generally, excited states of ketones such as the T_1 state of **1** have n, π^* electronic configuration where n and π^* are singly occupied molecular orbitals (SOMOs) and can serve as electronaccepting and -donating orbitals, respectively.⁵⁹ Generally, such SOMOs interact with the highest occupied molecular orbital (HOMO) and lowest unoccupied molecular orbital (LUMO) of an adjacent molecule and create new orbitals into which electrons from both molecules are partially or fully transferred; such a charge transfer generally allows energetic stabilization and may lead to formation of an excited-state complex.⁵⁹ For the n, π^* state of ketones, such intermolecular interaction with a ground-state molecule such as a solvent molecule may cause hydrogen abstraction from the latter because of the half-filled



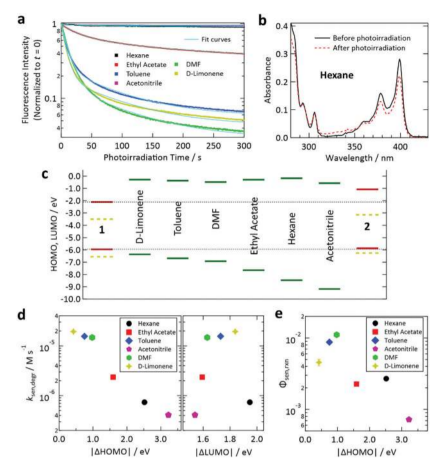


Fig. 3 (a) Temporal decay profiles of the fluorescence intensity of deaerated samples containing only sensitizer 1 under continuous irradiation at 405 nm. The excitation light intensity for each case (*cf.* Table 1) was chosen so that the T_1 state of 1 was generated at a rate of 1.9×10^{-3} M s⁻¹. (b) Decrease of the absorbance of 1 in deaerated hexane (optical path length: 1 mm) induced by photoirradiation at 405 nm using the setup and conditions described in Section S5 of the ESI.† (c) Calculated HOMO and LUMO levels of 1, 2, and the solvents. The SOMO levels of the T_1 states of 1 and 2 are shown as yellow dashed lines. (d) Degradation rates of fluorescence intensities determined from the results in panel (a) plotted against the difference between the HOMO levels of 1 and the solvent (left) and that between the LUMO levels of 1 and the solvent (right). (e) Reaction quantum yield between the T_1 state of 1 and solvent for the samples containing only 1 estimated using the optical absorption changes in Fig. S10 in the ESI.†

orbital on the oxygen atom of the ketone. Hydrogen abstraction by ketones has been widely studied.⁶⁹⁻⁷¹ Two factors are proposed to govern this intermolecular reaction: (i) the energetic proximity of the frontier orbitals of the two interacting molecules and (ii) the constructive spatial overlap of these orbitals.⁵⁹

To study factor (i), we calculated the HOMO and LUMO levels of 1, 2, and the solvents, as depicted in Fig. 3c. In this figure, SOMO levels of the T₁ states of 1 and 2 are also shown. From the relation between $k_{\text{sen,degr}}$ and the energetic separations of the HOMOs and LUMOs between 1 and the solvents (denoted as Δ |HOMO| and Δ |LUMO|, respectively), we found a clear correlation of $k_{\text{sen,degr}}$ with Δ |HOMO|, whereas no obvious correlation was found between $k_{\text{sen,degr}}$ and Δ |LUMO| (Fig. 3d). The same tendency was also observed when the difference between the ionization energies of 1 and the solvents (which physically corresponds to Δ |HOMO|) and that between their electron affinities (which corresponds to Δ |LUMO|) were plotted (Fig. S11, ESI†). These results reveal that the electron transfer from the solvent to 1 is the rate-limiting step of this

photodegradation, which can be interpreted as an electron transfer-initiated hydrogen abstraction process.^{71,72} We also estimated the quantum efficiency of the degradation of the T₁ state of **1** in each solvent ($\Phi_{\text{sen,rxn}}$) from the decease of the absorbance of **1** induced by the controlled photoirradiation (*cf.* Fig. S4, ESI†). The procedure followed to calculate $\Phi_{\text{sen,rxn}}$ is described in Section S15 of the ESI.† Although the scatter of the data points is larger than that in the case of $k_{\text{sen,egr}}$, a similar correlation with Δ |HOMO| was also found for $\Phi_{\text{sen,rxn}}$ (Fig. 3e).

