


 Cite this: *RSC Adv.*, 2023, **13**, 1757

Photophysical and anion sensing properties of a triphenylamine–dioxaborinine trimeric compound†

 Alexis Tigreros, ^a Camilo Bedoya-Malagón, ^b Alejandra Valencia, ^b Mayerlin Núñez-Portela ^b and Jaime Portilla ^{*a}

Herein, we report the synthesis and photophysical characterization of the novel tris(4-(2,2-difluoro-6-methyl-2*H*-1*λ*³,3,2*λ*⁴-dioxaborinin-4-yl)phenyl)amine trimeric probe (**A2**) *via* the reaction between triphenylamine (**1**), acetic anhydride, and $\text{BF}_3 \cdot \text{OEt}_2$ implying the twelve new bond formation in a one-pot manner. This highly fluorescent compound in solution (φ up to 0.91 at 572 nm) and solid state (φ = 0.24 at 571 nm) showed a better solvatofluorochromism than its analog monomeric **A1** due to symmetry-broken charge transfer, which is consistent with high solvent dipolarity (SdP) response in Catalán's multiparametric regression. Notably, **A2** had a high sensibility and selectivity for CN^- or F^- in solution ($\text{LOD CN}^-/\text{F}^- = 0.18/0.70 \mu\text{M}$), and CN^- can be discriminated from F^- by the reaction of **A2** with 3.0 equiv. of CN^- . In addition, **A2** was impregnated on filter paper to prepare test strips that were applied to naked-eye qualitative sensing of CN^- or F^- . Finally, the octupolar system in **A2** allows for better action of two-photon excitation cross-section values when compared with that of the dipolar structure in **A1**. These findings provide further information for the design of new efficient two-photon absorption dyes.

Received 25th November 2022

Accepted 29th December 2022

DOI: 10.1039/d2ra07498b

rsc.li/rsc-advances

Introduction

Boron complex-containing molecules are an interesting and extensively used class of organic fluorescent compounds. These dyes have valuable photophysical properties such as strong absorption bands, high fluorescent quantum yields (φ), good solubility in organic solvents, photostability, microenvironment-dependent emission, *etc.*^{1–3} Therefore, this important family of organic fluorophores can serve in numerous applications involving bioimaging probes,⁴ photosensitizers in photodynamic therapy,⁵ red-emitting complexes with a mega-large Stokes shift,⁶ multicolor fluorescent initiators,⁷ colored triboluminescence compounds,⁸ and fluorescent probes for mercury detection in living cells,⁷ among others.

Importantly, some characteristic fluorophores perform through two-photon absorption (TPA) phenomena that involve the simultaneous absorption of two photons from a laser light source,⁹ which has advantages over the classical one-photon process. For instance, a molecule can be excited with low-

energy photons *versus* the classical methods. The TPA transition probability increases with the excitation laser intensity's square giving the optical absorption's high spatial selectivity,¹⁰ leading to applications in three-dimensional (3D) data storage,¹¹ photodynamic therapy,^{12,13} 3D micro-fabrication,¹⁴ and high-resolution bioimaging.¹⁵ Thus, novel fluorophores development by efficient synthetic approach is highly desirable and a research-active field in chemistry, material sciences, and the industry.

Structurally, many architectures have been used to achieve good TPA responses; that is, conjugated donor (D) and acceptor (A) with different geometry organizations, dipolar (D–π–A), quadrupolar (D–π–A–π–D or A–π–D–π–A), and octupolar (D–(π–A)₃ or A–(π–D)₃),¹⁶ this appreciation has been verified by theoretical and experimentally results in the Jean-Luc Brédas research group.¹⁷ In this way, the electronic connecting between strategic fluorophores is a good tactic for designing dyes with TPA properties; for example, hybrid fluorophores containing triphenylamine (TPAm) or the boron complex dioxaborinine (DB) have been used for this purpose.^{9,18}

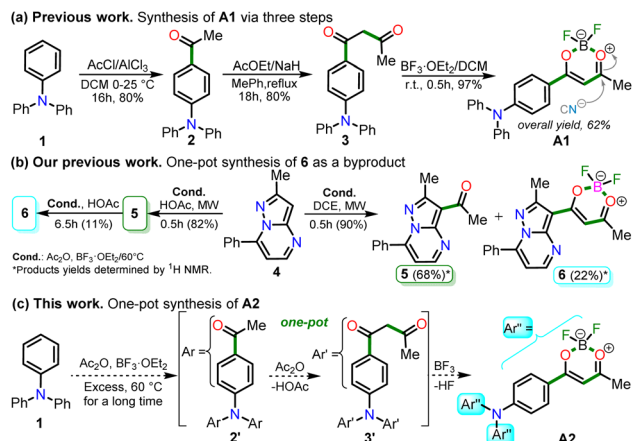
In 2020, Tamilarasan *et al.*¹⁹ synthesized the dipolar dye **A1** (Scheme 1a) based on a triphenylamine–dioxaborinine hybrid compound as a colorimetric and fluorimetric probe for the reversible detection of cyanide ($\text{LOD} = 0.36 \mu\text{M}$ in $\text{MeCN} : \text{H}_2\text{O}$ 98 : 2). The probe was obtained in 62% global yield by a three-step sequence starting from TPAm (**1**). This synthesis involves a sequential double acetylation reaction *via* the carbonyl compounds intermediates **2** and **3**, and the final formation of

^aBioorganic Compounds Research Group, Department of Chemistry, Universidad de Los Andes, Carrera 1 No. 18A-10, Bogotá 111711, Colombia. E-mail: jportill@uniandes.edu.co

^bQuantum Optics Laboratory, Department of Physics, Universidad de Los Andes, Carrera 1 No. 18A-10, Bogotá, Colombia

† Electronic supplementary information (ESI) available: Characterization data, experimental, spectra (NMR, HRMS, absorption, and emission), green metrics, and computational details of fluorophores. See DOI: <https://doi.org/10.1039/d2ra07498b>

