Full-colour solvatochromic fluorescence emitted from a semi-aromatic imide compound based on ESIPT and anion formation†

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A multi-colour solvatochromic fluorophore was developed by substituting a fully organic 3-amino-N-cyclohexyl phthalimide group with a p-toluenesulfonyl (tosyl) group. This compound is essentially colourless in the visible region and exhibits reddish fluorescence through excited-state intramolecular proton transfer (ESIPT) in the crystalline state with a considerably large Stokes shift of 9786 cm⁻¹, as well as bright fluorescence in the full-colour range, from purple to red, when dissolved in 10 different organic solvents. Excitation wavelength measurements, lifetime measurements, and time-dependent density functional theory (TD-DFT) calculations clarified that the full-colour solvatochromism of the fluorescence results from the combination of the fluorescence emissions from the three different excited states: locally excited (N*), intramolecular proton-transferred (T*), and anion states (A*). This unique solvatochromic fluorescence property is expected to be applied in organic electronic devices and biological imaging tools.

1. Introduction

Organic molecules which absorb short-wavelength ultraviolet (UV) light and emit long-wavelength light from violet to the near-infrared (NIR) region with a visible full-colour range are attracting considerable attention for their potential applications in solar spectral converters, fluorescent probes and labels in biology, and emitting materials for organic electronic devices, such as organic light-emitting diodes (OLEDs). 1–3 Of particular interest are fluorescent dyes with luminescent properties that are sensitive to physical changes in the local environment. 4 For instance, solvatochromic phosphors exhibit environmentally sensitive emission properties. 5 This dynamic behaviour is essential for the application of the dyes as imaging tools to provide information on the state of proteins. 6 To tune the fluorescence colour, molecules exhibiting dual or multiple emissions are useful. For instance, dual emission can be obtained one emitter with two emitting states, two independent emitters, or two correlated emitters, and their mechanisms and the electronic or geometrical conditions have been investigated for molecular design and applications. Such materials have been of great interest over the past few years. 7,8

In recent years, aromatic imide/amide molecules have attracted significant attention because of their excellent photothermal and chemical stabilities with rigid structures, high-fluorescence quantum yields, and characteristic electron-transporting properties. 9 Since the lowest energy electronic transition (LET) of phthalimide is a locally excited (LE) n–π* transition derived from a carbonyl group, 10 phthalimide and N-alklyphthalimide exhibit only weak fluorescence with minimal oscillator strengths. N-Alklyphthalimides without substituents on the benzene ring do not exhibit fluorescence; however, those substituted with electron-donating or electron-withdrawing substituents exhibit bright fluorescence because the LET can be changed to an LE π–π* transition with a large oscillator strength. 11 To achieve bright fluorescence, it is also important to suppress the aggregation caused quenching (ACQ) because the formation of aggregates induced by π–π stacking or charge transfer interactions in aromatic molecules frequently reduces the fluorescence quantum yields. The introduction of bulky functional groups 12, 13 and helical structures 14 is known to be effective to suppress or reduce the ACQ.

To achieve high UV absorption followed by full-colour emission with high efficiency, long-wavelength fluorescence with a very large Stokes shift (ν) is a prerequisite. An enhancement of Stokes shift helps to prevent self-absorption of emitted light, which leads to an increase in fluorescence quantum yield. We focused on the use of excited-state intramolecular proton transfer (ESIPT),
a process in which the excited state is relaxed by the structural relaxation (tautomerisation) from the normal (N*) form to the tautomer (T*) form through proton transfer in the excited state, and it can be used for white light emission and sensing probes. There are two types of ESIPT: hydroxyl (–OH) and amino (–NH) types, in which the proton donors are O and N, respectively. The >NH type has been found to undergo ESIPT more efficiently than the >OH type; it has a great advantage in that it can be substituted with various groups, such as –NH–R. In addition, ESIPT is sensitive to the ambient environment and can be easily tuned by changing the electron-donating or electron-withdrawing substituents. Thus far, by changing the chemical structures with adjusted electron-withdrawing substituents and extending the π-conjugated framework, various types of fluorophores that emit light covering the full-colour range from blue to the NIR region have been achieved. To achieve full-colour solvatochromic photoluminescence, blue, green, and red colour emissions (light’s three primary colours) are essential. Especially, red fluorescence is difficult to achieve by excitation in the UV region due to the very larger Stokes shift. Okamoto et al. reported the fluorescent properties of 3-aminophthalimides substituted with electron-donating and electron-withdrawing substituents. This article is licensed under a Creative Commons Attribution-NonCommercial 3.0 Unported Licence.