Next, we investigated photoinduced changes of samples containing both 1 and 2. For the sample with hexane, photodegradation of 1 was suppressed by the presence of 2×10^{-3} M of 2, as recognized from the invariance of the optical absorption spectrum of 1 during photoirradiation (Fig. 4a). This suppression is ascribed to prompt TET from 1 to 2 in hexane, which drastically shortens the lifetime of the T₁ state of 1, meaning that 2 strongly suppresses the probability of 1 reacting with the solvent. A similar tendency was also found for the samples with other solvents (Fig. S10 and S12 in the ESI†). However, for the sample with DMF, the decrease in the

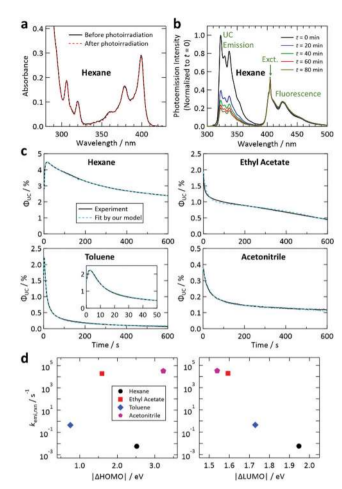


Fig. 4 (a) Demonstration of the suppression of the decrease of the absorbance of **1** by addition of **2** in hexane using the same conditions as in Fig. 3b. (b) Demonstration of the suppression of the decrease of the fluorescence of **1** upon addition of **2** in hexane, which was measured using the same conditions as in Fig. 3a. See also Fig. S9 and S13 in the ESI.† (c) Fit of the temporal decay curves of the UC quantum efficiency shown in Fig. 2e (solid curves) by the theoretical model proposed in this study (dashed curves). (d) Reaction rates between the T₁ state of **2** and the solvents, obtained from the fittings shown in (c), plotted against the difference between the HOMO levels of **2** and the solvent (left) and that between the LUMO levels of **2** and the solvent (right).

absorbance of **1** was not well suppressed (Fig. S12, ESI[†]); this could be because of inefficient TET from **1** to **2** caused by the relatively high polarity of DMF (*cf.* Fig. 2d and 1b). The suppressed photodegradation of **1** in hexane induced by addition of **2** was also evidenced by the invariance of the fluorescence emission intensity of **1** even after 80 min of photoirradiation (Fig. 4b); the similar tendency was also seen for the samples with other solvents (see Fig. S9 and S13 in the ESI[†]).

Our results reveal that by adding an energy-accepting emitter at sufficient concentration (of the order of 10^{-3} M), preferable aspects of ketones as the sensitizer (*cf.* first paragraph of the Experimental section) can be harnessed for UV-UC while effectively suppressing the drawback of using triplet ketones; *i.e.*, the relatively high reactivity of their T₁ state. Considering the viscosities of the solvents employed in this study (*ca.* 0.3–0.6 mPa s at room temperature), the diffusion-controlled rate constant $k_{\rm diff}$ was estimated to be 1–2 \times 10¹⁰ M⁻¹ s⁻¹ using the following equation^{16,59}

$$k_{\rm diff} = \frac{8RT}{3000\eta},\tag{1}$$

where *R*, *T*, and η are the gas constant, temperature, and solvent viscosity, respectively. From the concentration of the energy acceptor 2 (2 × 10⁻³ M) and assuming Case A in Fig. 1b, the lifetime of the T₁ state of 1 was estimated to be only 25–50 ns, which supports the results in Fig. 4a and b.

The rate of the reaction between the T_1 state of 1 and ground state of 2 was considered to be negligible, even though their ground-state HOMO levels are close (Fig. 3c), for the following reasons. First, a bimolecular reaction rate is proportional to the product of the concentrations of the two species involved. In our samples, the concentration of 2 (2×10^{-3} M) was much lower than that of the solvents (7–20 M). Second, the interaction time between the T_1 state of 1 and ground state of 2 should be very short because such an encounter immediately causes an exothermic TET,¹¹ unlike the interaction between the T_1 state of 1 and the solvent, which can last much longer.

