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Click and shift: the effect of triazole on solvatochromic dyes†

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Solvatochromic dyes are prime candidates for exploring complex polarity-dependent biological processes. The design of novel dyes for these applications typically presents long synthetic routines. Herein, we report a concise synthesis of azido-functionalised push–pull fluorenes, belonging to the most potent groups of fluoro-solvatochromic compounds. These fluorenes can be attached to multiple alkynes *via* the highly versatile copper-catalysed azide–alkyne cycloaddition (CuAAC). Our study focuses on the beneficial effect of the triazole, formed by CuAAC, comparing the simple push–pull fluorene FR1 with the novel model compound FR1TP. While the triazole in FR1TP is not conjugated to the π -system of the dye, the heterocycle surprisingly produces a bathochromic emission shift of around 50 nm. This effect, caused by triazole-induced LUMO stabilisation, was also observed for two other model compounds: FR2TP and FR1TM. All compounds display photophysical properties that are highly desired for *in vivo* imaging dyes, including remarkable two-photon absorption properties. The experimental results are supported by theoretical calculations. We anticipate that our findings will enable synthesis of new high-performance fluorosolvatochromic dyes for diverse applications.

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Introduction

Modulating the optical properties of fluorescent materials is key to the design of new sensing or imaging technologies.^{1–5} Prime examples of such materials are solvatochromic dyes, which are characterized by displaying pronounced changes in their photophysical properties with solvent polarity. This distinctive feature has been pivotal to their use in bioimaging to study protein aggregation,⁶ proteome stress,⁷ lipid organisation of cell membranes,⁸ and detection of pathogenic bacteria, among others.^{9,10} These advanced applications are based on environment-dependent biophysical parameters and require solvatochromic fluorophores suitable for imaging. Desired properties include (i) excitation in the visible region, (ii) high fluorescence quantum yields, (iii) good photostability, and (iv) high two-photon cross-sections (for multiphoton microscopy). Since the development of **PRODAN** by Weber and Farris in 1979,¹¹ several derivatives with all these properties have been developed.²

Fluorene-based dyes, such as **FR0**, developed by Klymchenko and coworkers are push–pull systems with notable solvatochromic properties, often comparable to or even exceeding those of

PRODAN-like dyes (Fig. 1).^{12,13} In this nomenclature, “FR” denotes “fluorene,” while the following number indicates the number of carbon atoms attached to the carbonyl group; a “0” specifies an aldehyde substituent. The behaviour of these push–pull fluorenes has been investigated by varying the nature of the acceptor groups and donor amines.¹⁴ The trends observed in their photophysical properties (solvatochromism, twisted internal charge-transfer, fluorescence in aqueous environments *etc.*) align with those observed for other classical push–pull systems.^{15–17}

However, unlike **PRODAN** and similar commercially marketed products (*e.g.*, solvatochromic pyrene derivatives¹⁸), these compounds remain underexplored, not merely due to the complexity of their syntheses (*i.e.* five steps for the aldehyde version of **FR0** and seven steps for the keto version of **FR8**; see below for structures),¹² but also that the synthetic routes lack flexibility. This

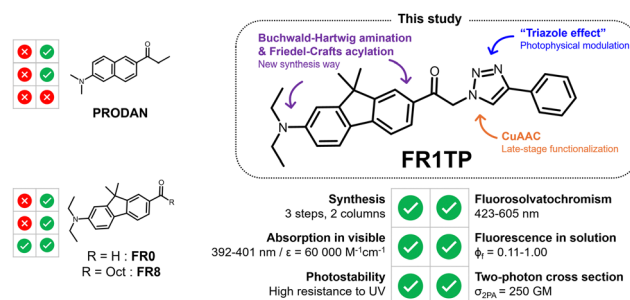


Fig. 1 Main features of **FR1TP** and parent compounds (**PRODAN**, **FR0** and **FR8**).

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limitation implies that *ad hoc* syntheses must be developed for each new application,^{19–21} which hinders broader studies. Click chemistry appears to be the most viable approach for overcoming these barriers of synthetic versatility.^{22,23} In this context, one of the most applied reactions is the Cu(I)-catalysed azide–alkyne cycloaddition (CuAAC), forming a 1,2,3-triazole derivative.