2. Experimental

2.1. Materials

The compound, 3-nitrophthalic anhydride (1), purchased from Kanto Chemical Co., Inc., was used as received. Cyclohexylamine, Pd/C (55% water), and p-toluenesulfonyl chloride, purchased from Tokyo Kasei Co., Ltd., were used as received.

2.2. Synthesis

The synthesis of 3-tosylamino-N-cyclohexylphthalimide (3TsAPI) is shown in Scheme 1.

Synthesis of 3-tosylamino-N-cyclohexylphthalimide (3TsAPI). Compound 2 (4.13 g, 15 mmol) was dissolved in a mixture of ethanol (50 mL) and ethyl acetate (95 mL), followed by the addition of 10% palladium on carbon (Pd/C, wetted with ca. 55% water) (0.917 g). This mixture was subsequently stirred at 20 °C under a hydrogen atmosphere for 1 day. The reaction solution was passed through Celite 535 to remove Pd/C, and the filtrate was concentrated using a rotary evaporator. The resulting solid was recrystallised from ethanol, and vivid yellow plate-like crystals were obtained (yield, 76%). For 1H NMR (400 MHz, CDCl3): δ [ppm] 1.24–1.41 (3H, br m, CH3), 1.66–1.73 (3H, br m, CH2), 1.84–1.87 (2H, br m, CH2), 2.13–2.23 (2H, br m, CH2), 4.04 (1H, m, N-CH), 5.22 (2H, br s, NH), 6.82 (1H, d, J = 8.4 Hz, aryl), 7.10 (1H, d, J = 7.6 Hz, aryl), 7.38 (1H, t, J = 15.6 Hz, aryl) (Fig. S1, ESI†).

Synthesis of 3-tosylamino-N-cyclohexylphthalimide (3TsAPI). Compound 3 (0.740 g, 3.02 mmol) and p-toluenesulfonyl chloride (1.93 g, 10 mmol) were dissolved in pyridine (1.0 mL) and heated at 80 °C for 1.5 h under a nitrogen atmosphere. After cooling to 20 °C, the white solid reprecipitated with excess ethanol was filtered and dried. White needle-like crystals were obtained by recrystallisation from ethanol (yield, 82%). For 1H NMR (400 MHz, CDCl3): δ [ppm] 1.25–1.39 (3H, br m, CH3), 1.72–1.73 (3H, br m, CH2), 1.86–1.87 (2H, br m, CH2), 2.15–2.25 (2H, br m, CH2), 4.15 (1H, m, N-CH), 7.90 (1H, t, J = 16.0 Hz, aryl), 8.08 (2H, t, J = 14.8 Hz, aryl) (Fig. S1, ESI†).
Synthetic procedures for other N-cyclohexylphthalimide compounds with different EWGs and those $^1$H NMR spectra are presented in ESI‡ (Fig. S4–S6).

2.3. Measurements

2.3.1. UV-Vis absorption and excitation/emission spectroscopy.

The concentrations of 3TsAPI in the organic solvents were set to 2.5 × 10^{-4} mol L^{-1}. The solvents, toluene (99.7%, Kanto Chemical Co., Inc. (Kanto), fluorescence grade (FG)), acetone (99.7%, Kanto, FG), ethanol (EtOH, 99.5%, FUJIFILM Wako Pure Chemical Corp. (Wako), FG), methanol (MeOH, 99.9%, Wako, FG), dimethyl sulfoxide (DMSO, 99.7%, Kanto, FG), ethyl acetate (EtOAc, 99.7%, Kanto, FG), acetonitrile (MeCN, 99.8%, Wako, FG), ethylene glycol (EG, 1,2-ethanediol, 99.5%, Kanto, FG), chloroform (CHCl₃, 99.0%, Tokyo Kasei Co., Ltd.) were used without further purification. The UV-vis absorption and PL excitation/emission spectra were recorded separately at 20 °C using a JASCO V-760 spectro-photometer (JASCO Co., Tokyo Japan) and a Hitachi F-7100 fluorescence spectrometer (Hitachi High-Technologies Co., Tokyo, Japan) equipped with an R928 photomultiplier tube (Hamamatsu Photonics Co., Japan), respectively. The front-face method was adopted for the film samples to reduce the self-absorption of the emitted luminescence.