We have reached the point to discuss the temporal decay curves of the UC emission intensity under continuous photoirradiation in Fig. 2e. To analyze these decay curves, we developed the theoretical model described below. First, we postulate that the triplet emitter (E^*) becomes a new species (ε) by reacting with a surrounding solvent molecule (sol) at a rate of $k_{\text{emi,rxn}}$ [s⁻¹]. This ε is assumed to quench both E^* and the triplet sensitizer (S^*) at the k_{diff} given by eqn (1). Therefore,

$$E^* + \operatorname{sol} \xrightarrow{k_{\operatorname{emi,rxn}}} \varepsilon (+ \operatorname{sol}')$$
 (2)

$$E^* + \varepsilon \xrightarrow{k_{\text{diff}}} E + \varepsilon^*; \quad S^* + \varepsilon \xrightarrow{k_{\text{diff}}} S + \varepsilon^*.$$
 (3)

Here, *E* and *S* are the ground states of the emitter and sensitizer, respectively, and ε^* is the excited state of ε . In this model, $E \gg E^*$ and $S \gg S^*$ are assumed and the reaction between S^* and solvent is neglected based on the considerations mentioned above. Photoirradiation of the sample was confirmed to shorten the triplet lifetime of 2 ($\tau_{\rm T}$) (Section S18 of the ESI†). Furthermore, it is assumed that ε^* converts into an inactive species ($\varepsilon_{\rm inactive}$) at a quantum yield of $\Phi_{\varepsilon,\rm rxn}$, presumably by reacting with the solvent as follows.

$$\varepsilon^* + \operatorname{sol} \xrightarrow{\phi_{\varepsilon,\operatorname{rxn}}} \varepsilon_{\operatorname{inactive}} (+ \operatorname{sol}'')$$
 (4)

$$\varepsilon^* \xrightarrow{1-\phi_{\varepsilon,\mathrm{rxn}}} \varepsilon.$$
 (5)

We also assume that ε and ε^* are non-emissive, which is evidenced by the invariance of the UC fluorescence spectrum under continuous photoirradiation (*cf.* Fig. 4b).

In addition, we consider initial impurity species in the solvent, Q, which quenches both E^* and S^* . Similar to the case of ε , we introduce the kinetic relations of

$$E^* + Q \xrightarrow{k_{\text{diff}}} E + Q^*; \quad S^* + Q \xrightarrow{k_{\text{diff}}} S + Q^*$$
 (6)

$$Q^* + \text{sol} \xrightarrow{\Phi_{Q,\text{rxn}}} Q_{\text{inactive}} (+ \text{sol}''')$$
 (7)

$$Q^* \xrightarrow{1-\Phi_{Q,\mathrm{rxn}}} Q. \tag{8}$$

We further assume that the second-order rate constant between E^* molecules for the TTA process (k_2) is close to k_{diff} (*i.e.*, $k_2 \approx k_{\text{diff}}$), which was found to be a quantitatively good approximation.³⁶ Although the degradation phenomenon considered here is transient, the timescales of the above-described kinetics are much shorter than those of the change of the UC emission intensity. Therefore, at each instantaneous moment during continuous photoirradiation, the quasi-steady-state approximation is considered to hold well for E^* ,

$$\frac{\mathbf{d}[E^*]}{\mathbf{d}t} \cong 0. \tag{9}$$

Combining all these relations, the proposed model describing the temporal change of UC emission intensity under continuous photoirradiation is obtained as

$$k_{\rm diff}[E^*]^2 + \{k_{\rm T} + k_{\rm diff}([\varepsilon] + [Q])\}[E^*] - \left(\frac{|E|}{[E] + [Q] + [\varepsilon]}\right)\Gamma = 0.$$
(10)

Here, $k_{\rm T}$ is the first-order decay rate of E^* (= $\tau_{\rm T}^{-1}$), which was determined by time-resolved photoemission measurements using light pulses (*cf.* Experimental section). Γ is the generation rate of the T₁ state of the sensitizer, which was 1.9×10^{-3} M s⁻¹. Eqn (10) is a quadratic equation of $[E^*]$ and thus $[E^*]$ at each moment could be expressed using the other parameters in this equation. These parameters were numerically calculated at various times (*t*) after the onset of the photoirradiation at t = 0(see Section S19 of the ESI† for details of the calculation). Because UC emission intensity is proportional to $[E^*]^2$, the theoretical curve of the UC emission intensity for t > 0 can be obtained. Finally, the theoretical curve was computationally fitted to the experimental curve by treating $k_{\rm emi,rxn}$, $\Phi_{c,rxn}$, $\Phi_{Q,rxn}$, and Q as adjustable parameters.