This synthetic strategy has already demonstrated its relevance in bioimaging, including a wide range of targeted applications such as the synthesis of fluorogenic probes,^{24–26} the fabrication of complex bioconjugates,^{27,28} and the visualisation of biologically active small molecules within living cells.²⁹

In this study, we introduce a new group of solvatochromic dyes synthesized from an azide compound (**FR1N3**) as a key intermediate. This approach facilitates the synthesis of **FRT** (“T” for triazole) derivatives *via* CuAAC (Fig. 1). Surprisingly, the triazole moiety, although not a constituent of the donor and acceptor π -system (commonly abbreviated as D– π –A), has a positive effect on the overall optical properties of these new dyes. This “triazole effect” was investigated in a comprehensive spectroscopic study, accompanied by a time-dependent density functional theory (TD-DFT) approach. These results demonstrate the considerable potential of these molecules in bioimaging, due to their excellent two-photon absorption (2PA) properties.

Results and discussion

Synthesis

Building on the seminal methodology developed by Klymchenko and coworkers to synthesise fluorene-based dyes,¹² Cho and coworkers notably simplified the synthetic process to obtain **FR0** (Fig. 2).¹⁹

Inspired by this approach, we applied a similar strategy to the commercially available 2-bromo-9,9-dimethyl-9H-fluorene (Fig. 2). Briefly, the Buchwald–Hartwig amination, followed by Friedel–Crafts acylation, provided a shortcut for the synthesis of the keto-terminated **FR1**, with just two synthetic steps and one chromatography column *vs.* seven steps and three chromatography columns as required for **FR8**. Additional details and characterization data are available in the ESI† (Sections S2 and S3, and Fig. S1–S15).

In analogy to this procedure, azide functionalisation of these fluorenes was envisaged by first introducing a bromoacetyl unit. Optimised Friedel–Crafts acylation of **FREt2N** with bromoacetyl bromide, followed by direct bromo-azido substitution, led to the clickable fluorene **FR1N3**. Finally, submitting **FR1N3** to the CuAAC click reaction with phenylacetylene provided an excellent yield of the model compound **FR1TP**. Importantly, this strategy permits the conjugation of diverse alkynes with our fluorosolvatochromic unit without a significant increase in synthetic effort.

Spectroscopic properties in solution

FR1 and **FR1TP** photophysics were characterised in a broad range of solvents using UV-vis absorption and steady-state as well as time-resolved fluorescence spectroscopy. The spectra are shown in Fig. 3, and the data are summarised in Table S1