The PL quantum yields were measured using a calibrated integrating sphere (C9920, Hamamatsu) connected to a multi-channel analyser (C7473, Hamamatsu) using an optical fibre link.

2.3.2. Time-resolved luminescence measurements.

Fluorescence lifetime measurements with a low time resolution of 1 ns were conducted using a fluorescence lifetime measurement system (Quantaurus-Tau, C11367-03, Hamamatsu Photonics, Japan) at 20 °C. The decay component was recorded using excitation by applying a flashing light-emitting diode (LED) light at wavelengths of 340, 365, or 405 nm. The fluorescence decay curves were accumulated until the peak intensity reached 1000 or 5000. The emission decay was well-fitted using one to three exponential functions. The average lifetime was calculated as 

\[ \tau = \frac{\sum_{i=1}^{n} A_i \tau_i}{\sum_{i=1}^{n} A_i} \]

where \( A_i \) is the pre-exponential for lifetime \( \tau_i \).

2.3.3. Other measurements. $^1$H NMR spectra were measured using a JEOl AL-400 spectrometer operating at a $^1$H resonance frequency of 400 MHz. The chemical shifts were calibrated in ppm (δH) using tetramethylsilane (TMS) as the standard (0 ppm).

2.4. Quantum chemical calculations

Density-functional theory (DFT) calculations were conducted using the Gaussian-16 software (Rev.C.01),\textsuperscript{27} as described in our previous studies.\textsuperscript{28,29} Geometry optimisation was independently performed with the B3LYP and CAM-B3LYP functionals with the 6-311G(d) basis set for the ground (S₀) and excited (S₁) states, respectively. The 6-311++G(d,p) basis set was used to calculate the vertical excitation wavelengths and oscillator strengths (f) for each of the S₀ and S₁ geometries. The S₀ → S₁ transition at the optimised S₀ geometry corresponds to the optical absorption of the ground state. Contrarily, that at the optimised S₁ geometry corresponds to the fluorescent emission according to Kasha’s rule. Each calculated transition was replaced by a Gaussian broadening function with a width of 0.10 eV, producing the shapes of the calculated spectra. The solvent effects in the ground and excited states were incorporated based on the polarisable continuum model (PCM) implemented in the Gaussian software.

2.5. Crystallography

A single crystal suitable for X-ray analysis was mounted on a fibre loop. Diffraction experiments were performed on a Rigaku Saturn CCD area detector with graphite-monochromated Mo-Kα radiation (λ = 0.71073 Å). Intensity data (6° < 2θ < 55°) were corrected for the Lorentz-polarisation effects and absorption. Structure solution and refinements were carried out using the CrystalStructure (2000–2018) program package (Rigaku Corp., Tokyo Japan). The heavy-atom positions were determined by a direct methods program (SIR92), and the remaining non-hydrogen atoms were found by subsequent Fourier syntheses and refined by full-matrix least-squares techniques against F² using the SHELXL-2014/7 programme.\textsuperscript{30} The position of the NH hydrogen atom was refined, whereas the remaining hydrogen atoms were included in the refinements using a riding model. Crystal data: C₉₃H₆₃N₄O₁₅S, M = 398.48, triclinic, a = 5.3459(13), b = 13.292(3), c = 14.387(3) Å, \( \alpha = 70.586(8) \), \( \beta = 81.639(9) \), \( \gamma = 84.599(10) \), \( \mu = 95.28(4) \) Å, \( T = 93 \) K, space group P\( \overline{1} \) (no. 2), Z = 2, 11 858 reflections measured, 4336 unique (\( R_{int} = 0.0623 \)), which were used in all calculations. The final wR(F²) was 0.1176 (all data). CCDC 2067299 contains the supplementary crystallographic data for this study."
The optimised conformations of 3TsAPI in the N and N* forms calculated under vacuum conditions were compared with those in the crystal lattice in Fig. S8(a)–(c) (ESI†). Moreover, the calculated absorption spectra of the N form and the fluorescence spectra of the N* form of 3TsAPI in various solvents are shown in Fig. S9 and S10 (ESI†), respectively. Intriguingly, the experimentally determined crystal structure of 3TsAPI is quite similar to the optimised N and N* structures, and the conformational change caused by one-electron excitation from N to N* is quite small, which leads to a small $\nu$ value of N* emission (5322 cm$^{-1}$ in toluene). In addition, the solvent-dependence of the N absorption and N* fluorescence peaks is also insignificant owing to their less polarisable electronic structures. Fig. S11 (ESI†) shows the spatial distributions of the HOMOs and LUMOs in the optimized geometries for the ground state (N), the excited state (N*), the excited proton transferred state (T*), and the excited anionic state (A*) of 3TsAPI. The very similar electronic structures of the N and N* states support the characters of the calculated absorption and emission spectra.