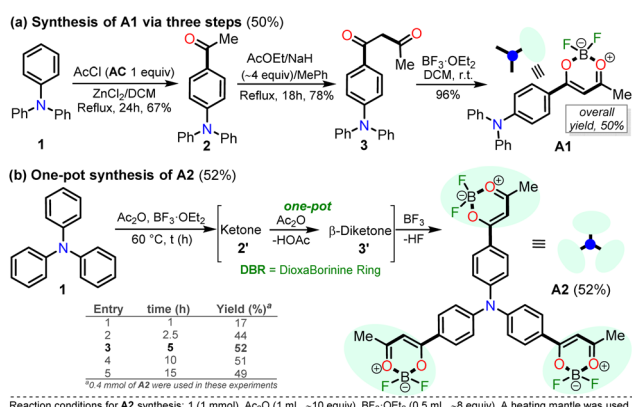




Scheme 1 Synthesis of fluorophores bearing dioxaborinine (a) A1,¹⁹ (b) 5, and (c) A2.²⁰

the boron complex in **A1**. Recently, we carried out a BF_3 -mediated synthesis of 3-acetylpyrazolo[1,5-*a*]pyrimidines (similar to 5) under microwave (MW) irradiation and using acetic anhydride as an acetylating agent (Scheme 2b). During the optimizing reaction conditions process, we observed that the fluorescent by-product 6, bearing the pyrazolo[1,5-*a*]pyrimidine-dioxaborinine hybrid system resulting from the formation of four new bonds in a one-pot manner, is favored in long reaction times.²⁰

Within the most conventional photophysical applications of organic fluorophores, ion recognition has been the subject of extensive study in the last two decades;^{21,22} in particular, cyanide (CN^-) is one of the most concerning environmental ions. Cyanide toxicity is known because it can inhibit the mitochondrial cytochrome C oxidase and suppress oxygen transport.^{23,24} Consequently, the development of sensitive and selective probes for the sensing of this anion has been gaining considerable attention in recent years because of its essential role in a wide variety of biological,²⁵ environmental,²⁶ biochemical,²⁷ clinical,²⁸ synthetical due to design and preparation of the respective chemosensors and industrial applications.^{21,22}



Scheme 2 Synthesis of dioxaborinine-triphenylamine hybrid dyes (a) A1 and (b) A2.

Considering the above background and our continuing interest in obtaining new functional fluorophores,^{20,29-31} we planned to develop the octupolar dye **A2** by a synthesis implying the twelve new bonds formation in a one-pot manner (Scheme 2c). Fluorophore **A2** is a tris-dioxaborinine-triphenylamine hybrid compound with promising photophysical properties in solution and solid-state, which we wish to study and compare with those for the similar dipolar system **A1**. Explicitly, **A2** is expected to have good TPA responses with improved results over other octupolar dyes synthesized even by difficult methods involving various reaction steps;³² for this purpose and as a proof-of-concept, by measuring the two-photon excitation action cross section when exciting the probes **A1** and **A2** with a continuous-wave (CW) laser at 810 nm. Ultimately, similar to **A1**,¹⁹ the novel dye **A2** could be applied for cyanide sensing by fluorescent changes after a chemical reaction with the anion.

Results and discussion

Synthesis

For this research, the dioxaborinine-triphenylamine hybrid dyes **A1** and **A2** were synthesized. The synthetical approaches for preparing these dyes starting from triphenylamine (1, TPAm), are simple and proceed in good yields (Scheme 2). According to the literature,^{19,33,34} compound **A1** was synthesized in 50% global yield *via* a three-step sequence starting from triphenylamine (1). Nevertheless, for **A2** synthesis, a new approach was used to convert substrate 1 into the trisubstituted derivate *via* the twelve new bonds formation in a one-pot manner in good yield (52%). In this respect, the mixture of acetic anhydride (AA) with boron trifluoride diethyl etherate ($\text{BF}_3 \cdot \text{OEt}_2$) was used as multiple acetylating agents six times to give the respective β -diketone intermediate 3' and the complexing agent of the last step. Structures for the hybrid dyes **A1** and **A2** were established by NMR spectroscopy (^1H , ^{13}C , and ^{19}F) and HRMS analysis (Fig. S1-S9†).

Remarkably, although the hybrid compounds **A1** (50%) and **A2** (52%) were obtained with close yields starting from triphenylamine (1), the octupolar dye **A2** requires only one step for its formation in 5 hours at 60 °C and with the twelve new bonds formation. In contrast, the dipolar derivative **A1** synthesis (Scheme 2a) consumptions three steps (*i.e.*, ring acetylation/one new bond, ketone α -acetylation/one new bond, and complexation with BF_3 /two new bond) through the intermediates (4-acetylphenyl)diphenylamine (2) and 1-(4-diphenylamino)phenylbutane-1,3-dione (3). Thus, the synthesis of **A2** implies a much better operational simplicity, lower consumption of solvents due to the solvents used as reaction medium and in the purification steps and a more excellent atomic economy concerning the probe **A1** as a result of the one-pot synthesis of **A2**.

It is important to note that the reaction time played a crucial role in **A2** synthesis since, after 1 hour at 60 °C, the product was obtained with only a 17% yield; however, the yield increased to 44% after 2.5 hours of reaction. The optimal conditions for this reaction turned out to be 5 hours at 60 °C because the yield increased to 52%, and no noticeable changes were observed during longer reaction times; with lower temperatures, the

reaction does not progress much, and with higher temperatures, by TLC, a complex mixture of products was observed (Scheme 2b). Consequently, our previous results on constructing the dioxaborinine ring are a reliable starting point for accessing various derivatives of this heterocyclic core.²⁰ Additionally, it was possible to establish that the optimum temperature to treat the acetylating mixture (an excess of $\text{Ac}_2\text{O}/\text{BF}_3\cdot\text{OEt}_2$) is 60 °C. Finally, we could verify our hypothesis that the formation of the DB ring in a one-pot manner is enhanced when long reaction times are used.