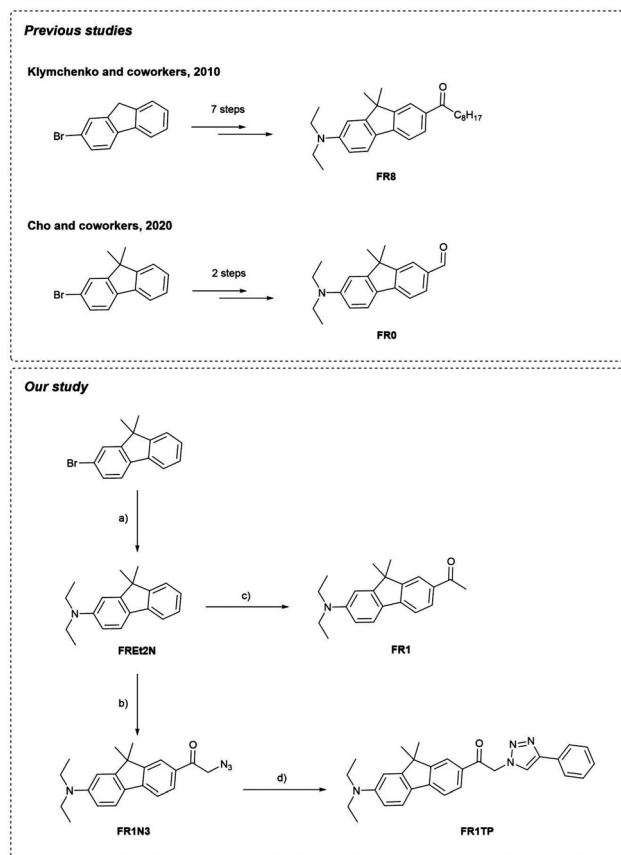


Fig. 2 Synthesis of previously described fluorophores **FR0**, **FR8** and new synthesis of **FR1** and **FR1TP**. Reagents and conditions: (a) Et₂NH, Pd(OAc)₂, BINAP, NaO^tBu, toluene, 110 °C, overnight, 76%; (b) 1. Bromoacetyl bromide, TfOH, 0 °C, 1 h; 2. NaN₃, DMSO, rt, 10 min., 66%; (c) acetyl chloride, AlCl₃, DCM, 0 °C to rt, overnight, 59%; (d) phenylacetylene, [Cu(CH₃CN)₄]PF₆, TBTA, CHCl₃, rt, overnight, 86%.

(ESI†). Additional details and spectra are included in the ESI† (Section S1 and Fig. S16–S21). Like **FR0**, the absorption bands of **FR1** and **FR1TP** do not shift significantly with solvent polarity (Table S1, and Fig. S16–S19, ESI†). Therefore, in Fig. 3, the

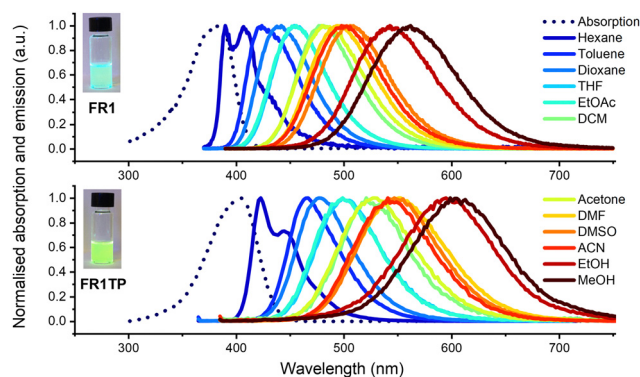


Fig. 3 Absorption (dotted line) in toluene and emission (solid lines) of **FR1** and **FR1TP** in a series of solvents. Concentration around 10^{−5} M. Excitation for emission readout at 360 nm in *n*-hexane, toluene, and dioxane, and at 380 nm for all other solvents. Pictures of acetone solutions of **FR1** and **FR1TP** under irradiation at 365 nm are displayed in the left insets.



absorption spectra in toluene only are shown as an example. Based on the spectral position of the emission bands, a pronounced fluorosolvatochromism was observed for **FR1** and **FR1TP**, similar to that of the parent compounds **FR0** and **FR8**.¹² The structured emission bands in hexane, ascribed to fluorescence originating from a local excited state, further prove the ICT character of these compounds. Note that by varying the solvent polarity, **FR1** and **FR1TP** display emission spectra across the entire visible region.

Surprisingly, the emission bands of **FR1TP** are systematically redshifted by 47 nm on average, compared to those of **FR1** for a given solvent. This desired property was not anticipated since the inherent properties of push-pull dyes are mainly dictated by the composition of the π -system, *i.e.* (i) the nature of the π -bridge and (ii) the D-A system.³⁰ However, **FR1** and **FR1TP** share a common electronic system: only the additional 4-phenyl-1,2,3-triazole differentiates **FR1TP** from **FR1**. Since this unit is isolated from the main π -system by an sp^3 carbon, no significant electronic communication is expected between the triazole and the fluorophore moieties. Therefore, the triazole group must influence the π -system without participating in the overall π -conjugation. To our knowledge, such an effect attributed to non-conjugated triazole has not been described previously. To verify this hypothesis, two other derivatives with the same D- π -A structure were synthesised (Fig. 4a): (i) **FR1TM**, a mesityl derivative, to characterise the impact of the steric hindrance in position 4 of the triazole; and (ii) **FR2TP**, a