In contrast, the optimised conformations of the T*, A, and A* forms are quite different from those of N and N*, as displayed in Fig. S8(d)–(f) (ESI†). The dihedral angles of C5–N2–S1–C15 are almost 180°, and the benzene ring of the tosyl group in the T* form is located orthogonally to the phthalimide plane, whereas it is located in the same plane in the A form. The large conformational change in T* significantly enhances the Stokes shift, whereas the limited change in A* reduces it. For instance, the calculated $\nu$ value for the emission from T* in toluene (13 066 cm$^{-1}$) is 3.4 times larger than that from A* (3816 cm$^{-1}$). However, as seen in Fig. S11 (ESI†), it is intriguing to note that the electronic structures of T* and A* look very similar to each other, but totally different from those of N and N*. The HOMOs and LUMOs are mainly localised at the phthalimide moiety, and both the excited states can emit bright fluorescence at longer wavelengths via LE($\pi^*-\pi$) transitions once T* or A* are generated.

The calculated T* spectra of 3TsAPI in various organic solvents are shown in Fig. S12 (ESI†). The shortest and longest fluorescences were predicted to appear at 572 nm in MeOH/MeCN and 608 nm in cyclohexane, respectively, owing to the dielectric effects (effect of the polarity of solvents) on the ESIPT process. The predicted order of the T* emission agrees well with the relative permittivity values of the solvents adopted in the Gaussian software. It is well known that the T* fluorescence shows a bathochromic shift in less polar solvents, such as toluene, and a hypsochromic shift in more polar solvents, such as EtOH.$^5$

Moreover, the calculated absorption and emission spectra of A and A* of 3TsAPI in various solvents are shown in Fig. S13 and S14 (ESI†), respectively. The solvent effects and the predicted order of the solutions are similar to those of the calculated T* emission, which indicates that the anion absorptions and emissions also reflect the dielectric effects of the solvents. Fig. S15 (ESI†) shows the relationships between the relative permittivity ($\varepsilon_r$) and the calculated deprotonation energy of 3TsAPI under vacuum and in various solvents. The inverse proportion indicates that the anion of 3TsAPI can be formed readily in polar solvents that are between acetone ($\varepsilon_r = 20.49$) and DMSO (46.83) in the figure. Notably, the optimised conformation of the A form gradually changes depending on the polarity of the solvent as shown in Fig. S16 (ESI†). The coplanar conformation between the
3.3 Optical properties of 3TsAPI crystal

The crystalline structure of 3TsAPI, analysed by single-crystal X-ray diffraction, is shown in Fig. 2. The needle-shaped crystals obtained from the ethanol solution were used for the measurement. The sulfonamide N2 atom lies in the phthalimide plane in close contact with the imide carbonyl oxygen O1 atom (2.849(2) Å), indicating a strong intramolecular hydrogen bond in the quasi-6-membered ring. Meanwhile, the S1 atom deviated slightly from the phthalimide plane (0.525(2) Å), and the two benzene rings are orthogonal to each other with an interplane angle of 85.17(7)°. Thus, the skeletal structure of 3TsAPI has a bent ‘L-shape’ in the crystal.

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3.3 Optical properties of 3TsAPI crystal