Photophysical properties

Solvatofluorochromism. Solvent-dependent optical properties of compounds **A1** and **A2** were evaluated by UV-vis absorption and emission measurements in a set of non-protic solvents (Fig. 1, Table 1, and eqn (1); see the Experimental section) such as toluene (PhMe), *tert*-butyl methyl ether (TBME), tetrahydrofuran (THF), ethyl acetate (EA), chloroform (CHCl₃), tetrahydrofuran (THF), ethyl acetate (EA), chloroform (CHCl₃),

N,N-dimethylformamide (DMF), and acetonitrile (MeCN). For instance, in toluene, two absorption bands were observed for **A1** (295 and 427 nm) and **A2** (315 and 446 nm), the latter having a molar absorptivity at least twice that of **A1**. This feature is ascribed to the degeneracy of the S1 state of **A2** and the increased number of DB groups that alter the electronic energy levels of the triphenylamine core by π -conjugation.³⁵ Meanwhile, little to non-interaction at the ground state was noticed since there was no solvatochromism in **A1** and **A2**.

Moreover, emission bands of **A1** shift bathochromically from the weakly polar toluene ($\lambda_{\text{em}} = 461$ nm) to the strongly polar acetonitrile ($\lambda_{\text{em}} = 492$ nm); while **A2** displays an intense solvatofluorochromism ($\lambda_{\text{em}}/\text{MePh} = 484$ nm to $\lambda_{\text{em}}/\text{MeCN} = 575$ nm), exhibiting a more dipolarity in the excited state, which may correspond to a symmetry broken dipolar state.³⁶ Catalán multiparametric regression analysis was performed to evaluate the polarity at the excited state of **A1** and **A2** (eqn (2), Table S1†). The solvent dipolarity (SdP) stabilized best the excited state of **A1** (slope = -1937.90 cm^{-1} , $R^2 = 0.8862$) and **A2** (slope = -4607.54 cm^{-1} , $R^2 = 0.9438$). In general, compound **A2** displays a more polar structure at the excited state when compared with **A1** (Fig. S12 and S13†). This result is consistent with previous studies demonstrating the solvent-induced symmetry-breaking charge transfer in an octupolar chromophore.³⁵ It is important to note that the branching increases the fluorescence quantum yield in a low to medium-polarity solvent (Table 1).

Solid-state emission. The fluorescence spectra in the solid-state of compounds **A1** and **A2** were registered under excitation at 300 nm at room temperature, and the results were fortunately very satisfactory (Fig. 2 and Table 1, entry 8). Emission of **A1** in the solid-state falls in the orange-red region (608 nm) and that of **A2** in the yellow-orange region (571 nm). Curiously, the solid-state emission spectrum of **A2** resembles that of **A2** dissolved in a high-polarity solvent. In contrast, the solid-state emission spectrum of the dipolar chromophore **A1** is highly red-shift from that in solution (498 nm in MeCN). Such a marked difference in solid-state emission of the dyes **A1** and **A2** can be related possibly to the different packing of the dipolar and octupolar molecules in the solid state.³⁷ Consequently, a strong intermolecular donor–acceptor interaction is expected

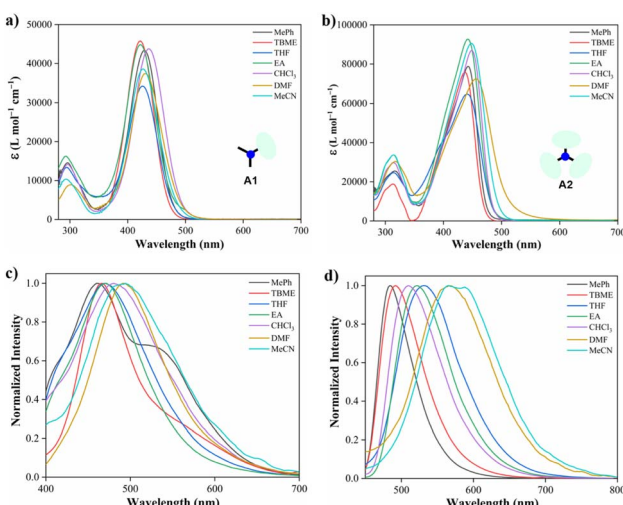


Fig. 1 Absorption and emission ($\lambda_{\text{ex}} = 350$ nm) spectra in various solvents (10.0 μM) at 20 °C of hybrid compounds **A1** (a and c) and **A2** (b and d).

Table 1 Photophysical properties of the hybrid compounds **A1** and **A2**^a

Entry	Solvent	A1		A2	
		λ_{abs} (nm), ε ($\text{L mol}^{-1} \text{cm}^{-1}$)	λ_{em} (nm), ϕ (%)	λ_{abs} (nm), ε ($\text{L mol}^{-1} \text{cm}^{-1}$)	λ_{em} (nm), ϕ (%)
1	PhMe	429, 43 300	461, 68	443, 78 700	484, 91
2	TBME	421, 45 700	467, 25	437, 76 000	492, 90
3	THF	425, 34 200	470, 12	440, 64 800	510, 29
4	EA	422, 44 800	470, 40	441, 92 600	522, 74
5	CHCl ₃	436, 43 800	480, 53	449, 87 100	531, 85
6	DMF	430, 37 300	492, 22	456, 72 400	567, 05
7	MeCN	426, 38 600	492, 13	457, 90 300	572, 08
8 ^b	—	—	608, 11	—	571, 24

^a Quantum yields (ϕ) were determined using Prodan as a reference standard (see eqn (1)). ^b Solid-state.