derivative containing an ethyl linker, to explore the impact of the triazole-fluorophore distance. The synthetic routes of **FR1TM** and **FR2TP** together with the absorption and emission spectra of these compounds are provided in the ESI† (Scheme S1 and Fig. S20 and S21). The absorption bands of these fluorene derivatives show only small variations with solvent polarity (Fig. S22 and Table S2, ESI†). Importantly, the fluorosolvatochromism trends outlined above for **FR1** and **FR1TP** are observed also for these new analogues (see Fig. S20 and S21, ESI†). The fluorescence properties of all the newly synthesised **FR**-analogues were further investigated, and their fluorescence quantum yields were measured across all the solvents studied. From the data presented in Fig. 4b, the following observations can be made:

(i) The influence of the alkyl chain in the α -position on the ketone is negligible: both **FR1** and **FR8** display similar emission bands and fluorescence quantum yields.

(ii) The effect of substituting the triazole at position 4 is marginal: **FR1TM** has a slightly blue-shifted emission compared to **FR1TP** (by 3 nm on average for the same solvent), indicating that steric hindrance in this position is not relevant to the fluorescence properties.

(iii) The “triazole effect” depends strongly on the fluorene-triazole distance: the emission maxima of **FR2TP** and **FR1TP** are redshifted by 20 nm and 47 nm on average, respectively, for a given solvent compared to **FR1** (lacking the triazole). This observation clearly demonstrates that an increase in the fluorene-triazole distance significantly reduces the redshift, and strengthens the hypothesis that the triazole unit induces the bathochromic effect.

(iv) In hexane, **FR1** (as **FR8**) display weak fluorescence only ($\phi_f < 0.05$), whereas the triazole derivatives show a substantial increase in the fluorescence quantum yield ($\phi_f > 0.70$). **FR0** and **PRODAN** were previously found to be prone to aggregation in hexane.³¹ The induced excitonic couplings also observed for **FR1**, which strongly favour non-radiative decay, are significantly inhibited by the triazole for **FR1TP** (see the fluorescence lifetime study below).

(v) All analogues are characterised by exceptional fluorescence quantum yields in nonpolar or slightly polar solvents, which gradually decrease with increasing solvent polarity. This behaviour, classically observed in push-pull architectures,^{12,32} is observed also for the triazole-decorated derivatives.

Additionally, fluorescence lifetimes were determined in a few representative solvents and are shown in Fig. S23 and Tables S3 and S4 (ESI†). In 1,4-dioxane, the radiative and non-radiative rate constants (k_r and k_{nr} , respectively) of these compounds, with or without triazole, are of the same order globally ($\langle k_r \rangle = 0.43 \text{ ns}^{-1}$ and $\langle k_{nr} \rangle = 0.55 \text{ ns}^{-1}$). In hexane, the triazole compounds showed on average a slightly more efficient non-radiative deactivation compared to 1,4-dioxane ($\langle k_r \rangle = 0.52 \text{ ns}^{-1}$ and $\langle k_{nr} \rangle = 0.94 \text{ ns}^{-1}$), likely due to their lower solubility in hexane. The negative influence of hexane on **FR1** resulted in a very short lifetime ($< 0.05 \text{ ns}$), with a non-radiative rate at least 6 times higher than the triazole derivatives.

On the other hand, using protic solvents such as methanol or ethanol implies solvent-specific interactions for the triazole