Fig. 4c shows the results of the fittings of this model to the experimental curves for the samples with hexane, ethyl acetate, toluene, and acetonitrile. In all cases, the agreement between the model and experimental curves was good, suggesting that the model has captured the physicochemical characteristics of the present system. All these fittings resulted in $Q \leq 5 \times 10^{-4}$ M (*i.e.*, molar fraction of 0.005% or lower), which does not contradict the certified purities of the solvents. It is noted that processes represented by eqn (4) and (7) are necessary to fit our theoretical model to the experimental curves.

In Fig. 4d, the values of $k_{\rm emi,rxn}$ obtained from the fitting are plotted against $|\Delta$ HOMO| and $|\Delta$ LUMO|. It is noted that the magnitude of $k_{\rm emi,rxn}$, which represents the chemical instability of the triplet emitter, strongly depended on the solvent. In this figure, a correlation was found only for $|\Delta$ LUMO| and $k_{\rm emi,rxn}$. The same tendency was also observed when the difference

between the ionization energies of **2** and the solvents and that between the electron affinities of **2** and the solvents were plotted (Fig. S15, ESI[†]). These results suggest that the process described by eqn (2) is limited by electron transfer from **2** to the solvent; *i.e.*, electron transfer in the opposite direction to that in the reaction between **1** and the solvent discussed above. We did not carry out further detailed investigation of the reaction mechanism because it is beyond the scope of the present study. Nevertheless, the findings acquired from our experimental and theoretical investigations revealed that the photostability of this UV-UC system is controlled by the energetic difference between the relevant frontier orbital levels of the solute (**1** or **2**) and solvent, and that these photodegradation reactions are rate-limited by the electron transfer between molecules.

Conclusions

Using sensitizer **1** and emitter **2**, UV-UC to a shorter wavelength than 340 nm (maximum intensity at 322 nm) was achieved in various solvents. Both $\Phi_{\rm UC}$ and the photostability of **1** under continuous photoirradiation depended on the solvent. The use of hexane yielded the highest $\Phi_{\rm UC}$ of 8.2%, which is close to that of 10.2% reported for UV-UC in the 350–400 nm range achieved using a nanocrystal sensitizer and PPO,⁵¹ and also the highest photostability among the tested solvents. We found that $\Phi_{\rm UC}$ was mainly governed by solvent polarity, which varied the relative T₁ energy-level matching between **1** and **2** because of the solvatochromic shift imposed on **1**. The solvent dependence of $\Phi_{\rm T,sen}$ and $\Phi_{\rm F,emi}$ should also affect $\Phi_{\rm UC}$.

When the samples were prepared without 2, $k_{\text{sen,degr}}$ was large in most of the tested solvents and found to be correlated with Δ |HOMO| between 1 and the solvent. This correlation indicated that the photodegradation of 1 was rate-limited by electron transfer from the solvent to 1 and likely to be an electron transfer-initiated hydrogen abstraction process. However, when the energy acceptor 2, which quenches the T₁ state of 1, was added to the samples, the degradation of 1 was effectively suppressed. This finding justifies the use of a ketonic sensitizer for UV-UC as long as the emitter concentration is higher than the order of 10⁻³ M in non-viscous solvents.

We developed a theoretical model and the curves generated by this model fitted the experimentally acquired temporal decay curves of the UC emission intensity well. This fitting provided several insights into the characteristics of the present UV-UC system. For example, the initial rapid rise of the UC emission intensity for the sample with hexane (*cf.* Fig. 4c) was ascribed to the presence of a trace amount of impurities ($Q \sim 1.9 \times 10^{-7}$ M). Furthermore, $k_{\rm emi,rxn}$ obtained from the fitting was correlated with Δ |LUMO|, which revealed that the photodegradation of 2 was rate-limited by electron transfer to the solvent. These findings indicate that the energetic difference between the frontier orbitals of the solute and solvent is the primary factor determining the photostability. Besides this viewpoint, the frontier orbital theory also addresses the importance of spatial overlap between two frontier orbitals involved in a reaction. Decreasing such overlap by addition of bulky groups to solutes may enhance their photostability.

Overall, this experimental and theoretical study has provided several fundamental insights regarding UV-UC in solvents. As general design guidelines for sample development, one should optimize solvent polarity to maximize $\Phi_{\rm UC}$ and use a combination of solvent and solute whose frontier energy levels are as far apart as possible to enhance solute photostability. These guidelines have not previously been explicitly proposed for UV-UC or visible-to-visible UC. The physicochemical insights obtained from this study will help to establish stable and efficient UV-UC systems in the future.