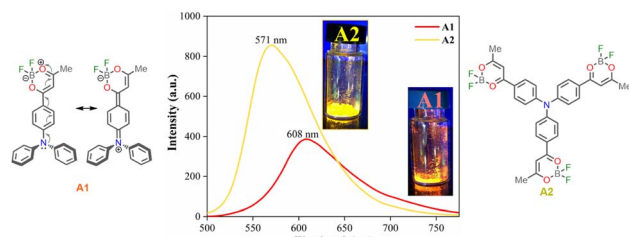


Fig. 2 Fluorescence spectra for dyes **A1** and **A2** in the solid-state at 20 °C, $\lambda_{\text{ex}} = 350$ nm.

in the solid phase for the dipolar dye **A1**. Another explanation could rely on the formation of J-aggregates that usually show different optical properties from dyes in solution, including red-shifted absorption and emission spectra and enhanced fluorescence quantum yields.³⁸

Response to anions sensing in solution. Recently, Tamilarasan *et al.*¹⁹ studied compound **A1** as a probe for cyanide sensing applications due to the presence of the dioxaborinine unit as an excellent cyanide-acceptor group. Inspired by these previous studies and our interest in developing molecular probes to detect cyanide,^{21,22,31} we envisioned dye **A2** could show an interesting behavior in this sensing field. Thus, the anion sensing property of **A2** was evaluated by treating the probe with CN^- and other anions of the potassium or sodium salts (100 μM), including F^- , Cl^- , Br^- , I^- , AcO^- , IO_4^- , PO_4^{3-} , HSO_4^- , HCO_3^- , and ClO^- using MeCN/Tris (9 : 1, 1.0 mM at pH 7.5) as a solvent. When CN^- (1.4 equiv.) was added to the dye solution, the emission band around 572 nm disappeared, and the color of the solution changed from orange to pale yellow; in contrast, by adding other different anions, except by fluorine (F^-), no substantial changes in the emission spectra were observed (Fig. 3a).

Due to the good preliminary results using the hybrid compound **A2** for anions recognition, titration of **A2** with F^- in acetonitrile/Tris was carried out the interaction was monitored by fluorescence at 572 nm (Fig. 3b). Upon the addition of 6.0 equiv. of F^- , the emission band at 572 nm decreased linearly with a concentration (conc.) increased from 0 to 60.0 μM ; the limit of detection (LOD), calculated by the expression $\text{LOD} = 3 \times \text{SD}/\sigma$, where σ is the slope of the titration curve, and SD is the standard deviation of ten measurements of the blank, for F^- was found to be 0.70 μM . These preliminary results allow us to

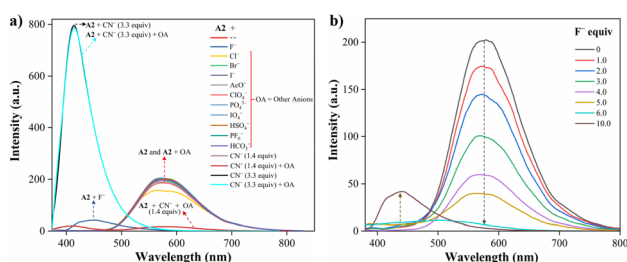


Fig. 3 Emission spectra of **A2** (10.0 μM , $\lambda_{\text{ex}} = 350$ nm) in MeCN/Tris (9 : 1) at 20 °C (a) with various anions (100.0 μM) and CN^- (14.0 and 33.0 μM), and (b) with F^- (0–100.0 μM).

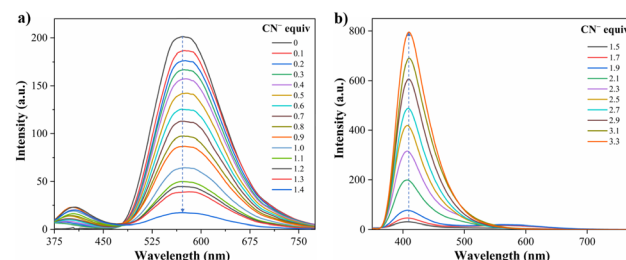


Fig. 4 Emission spectra of **A2** (10.0 μM , $\lambda_{\text{ex}} = 350$ nm) in MeCN/Tris (9 : 1) at 20 °C in the presence of CN^- (a) 0–14.0 μM and (b) 15.0–33.0 μM .

conclude that the sensitivity of **A2** toward CN^- is much higher than that found for F^- ; indeed, high fluorine concentrations are needed to complete the decrease in the emission band. Thus, titration with CN^- was also carried out to establish dye **A2** as an efficient probe to detect cyanide (Fig. 4). The fluorescence intensity of the emission band at 572 nm in **A2** decreased linearly when the CN^- concentration in the range of 0 to 14.0 μM (Fig. 4a). Noticeable, after 1.4 equiv. of CN^- , a new emission band appears around 410 nm, and the fluorescence intensity of the band increased linearly with the concentration of CN^- in the range of 15.0 to 33.0 μM (Fig. 4b). Finally, the LOD for CN^- was evaluated to be 0.18 μM from Fig. S3 and S6.† These results indicate that **A2** is a sensitive probe for CN^- sensing by fluorimetric methods, showing a LOD far below the WHO suggestion (1.9 μM) for drinking water.³⁹ Moreover, comparing these results with those reported for **A1** (LOD = 0.36 μM)¹⁹ indicates that an octupolar architecture improves the cyanide sensing performance.

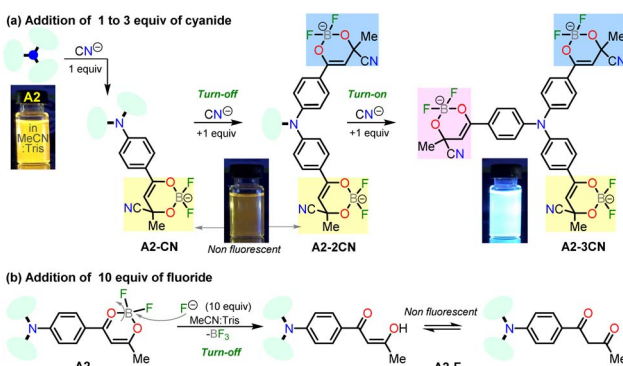
Several chemosensors containing dioxaborinine core as the signaling subunit have been described recently; therefore,

Table 2 Comparison of representative cyanide chemosensors, based on dioxaborinine core, with the new probe **A2**^a

Compound	Features, LOD, and solvent	Reference
DMAPh-DB	“Turn on” 2.23 μM , H_2O $\lambda_{\text{em}} = 620$ nm	Gao <i>et al.</i> ⁴⁰
Ar ₂ -DB	“Turn on” 0.14 μM , $\text{H}_2\text{O}/\text{MeCN}$ (1 : 4) $\lambda_{\text{abs}} = 649$ nm	Chaicham <i>et al.</i> ⁴¹
Coumarin-DB	“Turn off” 72 nM, PBS/DMSO (3 : 2, pH 7.4) $\lambda_{\text{em}} = 565$ nm	Li <i>et al.</i> ⁴²
A1	“Turn on” 0.36 μM , MeCN/ H_2O (98 : 2) $\lambda_{\text{em}} = 402$ nm	Tamilarasan <i>et al.</i> ¹⁹
A2	“Turn on-off-on” 0.18 μM , MeCN/Tris (9 : 1, pH 7.5) $\lambda_{\text{em}} = 572$ nm	This work

^a Data were recorded at different concentrations of **A1** and **A2** in THF at 20 °C (see eqn (3)).