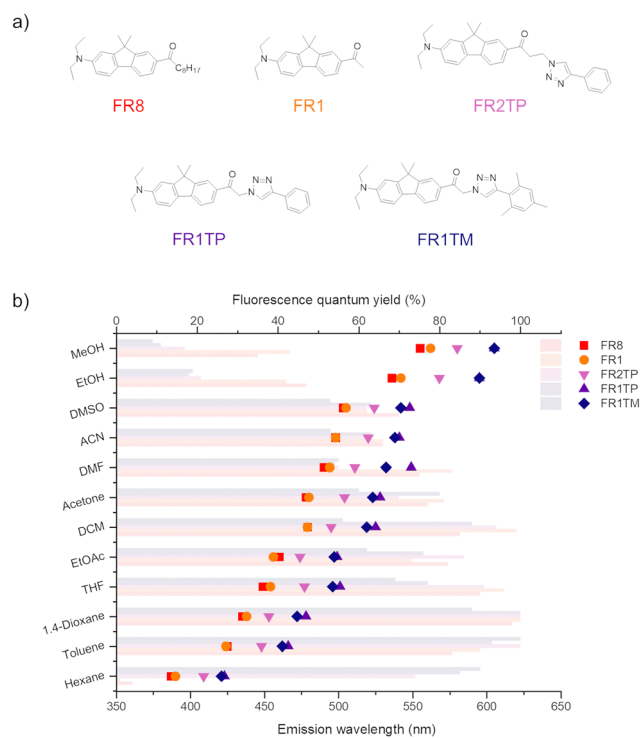


Fig. 4 (a) Structures of the fluorene derivatives studied in this work. (b) Fluorescence quantum yields (faded horizontal bars) and emission maxima (symbols) of **FR8** (red), **FR1** (orange), **FR1TP** (pink), **FR2TP** (purple) and **FR1TM** (dark blue) in different solvents.



derivatives.³³ In ethanol, **FR1TP** and **FR1TM** display shorter fluorescence lifetimes than **FR1** (1.89 and 1.85 ns for **FR1TP** and **FR1TM** vs. 2.99 ns for **FR1**). More interestingly, **FR2TP** has a similar pattern to **FR1**: consequently, the proximity of the triazole to the fluorene moiety is detrimental to proticity-induced non-radiative relaxation. In aprotic polar solvents however, such as dimethylsulfoxide, the differences in the lifetimes are negligible, proving the isolated effect of the proticity. To further investigate the impact of solvent proticity on the fluorescence properties, deuterated methanol was employed to minimize proton interaction with the dyes.³⁴ The shape and the spectral position of the emission bands of **FR1** and **FR1TP** were unchanged in deuterated methanol compared to regular methanol, indicating no influence on the electronic properties (Fig. S24a, ESI[†]). However, both **FR1** and **FR1TP** experienced a significant increases in the fluorescence quantum yields when changing from methanol to deuterated methanol: from 0.43 to 0.62 for **FR1** and from 0.11 to 0.26 for **FR1TP** (Fig. S24b, ESI[†]). This suggests that solvent proton interaction mainly contributes to fluorescence quenching in polar protic solvents, aligning with other dye systems.³⁴ Additionally, this effect was more pronounced for **FR1TP** than for **FR1**, with an increased quantum yield by factors of 2.4 for **FR1TP** and 1.4 for **FR1**. Consequently, the more efficient fluorescence quenching of **FR1TP** compared to **FR1** implies a distinctive interaction between the carbonyl/triazole system of **FR1TP** and the solvent proton.

Characterisation of the triazole effect

The experimental results described above show that the introduction of the triazole at different positions significantly influences the photophysical properties. To support the experimental observations, theoretical calculations were performed using the mPW1PW91/6-311+G(2d,p) level of theory. Calculations were conducted in vacuum and with various solvents (toluene, dichloromethane, and acetonitrile). The results are summarised in Table S9 (ESI[†]), and more details are provided in the ESI[†]. The experimental trends are satisfactorily reproduced, and it is also encouraging to note the good agreement between the calculated and the experimental values.

The lowest transition, $S_0 \rightarrow S_1$, is dominated in all cases by HOMO and LUMO frontier orbitals, as represented in Fig. 5. For all derivatives, the HOMO is mainly localised around the diethylamino moiety, whereas the electron density is strongly shifted towards the carbonyl moiety in the LUMO. This observation confirms the D- π -A configuration of these compounds. However, the LUMO distribution varies between the analogues, and a minor triazole contribution is identified from these calculations, especially for derivatives **FR1TP** and **FR1TM**. Importantly, the electronic distribution of these orbitals along the scaffolds remains consistent, irrespective of the solvent considered in the calculations (Fig. S32–S34, ESI[†]). A better representation of the electron-density changes and the intramolecular charge transfer (ICT) phenomena is given by the electron density difference (EDD) plots (Fig. S30, ESI[†]).