Fig. 3 shows the excitation and emission spectra of 3TsAPI in the crystalline state at 20 °C. The excitation and emission peaks are observed at 370 nm and 580 nm, respectively. Therefore, the crystal appears colourless under white light and exhibits bright orange fluorescence under UV irradiation at 365 nm. Based on the large Stokes shift ($\nu = 9786 \text{ cm}^{-1}$) and the considerably short fluorescence lifetime ($\tau = 1.1 \text{ ns}$, Fig. S17, ESI†), the observed emission is attributable to the T* fluorescence (Fig. 1(a) and (c)) via ESPT. Moreover, a high fluorescence quantum yield ($\Phi = 0.22$) was observed despite the large Stokes shift. We have also synthesized N-cyclohexylphthalimide compounds substituted at the 3-position with –NH$_2$, –NHCOCH$_3$, –NHCOCF$_3$, –NHSO$_2$CH$_3$ groups (the synthesis procedures are shown in the ESI† as stated in Section 2.2), and their experimental excitation and emission spectra in the crystalline state are shown in Fig. S18 (ESI†). In addition, the excitation and emission wavelengths ($\lambda_{\text{ex}}$, $\lambda_{\text{em}}$) and quantum yield ($\Phi$) are summarised in Table S1 (ESI†). As predicted by the TD-DFT calculations, the compounds substituted with –NH$_2$ and –NHCOCH$_3$ show green fluorescence at 503 nm and blue fluorescence at 456 nm, respectively, without ESPT emission, while the compound with –NHCOCF$_3$ shows yellow fluorescence at 542 nm and the one with –NHSO$_2$CH$_3$ demonstrates red fluorescence at 585 nm. These results indicate that the emission colour can be well tuned by the selection of EWGs. It should be noted that the $\Phi$ value of 3TsAPI is 2.75 times as high as another red-emitting compound with –NHSO$_2$CH$_3$. This should be originated from the bulky tosyl and cyclohexyl groups that reduce intermolecular aggregation and suppress ACQ. Accordingly, 3TsAPI could be the best compound exhibiting red fluorescence via ESPT with a large Stokes shift and a high quantum yield.

3.4 Optical properties of 3TsAPI in solutions

Fig. 4 shows the UV-vis absorption spectra of 3TsAPI dissolved in 10 different organic solvents with a wide polarity range. Strong absorption peaks are observed at around 340 nm for all solvents (Table 1). This wavelength correlates with the calculated highest occupied molecular orbital (HOMO)–lowest unoccupied molecular orbital (LUMO) transition at 338 nm (Fig. 1(a) and (c)). As depicted in Fig. S11 (ESI†), the N* form mainly contributes to the locally excited state (N*) generated by one-electron excitation from the HOMO to the LUMO. In addition, relatively weak absorption peaks were observed between 390 and 480 nm in solvents with proton-accepting abilities, such as DMSO, MeOH, and EtOH. Fig. 5 shows photographs of these solutions under white light. The nine solutions, except for DMSO, are almost colourless and transparent, although the DMSO solution shows a pale yellow colour, which is caused by the broad absorption at 419 nm. Since the calculated absorption of the anion in DMSO at 429 nm (Fig. 1(b) and (d)) is very close to this wavelength, these weak absorption bands may originate from the partly proton-dissociated anionic form. As predicted by the calculation, deprotonation is likely to occur in solvents that are more polar than THF. The significant dependence of the optimised conformations of the anionic state on the polarity of the solvents may account for the characteristic solvatochromism in the absorption spectra.

Excitation/emission spectra of 3TsAPI in organic solvents. Fig. 6 shows the excitation and emission spectra of 3TsAPI in 10 different organic solvents, and Fig. 5 shows the CIE coordinates estimated from the emission spectra and photographs of these
Curves in the solvents are presented in ESI†.

Table 1

| Solvent     | \(\lambda_{\text{ex}}/\text{nm}\) | \(\lambda_{\text{em}}/\text{nm}\) | \(\nu/\text{cm}^{-1}\) | \(\tau/\text{ns}\) | \(\Phi\)  |
|-------------|-------------------------------|-------------------------------|----------------|----------------|-------|
| Toluene     | 342                           | 346                           | 602            | 12 288         | 0.4   |
|             | 361                           | 409                           | 3263           | 4.8            | 0.07  |
| Acetone     | 337                           | 351                           | 599            | 11 796         | 0.2   |
|             | 351                           | 420                           | 4681           | 1.6            | 0.01  |
| EtOH        | 339                           | 395                           | 507            | 5585           | 7.1   |
|             | 400                           | 496                           | 4822           | 11.2           | 0.19  |
| MeOH        | 337, 400                      | 392                           | 508            | 5825           | 9.7   |
|             | 419                           | 528                           | 4927           | 5.6            | 0.21  |
| DMSO        | 339                           | 350                           | 600            | 11 905         | 0.1   |
|             | 400                           | 490                           | 4592           | 12.6           | 0.31  |
| EtOAc       | 338                           | 350                           | 600            | 11 905         | 0.1   |
|             | 400                           | 490                           | 4592           | 12.6           | 0.31  |
| MeCN        | 337                           | 345                           | 597            | 12 235         | 0.1   |
|             | 395                           | 493                           | 5049           | 7.7            | 0.14  |
| EG          | 337                           | 355                           | 592            | 11 277         | 0.9   |
|             | 355                           | 427                           | 4750           | 0.4            |       |
| CHCl₃       | 341                           | 350                           | 601            | 11 927         | 0.3   |
| Cyclohex    | 341                           | 346                           | 598            | 12 168         | 0.3   |