Conflicts of interest

There are no conflicts to declare.

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References

- 1 S. Baluschev, T. Miteva, V. Yakutkin, G. Nelles, A. Yasuda and G. Wegner, *Phys. Rev. Lett.*, 2006, **97**, 143903.
- 2 T. N. Singh-Rachford and F. N. Castellano, *Coord. Chem. Rev.*, 2010, **254**, 2560–2573.
- 3 A. Monguzzi, R. Tubino, S. Hoseinkhani, M. Campione and F. Meinardi, *Phys. Chem. Chem. Phys.*, 2012, **14**, 4322–4332.
- 4 V. Gray, D. Dzebo, M. Abrahamsson, B. Albinsson and K. Moth-Poulsen, *Phys. Chem. Chem. Phys.*, 2014, 16, 10345–10352.
- 5 T. F. Schulze and T. W. Schmidt, *Energy Environ. Sci.*, 2015, **8**, 103–125.
- 6 P. E. Keivanidis, S. Baluschev, T. Miteva, G. Nelles, U. Scherf, A. Yasuda and G. Wegner, *Adv. Mater.*, 2003, 15, 2095–2098.
- 7 D. V. Kozlov and F. N. Castellano, *Chem. Commun.*, 2004, 2860–2861.
- 8 S. Baluschev, P. E. Keivanidis, G. Wegner, J. Jacob, A. C. Grimsdale, K. Müllen, T. Miteva, A. Yasuda and G. Nelles, *Appl. Phys. Lett.*, 2005, 86, 061904.
- 9 R. R. Islangulov, D. V. Kozlov and F. N. Castellano, *Chem. Commun.*, 2005, 3776–3778.
- S. Baluschev, V. Yakutkin, T. Miteva, G. Wegner, T. Roberts, G. Nelles, A. Yasuda, S. Chernov, S. Aleshchenkov and A. Cheprakov, *New J. Phys.*, 2008, **10**, 013007.
- 11 A. Monguzzi, R. Tubino and F. Meinardi, *Phys. Rev. B:* Condens. Matter Mater. Phys., 2008, 77, 155122.
- 12 T. Miteva, V. Yakutkin, G. Nelles and S. Baluschev, *New J. Phys.*, 2008, **10**, 103002.