Analysis of the orbital energies further supports the role of triazole in the bathochromic shift. Notably, the relative energies of the HOMO levels are almost identical for all studied compounds.

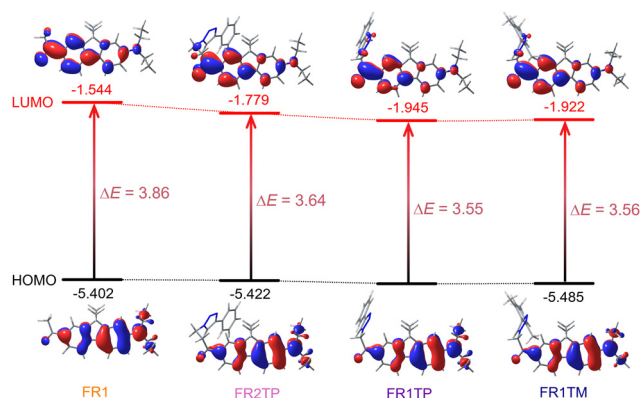


Fig. 5 Frontier molecular orbitals contour plots and energies for **FR1**, **FR2TP**, **FR1TP** and **FR1TM** at the mPW1PW91/6-311+G(2d,p)//mPW1PW91/6-31+G(d) level (isosurface value: 0.03 a.u.). Orbital energies indicated are in eV.

However, the LUMO levels differ substantially, clearly showing stabilisation by the triazole unit. More specifically, in vacuum, the energy associated with the LUMO of **FR1** was -1.544 eV, with a decrease of 14% for **FR2TP** and 24% for **FR1TP** and **FR1TM**. Thus, the proximity of the triazole motif to the fluorene part induces a decrease in the excitation energy. This behaviour is also well reproduced by TDDFT calculations performed in vacuum: both absorption and emission wavelengths are redshifted upon the incorporation of the triazole unit onto the scaffolds (Table S9, ESI[†]). For instance, the excitation energy is reduced from 3.86 eV for **FR1** to 3.56 eV for **FR1TP**. Furthermore, similar trends are observed in the relative energies of frontier molecular orbitals when solvent effects are included in the DFT calculations (Fig. S32–S34, ESI[†]).

The “triazole effect” on the fluorosolvatochromism of **FRT** derivatives can also be quantified at the experimental level. To evaluate the amplitude of the solvatochromic character of a given chromophore, two models are commonly used:¹² (i) the correlation between the solvent’s $E_T(30)$ parameter and the corresponding emission maxima, or (ii) the correlation between the Stokes shift and the orientation polarizability function for each solvent (known as the Lippert–Mataga plot^{35,36}). Using both approaches, the linear correlations obtained for all compounds suggest a noticeable effect of solvent polarity on their respective spectroscopic properties (Fig. S25 and S26/Tables S5–S7, ESI[†]). Furthermore, the difference in dipole moments between the ground- and excited states extracted from the Lippert–Mataga model reveals that the triazole derivatives display a slightly increased charge-transfer (CT) character: 16 D for **FR1**, 17 D for **FR2TP** vs. 20 D for **FR1TP** and **FR1TM**. Therefore, the triazole substitution in this chromophore family significantly affects the fluorosolvatochromism.

In addition to the bathochromic shift caused by the “triazole effect”, the molar absorption coefficients in ethanol are also substantially increased for the triazole-substituted analogues: 26 000, 60 000, and 57 000 $M^{-1} cm^{-1}$ for **FR1**, **FR1TP**, and **FR1TM**, respectively. The photostability of these dyes was investigated by monitoring the decrease in fluorescence intensity over time under constant irradiation at 365 nm. This data shows an overall equal or