Stokes shifts (\(\nu = 11 277–12 288 \text{ cm}^{-1}\)) and the considerably short lifetimes (\(\tau < 1.0 \text{ ns}\), Table 1). The observed emission wavelengths adequately reflect the calculated value for T* fluorescence. Conversely, the short-wavelength fluorescence observed at approximately 400 nm in toluene, acetone and EG can be attributed to the N* fluorescence due to the small Stokes shifts (\(\nu = 3263–4750 \text{ cm}^{-1}\)) and the moderately short fluorescence lifetimes (1.0 < \(\tau < 5.0 \text{ ns}\), Table 1). These wavelengths are well predicted by the calculated N* fluorescence (Fig. 1(a), (c) and Fig. S10, ESI†). The differences in the intensity ratio between T* and N* fluorescence can be explained by the solvent dependence of the energy barriers for the tautomerisation between N* and T*.

For the solvents, except for EtOH, MeOH, and DMSO, long-wavelength fluorescence is observed at approximately 600 nm when excited at approximately 350 nm. This is attributable to the T* fluorescence via ESIPT because of the considerably large solutions under UV irradiation at 365 nm (UV on). In addition, the excitation and emission wavelengths (\(\lambda_{\text{ex}}, \lambda_{\text{em}}\)), Stokes shifts (\(\nu\)), fluorescence lifetimes (\(\tau\)), and quantum yields (\(\Phi\)) are summarised in Table 1. The differences in the intensity ratio between T* and N* fluorescence can be explained by the solvent dependence of the energy barriers for the tautomerisation between N* and T*.32

In addition to the fact that the T* fluorescence via ESIPT was not observed in EtOH, MeOH, and DMSO, other fluorescence bands with small Stokes shifts (4592 \text{ cm}^{-1} < \nu < 5825 \text{ cm}^{-1}) were observed between 490 nm and 528 nm in acetone, EtOAc, and MeCN when excited at approximately 400 nm (392 nm < \(\lambda_{\text{ex}} < 419 \text{ nm}\)). Moreover, their excitation wavelengths, \(\lambda_{\text{ex}}\), were longer than those of N* (345 nm < \(\lambda_{\text{ex}} < 361 \text{ nm}\)), and the fluorescence lifetimes (\(\tau\)) were significantly elongated up to > 5.0 ns compared with those of N* and T* (\(\tau < 5.0 \text{ ns}\)), which are quite different from the N* and T* emissions.

To investigate the origin of the fluorescence characterised by long excitation wavelengths and long lifetimes, a small portion of trifluoroacetic acid (TFA) was added to the solutions. Fig. 7 shows the emission spectra of 3TsAPI in acetone and EtOH before and after the addition of TFA, with their photographs under UV irradiation. In addition, the excitation and emission...
wavelengths ($\lambda_{\text{ex}}$, $\lambda_{\text{em}}$), Stokes shifts ($\nu$), fluorescence lifetimes ($\tau$), and quantum yields ($\Phi$) are summarised in Table S2 (ESI†).

As shown in Fig. 7(a), the addition of TFA to the acetone solution effectively suppresses the fluorescence at 496 nm with a long $\tau$ (11.9 ns), and the fluorescence from T* via ES IPT at 605 nm increases when excited at 351 nm (Fig. S20, ESI†). The fluorescence colour under UV irradiation immediately changed from light blue to orange upon the addition of TFA. As shown in Fig. 7(b), the addition of TFA effectively suppressed the fluorescence at 507 nm in EtOH with a long $\tau$ (7.1 ns), and subsequently, the fluorescence from T* at 601 nm increased upon excitation at 350 nm. The fluorescence colour changed from green to orange. These changes in the emission spectra and lifetimes clearly indicate that the original state of 3TsAPI in these solutions included the anionic form, implying that deprotonation occurred in the ground state. The experimental $\lambda_{\text{ex}}$ and $\lambda_{\text{em}}$ values correspond well with the calculated values for the anionic forms, as shown in Fig. 1(b) and (d) ($\lambda_{\text{ex}} = 429$ nm, $\lambda_{\text{em}} = 513$ nm in DMSO). The lifetimes of the fluorescence at short-wavelengths of 414 nm in acetone/TFA and 400 nm in EtOH/TFA are very short (1.7 ns and 0.4 ns, respectively), and the emission wavelengths correlate with the calculated wavelengths for the N*. Only A* exhibited fluorescence in EtOH, MeOH, and DMSO without T* fluorescence because the tosyl-substituted amide group of 3TsAPI forms strong hydrogen bonds with these proton-accepting solvents, as shown in Fig. S21 (ESI†), thus suppressing the ES IPT process.31,33–35 As shown in Fig. 4, these solutions demonstrated partial absorption of the A form in the visible region, also indicating that 3TsAPI forms anions in solvents with a strong proton-accepting ability.