- 13 A. Monguzzi, R. Tubino and F. Meinardi, *J. Phys. Chem. A*, 2009, **113**, 1171–1174.
- 14 T. N. Singh-Rachford, J. Lott, C. Weder and F. N. Castellano, J. Am. Chem. Soc., 2009, 131, 12007–12014.
- 15 Y. Y. Cheng, B. Fückel, T. Khoury, R. G. C. R. Clady, M. J. Y. Tayebjee, N. J. Ekins-Daukes, M. J. Crossley and T. W. Schmidt, *J. Phys. Chem. Lett.*, 2010, 1, 1795–1799.
- 16 Y. Murakami, Chem. Phys. Lett., 2011, 516, 56-61.
- 17 A. Monguzzi, M. Frigoli, C. Larpent, R. Tubino and F. Meinardi, *Adv. Funct. Mater.*, 2012, **22**, 139–143.
- 18 Y. Y. Cheng, B. Fückel, R. W. MacQueen, T. Khoury, R. G. C. R. Clady, T. F. Schulze, N. J. Ekins-Daukes, M. J. Crossley, B. Stannowski, K. Lips and T. W. Schmidt, *Energy Environ. Sci.*, 2012, 5, 6953–6959.
- 19 C. E. McCusker and F. N. Castellano, *Chem. Commun.*, 2013, 49, 3537–3539.
- 20 X. Cao, B. Hu and P. Zhang, J. Phys. Chem. Lett., 2013, 4, 2334–2338.
- 21 Y. Murakami, H. Kikuchi and A. Kawai, J. Phys. Chem. B, 2013, 117, 2487-2494.
- 22 Y. Murakami, H. Kikuchi and A. Kawai, J. Phys. Chem. B, 2013, 117, 5180–5187.
- 23 Y. Murakami, T. Ito and A. Kawai, *J. Phys. Chem. B*, 2014, **118**, 14442–14451.
- 24 S. H. C. Askes, A. Bahreman and S. Bonnet, Angew. Chem., Int. Ed., 2014, 53, 1029–1033.
- 25 R. Vadrucci, C. Weder and Y. C. Simon, *Mater. Horiz.*, 2015, 2, 120–124.
- 26 X. Yu, X. Cao, X. Chen, N. Ayres and P. Zhang, *Chem. Commun.*, 2015, **51**, 588–591.
- 27 S. Hisamitsu, N. Yanai and N. Kimizuka, Angew. Chem., Int. Ed., 2015, 54, 11550–11554.
- 28 Y. Murakami, Y. Himuro, T. Ito, R. Morita, K. Niimi and N. Kiyoyanagi, J. Phys. Chem. B, 2016, **120**, 748–755.
- 29 H. Kim, O. S. Kwon, S. Kim, W. Choi and J. H. Kim, *Energy Environ. Sci.*, 2016, **9**, 1063–1073.
- 30 K. Börjesson, P. Rudquist, V. Gray and K. Moth-Poulsen, *Nat. Commun.*, 2016, 7, 12689.
- 31 K. Kamada, Y. Sakagami, T. Mizokuro, Y. Fujiwara, K. Kobayashi, K. Narushima, S. Hirata and M. Vacha, *Mater. Horiz.*, 2017, 4, 83–87.
- J. Han, F. Zhang, J. You, Y. Hiroaki, S. Yamada, T. Morifuji,
 S. Wang and X. Li, *Photochem. Photobiol. Sci.*, 2017, 16, 1384–1390.
- 33 K. Narushima, S. Hirata and M. Vacha, *Nanoscale*, 2017, 9, 10653–10661.
- 34 R. Vadrucci, A. Monguzzi, F. Saenz, B. D. Wilts, Y. C. Simon and C. Weder, *Adv. Mater.*, 2017, 29, 1702992.
- 35 Ł. Bujak, K. Narushima, D. K. Sharma, S. Hirata and M. Vacha, J. Phys. Chem. C, 2017, 121, 25479–25486.
- 36 Y. Murakami, S. K. Das, Y. Himuro and S. Maeda, *Phys. Chem. Chem. Phys.*, 2017, **19**, 30603–30615.
- S. Hisamitsu, N. Yanai, H. Kouno, E. Magome, M. Matsuki,
 T. Yamada, A. Monguzzi and N. Kimizuka, *Phys. Chem. Chem. Phys.*, 2018, 20, 3233–3240.

- 38 A. Abulikemu, Y. Sakagami, C. Heck, K. Kamada, H. Sotome, H. Miyasaka, D. Kuzuhara and H. Yamada, ACS Appl. Mater. Interfaces, 2019, 11, 20812–20819.
- 39 Y. Che, X. Yuan, F. Cai, J. Zhao, X. Zhao, H. Xu and L. Liu, Dyes Pigm., 2019, 171, 107756.
- 40 A. Tokunaga, L. M. Uriarte, K. Mutoh, E. Fron, J. Hofkens, M. Sliwa and J. Abe, *J. Am. Chem. Soc.*, 2019, 141, 17744–17753.
- 41 K. Hashimoto, H. Irie and A. Fujishima, *Jpn. J. Appl. Phys.*, 2005, 55, 8269–8285.
- 42 W. Zhao and F. N. Castellano, J. Phys. Chem. A, 2006, 110, 11440-11445.
- 43 T. N. Singh-Rachford and F. N. Castellano, *J. Phys. Chem. A*, 2009, **113**, 5912–5917.
- 44 P. B. Merkel and J. P. Dinnocenzo, *J. Lumin.*, 2009, **129**, 303–306.
- 45 P. Duan, N. Yanai and N. Kimizuka, *Chem. Commun.*, 2014, 50, 13111–13113.
- 46 M. Majek, U. Faltermeier, B. Dick, R. Pérez-Ruiz and A. J. Wangelin, *Chem. – Eur. J.*, 2015, 21, 15496–15501.
- 47 N. Yanai, M. Kozue, S. Amemori, R. Kabe, C. Adachi and N. Kimizuka, *J. Mater. Chem. C*, 2016, 4, 6447–6451.
- V. Gray, P. Xia, Z. Huang, E. Moses, A. Fast, D. A. Fishman,
 V. I. Vullev, M. Abrahamsson, K. Moth-Poulsen and
 M. L. Tang, *Chem. Sci.*, 2017, 8, 5488–5496.
- 49 M. Barawi, F. Fresno, R. Pérez-Ruiz and V. A. de la Peña O'Shea, ACS Appl. Energy Mater., 2019, 2, 207–211.
- 50 H. L. Lee, M. S. Lee, H. Park, W. S. Han and J. H. Kim, *Korean J. Chem. Eng.*, 2019, 36, 1791–1798.
- 51 S. He, X. Luo, X. Liu, Y. Li and K. Wu, *J. Phys. Chem. Lett.*, 2019, **10**, 5036–5040.
- 52 S. Hisamitsu, J. Miyano, K. Okumura, J. Ka-Ho Hui, N. Yanai and N. Kimizuka, *ChemistryOpen*, 2020, **9**, 14–17.
- 53 Y. Murakami, T. Ito, R. Morita, K. Niimi and N. Kiyoyanagi, worldwide patent application WO2015115556A1, international publication date, Aug 6, 2015, https://patents.google.com/ patent/WO2015115556A1/en; Jpn Pat., JP2014023668, JP6436356B2; US Pat., US20170084758A1, US10175557B2; CN Pat., CN201580006015.1A, CN105940083B.
- 54 B. Pfund, D. M. Steffen, M. R. Schreier, M. S. Bertrams, C. Ye, K. Börjesson, O. S. Wenger and C. Kerzig, *J. Am. Chem. Soc.*, 2020, **142**, 10468.