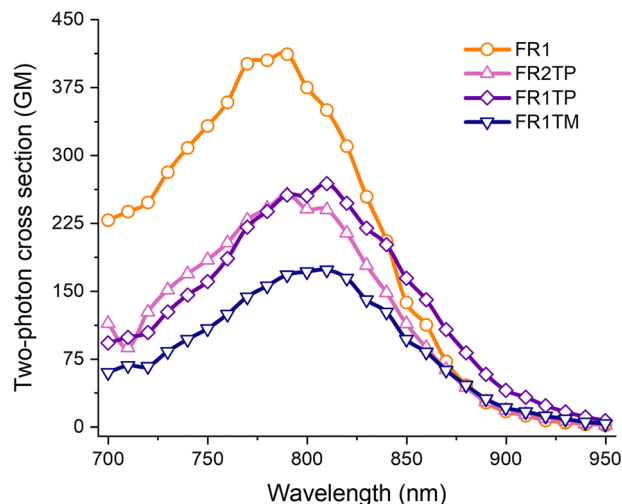


Fig. 6 2PA spectra of **FR1** (orange), **FR2TP** (pink), **FR1TP** (purple) and **FR1TM** (dark blue) in ethanol (*ca.* 1 μ M air-equilibrated solutions).

superior photostability of the fluorene dyes compared to **PRODAN** (see Fig. S32 and Table S8, ESI[†]). In ethanol, **FR1TP** shows minimal photodecomposition compared to **PRODAN** or **FR1**. **FR1** and **FR1TP** are more stable than **PRODAN** in toluene. Only the photostability of **FR1TP** is comparable to **PRODAN** in THF.

Altogether, these compounds display photophysical properties highly desired for applications in bioimaging. This led us to study the 2PA properties (Fig. 6). **FR0** has a high cross section (σ_{2PA} up to 350 GM in ethanol), indicating that **FR1TP** and its derivatives could be suitable for applications such as two-photon fluorescence microscopy (2PFM). The two-photon excited fluorescence spectra of **FR1**, **FR1TP**, **FR1TM** and **FR2TP** were recorded in ethanol solution in the 700–950 nm range using a femtosecond pulsed laser source, and the two-photon absorption cross section was calculated according to the two-photon induced fluorescence (2PIF) method (more details are provided in the ESI[†]). The 2PA spectra of all analogues feature a maximum between 780 and 800 nm, essentially coinciding with the respective $S_0 \rightarrow S_1$ absorption bands. These 2PA bands show high intensity, manifested in the high cross section values: 400 GM for **FR1** (in agreement with that observed for **FR8**),¹² 250 GM for **FR1TP** and **FR2TP**, and 170 GM for **FR1TM**. These values indicate that the presence of triazole slightly reduces the 2PA capability. The emission observed was generated in all cases by 2PA, as demonstrated by the slopes of *ca.* 2 determined from the double logarithmic plot representing emission intensity *vs.* laser excitation power (Fig. S28 and Table S9, ESI[†]). Furthermore, the fluorescence bands were not affected in spectral position or shape by the excitation mechanism. These observations confirm that the same electronic states are involved in both the absorption and emission processes, regardless of the nature of the excitation (Fig. S29, ESI[†]).

Conclusions

In summary, we have developed a short synthesis of clickable solvatochromic fluorene derivatives. The new model derivative

FR1TP shows a dramatic effect of the triazole on the spectroscopic properties. We demonstrated experimentally and by TD-DFT that the non-conjugated triazole is responsible for a bathochromic shift in the fluorosolvatochromism of **FR1TP** and similar analogues. Specifically, the triazole stabilizes the LUMO of the fluorene moiety, reducing the overall excitation energy. The pronounced photostability, high molar absorption coefficients and two-photon cross section of these triazole derivatives make them excellent candidates for bioimaging. We anticipate that these results will facilitate the development of new bioconjugates or other types of dyads, to modulate the photophysics of these fluorophores. Studies in this direction are underway in our group.

Author contributions

J. R.: conceptualization, formal analysis, investigation, methodology, visualization, writing – original draft. C. B. M.: formal analysis, investigation, methodology, visualization, writing – review & editing. J. A.: formal analysis, funding acquisition, methodology, project administration, supervision, writing – review & editing. M. G.: formal analysis, funding acquisition, methodology, project administration, supervision, writing – review & editing.

Data availability

Relevant data selected for open access will be shared *via* Zenodo.

Conflicts of interest

There are no conflicts of interest to declare.

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