Scheme 3 Plausible mechanisms for anions sensing upon addition of (a) 1 to 3 equiv. of CN⁻ or (b) 10 equiv. of F⁻ to solutions of A2.

a representative summary of this type of probe compared to the synthesized dye in this work was carried out (Table 2).^{19,40-42}

Notably, the new hybrid dye A2 reported here demonstrates good sensitivity and selectivity with a relatively simple chemical structure and synthetic pathway. Likewise, and due to its trimeric molecular architecture, A2 is one of the few probes that can detect cyanide ions sequentially through a “turn on-off-on” process (*i.e.*, detection by adding 1 equiv. and then completing up to 3 equiv.). Ultimately, the synthesis of A2 is carried out by an operationally simple and efficient process compared to other probes synthesized for cyanide recognition (Schemes 2 and 3).

Proposed mechanism for anions recognition. The cyanide sensing mechanism for the dipolar dioxaborinine receptor A1 has already been described as a nucleophilic addition of CN⁻ to the sterically less hindered electrophilic carbon (Scheme 1a).¹⁹ Upon adding 1 or 2 equiv. of CN⁻ ions to the solution of A2, the probe emission band at \sim 570 nm disappears with a turn-off fluorescence due to the symmetry broken into the formed complexes A2-1CN and A2-2CN. However, by adding 3.3 equiv. of CN⁻, the donor-acceptor architecture in A2 disappears, and the symmetry is restored; thus, the emission properties now rely on the less π -conjugated tris-vinyl-TPAm moiety in A2-3CN (λ_{em} = 410 nm, Fig. S16†), favoring a redshifted turn-on fluorescence concerning A2 (Scheme 3a). On the other hand, as reported by Yan *et al.*,⁴³ the F⁻ addition is presumed to proceed with the opening of the dioxaborinine ring in A2 due to the attack on the boron atom in the probe molecule (Scheme 3b). The mechanistic assumptions cited were tracked by mass spectrometry analysis for adducts A2-1CN, A2-2CN, and A2-F (Fig. S10 and S11†).

Test strips. Based on the distinct emission color change of A2 under a 365 nm hand-held UV lamp in an acetonitrile

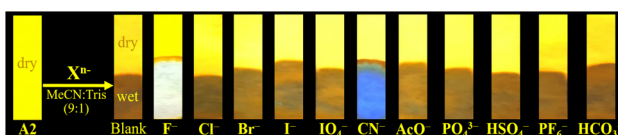


Fig. 5 Images of A2 test strips prepared on filter paper for the selective detection of CN⁻ and F⁻ in MeCN/Tris (9:1) using a hand-held UV lamp (λ_{ex} = 365 nm).

solution by adding cyanide, filter paper immersed with A2 was proposed to detect this anion. The test strip was prepared by simple immersion of qualitative filter paper (Filter Disc, Ref. 3.303.125, Boeco) in a solution of A2 in THF (10 mL at 1.2 mM) followed by air-drying under atmospheric conditions (Fig. 5). Subsequent, anions spiked acetonitrile sample (0.1 mM) was dropped onto the strips for naked eye detection under illumination with a hand-held UV lamp without the paper being completely dry. As depicted, the paper containing A2 displayed bright blue emission only under exposure to cyanide solution. However, the color changed to white after the test strips stained with A2 were immersed into the acetonitrile solutions with fluorine F⁻ (100.0 μ M). Based on the distinctive color change of A2 when exposed to CN⁻ this compound proved that the test strips could be applied to detect CN⁻ and F⁻ anions qualitatively in a rapid way.

Preliminary two-photon absorption properties. The preliminary TPA properties of A1 and A2 in THF were studied by detecting two-photon-induced emissions. The detected fluorescence is shown as a function of the laser light power for different concentrations of dyes (Fig. 6). The dots correspond to experimental measurements, and the dashed lines are quadratic fits to the data (R^2 /conc. in mM: A1 = 0.999/1.0, 0.996/5.0, and 0.999/10.0; A2 = 0.968/1.0, 0.9982/0.1, 0.999/1.0, and 0.997/5.0). From the fitting parameters, the experimental value for the two-photon excitation action cross-section (σ') can be obtained according to eqn (3) (see the Experimental section). The values obtained for σ' for diverse concentrations of A1 and A2 are also reported (Table 3). The technique was verified by measuring the σ' using rhodamine B (RB) in methanol to

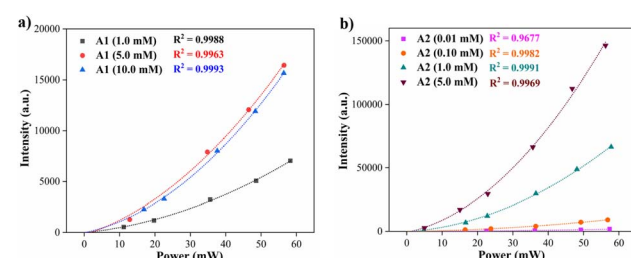


Fig. 6 Two-photon induced fluorescence signal as a function of the laser power of compounds (a) A1 and (b) A2 at different concentrations in THF (20 °C).

Table 3 Two-photon excitation action cross section for dyes A1 and A2^a

Concentration (mM)	σ' (GM)	
	A1	A2
0.01	—	20 \pm 4
0.10	—	12 \pm 2
1.00	0.49 \pm 0.09	8.8 \pm 1.6
5.00	0.25 \pm 0.05	4.3 \pm 0.8
10.00	0.12 \pm 0.02	—

^a Data were recorded at different concentrations of A1 and A2 in THF at 20 °C (see eqn (3)).