- 55 R. López and R. Gómez, J. Sol-Gel Sci. Technol., 2012, 61, 1–7.
- 56 M. Fox, *Optical Properties of Solids*, Oxford University Press, Oxford, 2001.
- 57 D. Reyes-Coronado, G. Rodriguez-Gattorno, M. E. Espinosa-Pesqueira, C. Cab, R. de Coss and G. Oskam, *Nanotechnology*, 2008, **19**, 145605.
- 58 H. Kato, K. Asakura and A. Kudo, J. Am. Chem. Soc., 2003, 125, 3082–3089.
- 59 N. J. Turro, V. Ramamurthy and J. C. Scaiano, *Principles of Molecular Photochemistry: An Introduction*, University Science Books, Sausalito, 2009.
- 60 J. Ma, S. Chen, C. Ye, M. Li, T. Liu, X. Wang and Y. Song, *Phys. Chem. Chem. Phys.*, 2019, **21**, 14516–14520.
- 61 A. Haefele, J. Blumhoff, R. S. Khnayzer and F. N. Castellano, J. Phys. Chem. Lett., 2012, 3, 299.
- 62 J. P. Cerón-Carrascoa, D. Jacquemin, C. Laurence, A. Planchat, C. Reichardt and K. Sraïdi, *J. Phys. Org. Chem.*, 2014, 27, 512–518.
- 63 L. Komsta, M. Waksmundzka-Hajnos and J. Sherma, *Thin Layer Chromatography in Drug Analysis*, CRC Press, Boca Raton, 2013.
- 64 F. Marsico, A. Turshatov, R. Peköz, Y. Avlasevich, M. Wagner, K. Weber, D. Donadio, K. Landfester, S. Baluschev and F. R. Wurm, *J. Am. Chem. Soc.*, 2014, 136, 11057-11064.
- 65 C. Mongin, J. H. Golden and F. N. Castellano, ACS Appl. Mater. Interfaces, 2016, 8, 24038–24048.
- 66 D. Dzebo, K. Moth-Poulsen and B. Albinsson, *Photochem. Photobiol. Sci.*, 2017, 16, 1327–1334.
- 67 Q. Liu, M. Xu, T. Yang, B. Tian, X. Zhang and F. Li, ACS Appl. Mater. Interfaces, 2018, 10, 9883–9888.
- 68 S. Wan, J. Lin, H. Su, J. Dai and W. Lu, Chem. Commun., 2018, 54, 3907–3910.
- 69 C. Walling and M. J. Gibian, J. Am. Chem. Soc., 1965, 87, 3361–3364.
- 70 W. J. Leigh, E. C. Lathioor and M. J. St. Pierre, J. Am. Chem. Soc., 1996, 118, 12339–12348.
- 71 E. C. Lathioor and W. J. Leigh, *Photochem. Photobiol.*, 2006, 82, 291–300.
- 72 S. Inbar, H. Linschitz and S. G. Cohen, *J. Am. Chem. Soc.*, 1981, **103**, 1048–1054.