Finally, the origin of the full-colour fluorescence represented in the CIE coordinates and the photographs of the 3TsAPI solutions (Fig. 5) is discussed. In the acetone solution, the fluorescence bands of the anionic form and T* were observed in addition to the N* fluorescence, and the combination of these emissions led to a light blue colour. Although only the anion fluorescence was observed in EtOH, MeOH, and DMSO, different fluorescence colours were achieved, ranging from green to yellow. Considering the order of the relative permittivity ($\varepsilon_r$ dielectric constants) of these solvents: DMSO ($\varepsilon_r = 46.8$) > MeOH (32.6) > EtOH (24.9) (Fig. S15, ESI†), the excited state of the anionic forms is more stable in high $\varepsilon_r$ solvents, which results in long-wavelength emissions resulting in different colours. As shown in Table 1, anion fluorescence was observed in polar solvents of EtOAc, acetone, EtOH, MeOH, MeCN and DMSO, which coincides well with the order of $\varepsilon_r$ in Fig. S15 (ESI†).
except EG. In the EtOAc and MeCN solutions, T* fluorescence was observed in addition to the anion fluorescence, after which white and pale yellow fluorescence colours were observed. In particular, due to the good balance between the relative fluorescence intensities of the anionic and T* forms, the emission spectrum covered the entire visible region, which endowed the EtOAc solution with white fluorescence (Fig. S22, ESI† λex = 360 nm). In the EG, CHCl3, and Cyclohex solutions, the intense T* fluorescence led to the characteristic orange-red fluorescence because the anionic form is destabilised in non-polar solvents with small ε values (4.7 for CHCl3 and 2.0 for Cyclohex) (Fig. S15, ESI†). However, the reason for the orange fluorescence in EG with a high ε of 40.2 is unclear at present.

In summary, the novel imide compound, 3TsAPI, exhibits both small and large Stokes shifted fluorescence derived from N* and T*, as well as another fluorescence derived from the anionic form. These three types of fluorescence compete through interactions in various solvents, resulting in multi-colour fluorescence in the entire visible region. The compound, 3TsAPI, is an extraordinary fluorophore that exhibits bright fluorescence in the full-colour range, from purple to red, owing to the exquisite balance of the three fluorescence species generated in various organic solvents.

4. Conclusions

We developed a new fluorophore that absorbs only UV light and emits fluorescence in the long-wavelength region of visible light. The TD-DFT calculations predicted that an imide compound with an –R–N–H…O=–C= type intramolecular hydrogen bond substituted with a strong electron-withdrawing tosyl group (3TsAPI) would absorb only UV light and exhibit reddish fluorescence through ESIPT. This compound exhibited a high quantum yield and a considerably large Stokes shift (9786 cm−1) in the crystalline state. The high quantum yield originated from the suppression of ACQ, resulting from the bulky tosyl and cyclohexyl groups. This compound shows an extremely unique solvatochromic fluorescence that exhibits a full-colour range from purple to red in 10 different organic solvents. Excitation wavelength measurements, lifetime measurements, and TD-DFT calculations clarified that the full-colour fluorescence results from the combination of the fluorescence emitted from the three states, namely N*, T*, and A*. The significant colour change caused by solvatochromic fluorescence and the characteristic dependence on the excitation wavelength is caused by the different stabilities and subtle balance among the states of N*, T*, and A*, arising from the interactions between the molecules and various solvents with different polarities. These properties have great potential for application in organic electronic devices, such as OLEDs and imaging tools in biology.

Conflicts of interest

There are no conflicts to declare.

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