0.05 mM.⁴⁴ The value obtained in the experiment is $\sigma' = 4.4 \pm 0.7$ GM. Considering the value of ϕ reported in the literature for RB,⁴⁵ herein we report a TPA cross-section $\delta = 6.8 \pm 1.8$ GM that agrees with the previously reported values (Table S2†). Notably, values found for **A2** are in the same order of magnitude as RB under similar conditions.

The results of two-photon absorption experiments demonstrate that the hybrid fluorophores **A1** and **A2** can induce such processes. The values obtained for the two-photon excitation action cross-section show that trimeric compound **A2** has a higher probability of a two-photon induced fluorescence process when compared with the analog monomeric **A1**. This effect can be attributed to the geometry differences between **A1** (dipolar) and **A2** (octupolar) since the TPA cross-section in octupoles could scale three times the corresponding values in the isolated dipolar analogs.^{17,46–48} Values of the cross sections in Table 3 change due to variation in the fluorescence quantum yield with the concentration⁴⁶ and the possibility of aggregation effects at higher concentrations. Finally, the respective representative diagrams of the z-scan system and the fluorescence process were made (Fig. S18†) to clarify the optical route by which the fluorescence induced by two-photon occurs. Specifically, Jablonski diagrams in Fig. S18b† show the difference between linear fluorescence and two-photon excitation, clarifying the two-photon absorption mechanism of the fluorophore **A2**.

Conclusions

In summary, a highly fluorescent trimeric probe **A2** was synthesized using a one-pot methodology, and its photophysical properties were examined. This compound displays high fluorescence quantum yields in solvents with low to medium polarity (ϕ of 0.91 MePh to 0.85 CHCl₃), interesting emission quantum yield at solid-state ($\phi = 0.24$ to 571 nm), and moderate solvatofluorochromism from toluene (485 nm) to acetonitrile (572 nm). The nucleophilic addition reaction of CN[−] or F[−] on the dioxaborinine ring changes the emission properties of the **A2** solution, giving selective fluorometric detection of these anions with limits of detection of 0.18 and 0.70 μ M, respectively. In addition, a test strip assay using **A2** has also been applied to detect CN[−] or F[−] in organic solutions. Noticeably, CN[−] can be discriminated from F[−] by tracking the emission at 410 nm. Ultimately, a clear two-photon induce process was observed for **A1** and **A2**. In particular, the two-photon excitation action cross section of **A2** shows that this probe can be exploited, as rhodamine B, in different applications, *e.g.*, two-photon microscopy or non-linear optics for the ion sensing field. The extension of the π -conjugation and the possibility of a symmetry-broken dipolar state induce better photophysical properties in compound **A2** than its dipolar analog **A1**.

Experimental section

Reagents and materials

Synthesis. Reagents were purchased from commercial sources and used without further purification; these were weighed

and handled in the air at room temperature. The reaction was monitored by thin-layer chromatography (TLC), visualized by a UV lamp (254 or 365 nm), and flash chromatography was performed on silica gel (230–400 mesh). The methyl ketone precursor (4-acetylphenyl)diphenylamine (2) was obtained in a 67% yield using a known method starting from triphenylamine (1).³³ Subsequently, the β -diketone intermediate 1-(4-(diphenylamino)phenyl)butane-1,3-dione (3) was prepared in a 78% yield from methyl ketone 1 (Scheme 2a).¹⁹

Characterization. For the structural characterization of the hybrid compounds **A1** and **A2**, their NMR spectra (Fig. S1–S7†) were recorded at 400 MHz (¹H), 101 MHz (¹³C/¹H}), and 374 MHz (¹⁹F) at 25 °C using CDCl₃ or DMSO-*d*₆ as solvents and tetramethylsilane (TMS, δ : 0 ppm) as the internal reference. The chemical shifts (δ) are reported in ppm, and the coupling constants (J) are reported in Hz. The following abbreviations are used for multiplicities: s = singlet, d = doublet, and m = multiplet. The melting point was collected using a capillary melting point apparatus and is uncorrected. The high-resolution mass spectra (HRMS) for the hybrid dyes **A1** and **A2** (Fig. S8 and S9†) were obtained on an Agilent Technologies Q-TOF 6520 spectrometer *via* electrospray ionization (ESI). The mass spectra for adducts **A2-1CN**, **A2-2CN**, and **A2-F** (Fig. S10 and S11†) were recorded on a Thermo-Scientific LCQ Fleet™ ion-trap mass spectrometer using positive ion mode ESI and a direct inlet system.

Regarding the photophysical studies of **A1** and **A2**, absorption (UV-vis) and emission spectra were recorded at room temperature (20 °C) in an air-equilibrated solution on Varian Cary 100 and Cary Eclipse spectrophotometers, respectively (both are Agilent Technologies devices) using quartz cuvettes with a path length of 1 cm. For fluorescence studies, both the excitation and emission slit widths were 5 nm. The fluorescence quantum yields (ϕ) were determined using Prodan⁴⁹ as a reference standard by eqn (1)

$$\phi_{f,x} = \phi_{f,st} \frac{F_x}{F_{st}} \frac{A_{st}}{A_x} \frac{n_x^2}{n_{st}^2} \quad (1)$$

where F is the integral photon flux, A is the absorption factor, n is the solvent refractive index, ϕ_f is the quantum yield. The indexes x and st denote the sample and standard, respectively.⁵⁰

On the other hand, Catalán's multiparametric relationship can be formulated by eqn (2)

$$A = A_0 + bSA + cSB + dSP + eSdP \quad (2)$$

where A is a solvent-dependent physicochemical property in a given solvent, and A_0 is the statistical quantity agreeing to the value of the property in the gas phase; SA, SB, SP, and SdP represent independent yet complementary solvent parameters accounting for various types of solute–solvent interactions; and b to e are the regression coefficients relating the sensitivity of property A to the different solute–solvent interaction mechanisms.⁵¹

Finally, the two-photon excitation action cross section (σ') was measured using a continuous-wave (CW) laser at 817 nm



when driving the two-photon transition. The laser light was focused on the sample contained in a 1 mm thick quartz cuvette, employing an objective microscope lens. The sample was placed on a translational stage that allows for implementing the z-scan technique. The fluorescence was detected as a function of the position of the sample by a photomultiplier tube. The dependence of the fluorescence signal (F) with the power of the laser light (P) is given by eqn (3)

$$F \approx g\eta\sigma' C \frac{4nP^2}{\pi\lambda} \quad (3)$$

where g is the temporal second-order correlation function of the incoming light source, η is overall fluorescence collection efficiency, n is the refractive index of the solvent, C is the sample concentration, and $\sigma' = \phi\delta$ (ϕ is the quantum yield and δ the TPA cross-section).⁴⁵

Synthesis and characterization

4-(2,2-Difluoro-6-methyl-2H-1λ³,3,2λ⁴-dioxaborinin-4-yl)-N,N-diphenylaniline (A1). To a stirred solution of the freshly synthesized (54% for the two steps from 1) β-diketone 3 (0.12 g, 0.36 mmol) in dichloromethane (DCM, 5.0 mL), $\text{BF}_3 \cdot \text{OEt}_2$ (0.054 mL, 0.44 mmol) was added dropwise at room temperature and maintained for 30 min. The reaction mixture color changed instantly to red during the $\text{BF}_3 \cdot \text{OEt}_2$ addition. The reaction was quenched with aqueous NaOH solution 0.5 M and extracted with DCM \times 3. The crude product was purified by column chromatography in silica gel (DCM as eluent) to obtain A1 as a red solid in a 96% yield. Mp: 209–212 °C. ¹H NMR (401 MHz, CDCl_3): δ = 2.33 (s, 3H), 6.37 (s, 1H), 6.94 (d, J = 8.9 Hz, 2H), 7.26–7.17 (m, 6H), 7.38 (t, J = 7.7 Hz, 4H), 7.94–7.80 (m, 2H) ppm. ¹³C NMR (100 MHz, CDCl_3): δ = 24.4, 95.9, 118.3, 121.5, 126.0, 126.7, 130.0, 131.3, 145.3, 154.5, 180.9, 188.2 ppm. ¹⁹F NMR (374 MHz, CDCl_3): δ = −139.9 ppm. HRMS (ESI) m/z : [M − F]⁺ calcd for $\text{C}_{22}\text{H}_{18}\text{BFNO}_2^+$ 358.1415; found 358.1409. These data matched those previously reported.¹⁹

Tris(4-(2,2-difluoro-6-methyl-2H-1λ³,3,2λ⁴-dioxaborinin-4-yl)phenyl)amine (A2). To a stirred solution of triphenylamine (1, 0.245 g, 1.0 mmol) in acetic anhydride (1.0 mL), $\text{BF}_3 \cdot \text{OEt}_2$ (0.5 mL) was added dropwise at 60 °C and maintained for 5 h. The reaction mixture color changed instantly to black during the addition of $\text{BF}_3 \cdot \text{OEt}_2$. The reaction was quenched with aqueous NH_3 solution 0.5 M and extracted with DCM \times 3. The crude product was purified by column chromatography in silica gel (DCM as eluent) to obtain A2 as an orange solid in a 52% yield. Mp: 210–211 °C. ¹H NMR (400 MHz, $\text{DMSO}-d_6$): δ : 2.45 (s, 9H), 7.23 (s, 3H), 7.36 (d, J = 8.5 Hz, 6H), 8.20 (d, J = 8.5 Hz, 6H) ppm. ¹³C NMR (101 MHz, $\text{DMSO}-d_6$): δ : 24.4 (CH_3), 97.9 (CH), 124.9 (CH), 126.4 (C), 131.3 (CH), 151.4 (C), 179.7 (C), 193.2 (C=O) ppm. ¹⁹F NMR (374 MHz, CDCl_3): δ = −137.8 ppm. HRMS (ESI) m/z : [M − F]⁺ calcd for $\text{C}_{30}\text{H}_{24}\text{B}_3\text{F}_5\text{NO}_6^+$ 622.1798; found 622.1816.

Conflicts of interest

The authors declare no competing financial interest.

Acknowledgements

We wish to thank the Departments of Chemistry and Physics and Vicerrectoría de Investigaciones at the Universidad de Los Andes for financial support. M. N.-P. and J. P. acknowledge support from the science faculty (projects INV-2020-105-2083 and INV-2019-84-1800). We also acknowledge Sandra Ortiz of Universidad de Los Andes for acquiring the mass spectra.

Notes and references

- 1 D. Cappello, R. R. Maar, V. N. Staroverov and J. B. Gilroy, *Chem.-Eur. J.*, 2020, **26**, 5522–5529.
- 2 J.-L. Jin, L. Yang, X. Ding, L.-H. Ou, Y.-D. Chen, H.-Y. Gu, Y. Wu and Y. Geng, *ACS Omega*, 2020, **5**, 21067–21075.
- 3 A. Filarowski, M. Lopatkova, P. Lipkowski, M. van der Auweraer, V. Leen and W. Dehaen, *J. Phys. Chem. B*, 2015, **119**, 2576–2584.
- 4 M. Más-Montoya, M. F. Montenegro, A. E. Ferao, A. Tárraga, J. N. Rodríguez-López and D. Curiel, *Org. Lett.*, 2020, **22**, 3356–3360.
- 5 V. Ramu, S. Gautam, P. Kondaiah and A. R. Chakravarty, *Inorg. Chem.*, 2019, **58**, 9067–9075.
- 6 X. Ren, F. Zhang, H. Luo, L. Liao, X. Song and W. Chen, *Chem. Commun.*, 2020, **56**, 2159–2162.
- 7 W.-J. Shi, Y.-F. Wei, C.-F. Li, H. Sun, L.-X. Feng, S. Pang, F. Liu, L. Zheng and J. Yan, *Spectrochim. Acta, Part A*, 2021, **248**, 119207.
- 8 H. Zhang, J. Xing, J. Peng, J. Bai, J. Zhang, D. Fu and J. Jia, *J. Lumin.*, 2022, **241**, 118525.
- 9 M. Pawlicki, H. A. Collins, R. G. Denning and H. L. Anderson, *Angew. Chem., Int. Ed.*, 2009, **48**, 3244–3266.
- 10 R. Medishetty, J. K. Zaręba, D. Mayer, M. Samoć and R. A. Fischer, *Chem. Soc. Rev.*, 2017, **46**, 4976–5004.
- 11 Q. Geng, C. Gu, J. Cheng and S. Chen, *Optica*, 2017, **4**, 674.
- 12 F. Bolze, S. Jenni, A. Sour and V. Heitz, *Chem. Commun.*, 2017, **53**, 12857–12877.
- 13 V. Juvekar, C. S. Lim, D. J. Lee, S. J. Park, G. O. Song, H. Kang and H. M. Kim, *Chem. Sci.*, 2021, **12**, 427–434.
- 14 Q. Geng, D. Wang, P. Chen and S.-C. Chen, *Nat. Commun.*, 2019, **10**, 2179.
- 15 X. Lou, Z. Zhao and B. Z. Tang, *Small*, 2016, **12**, 6430–6450.
- 16 S. Pascal, S. David, C. Andraud and O. Maury, *Chem. Soc. Rev.*, 2021, **50**, 6613–6658.
- 17 D. Beljonne, W. Wenseleers, E. Zoyer, Z. Shuai, H. Vogel, S. J. K. Pond, J. W. Perry, S. R. Marder and J.-L. Brédas, *Adv. Funct. Mater.*, 2002, **12**, 631–641.
- 18 M. Collot, *Mater. Horiz.*, 2021, **8**, 501–514.
- 19 D. Tamilarasan, R. Suhasini, V. Thiagarajan and R. Balamurugan, *Eur. J. Org. Chem.*, 2020, **2020**, 993–1000.
- 20 S.-L. Aranzazu, A. Tigreros, A. Arias-Gómez, J. Zapata-Rivera and J. Portilla, *J. Org. Chem.*, 2022, **87**, 9839–9850.
- 21 A. Tigreros and J. Portilla, *Eur. J. Org. Chem.*, 2022, **2022**, e202200249.
- 22 M.-C. Ríos, N.-F. Bravo, C.-C. Sánchez and J. Portilla, *RSC Adv.*, 2021, **11**, 34206–34234.



23 G. Delhumeau, A. M. Cruzmendoza and C. G. Lojero, *Toxicol. Appl. Pharmacol.*, 1994, **126**, 345–351.

24 H. Kaur and P. Singh, *Bioorg. Chem.*, 2019, **82**, 229–240.

25 R. Kaushik, A. Ghosh, A. Singh, P. Gupta, A. Mittal and D. A. Jose, *ACS Sens.*, 2016, **1**, 1265–1271.

26 S. Malkondu, S. Erdemir and S. Karakurt, *Dyes Pigm.*, 2020, **174**, 108019.

27 P. Xing, Y. Xu, H. Li, S. Liu, A. Lu and S. Sun, *Sci. Rep.*, 2015, **5**, 16528.

28 N. Bortey-Sam, R. Jackson, O. A. Gyamfi, S. Bhadra, C. Freeman, S. B. Mahon, M. Brenner, G. A. Rockwood and B. A. Logue, *Anal. Chim. Acta*, 2020, **1098**, 125–132.

29 A. Tigreros, M. Macías and J. Portilla, *Dyes Pigm.*, 2021, **184**, 108730.

30 A. Tigreros, M. Macías and J. Portilla, *Dyes Pigm.*, 2022, **202**, 110299.

31 A. Tigreros, J. Zapata-Rivera and J. Portilla, *ACS Sustainable Chem. Eng.*, 2021, **9**, 12058–12069.

32 S.-J. Chung, K.-S. Kim, T.-C. Lin, G. S. He, J. Swiatkiewicz and P. N. Prasad, *J. Phys. Chem. B*, 1999, **103**, 10741–10745.

33 Q. Zhang, P. Jiang, K. Wang, G. Song and H. Zhu, *Dyes Pigm.*, 2011, **91**, 89–97.

34 L. D. Costa, S. Guieu, J. Rocha, A. M. S. Silva and A. C. Tomé, *New J. Chem.*, 2017, **41**, 2186–2192.

35 S. Easwaramoorthi, P. Thamaraiselvi, K. Duraimurugan, A. J. Beneto, A. Siva and B. U. Nair, *Chem. Commun.*, 2014, **50**, 6902–6905.

36 L. Wu, J. Liu, P. Li, B. Tang and T. D. James, *Chem. Soc. Rev.*, 2021, **50**, 702–734.

37 V. Parthasarathy, S. Fery-Forgues, E. Campioli, G. Recher, F. Terenziani and M. Blanchard-Desce, *Small*, 2011, **7**, 3219–3229.

38 K. Li, X. Duan, Z. Jiang, D. Ding, Y. Chen, G.-Q. Zhang and Z. Liu, *Nat. Commun.*, 2021, **12**, 2376.

39 X. He, X. Wang, L. Zhang, G. Fang, J. Liu and S. Wang, *Sens. Actuators, B*, 2018, **271**, 289–299.

40 Y. Gao, M. Li, X. Tian, K. Xu, S. Gong, Y. Zhang, Y. Yang, Z. Wang and S. Wang, *Spectrochim. Acta, Part A*, 2022, **271**, 120882.

41 A. Chaicham, S. Kulchat, G. Tumcharern, T. Tuntulani and B. Tomapatanaget, *Tetrahedron*, 2010, **66**, 6217–6223.

42 S. Li, F. Huo, K. Ma, Y. Zhang and C. Yin, *New J. Chem.*, 2021, **45**, 1216–1220.

43 N. Yan, F. Wang, J. Wei, J. Song, L. Yan, J. Luo, Z. Fang, Z. Wang, W. Zhang and G. He, *Dyes Pigm.*, 2019, **166**, 410–415.

44 C. B. Malagon, M. Pellaton, A. Tigreros, J. Portilla, A. Valencia and M. N. Portela, in *OSA Nonlinear Optics 2021*, Optica Publishing Group, Washington, D.C., 2021, p. NTh3A.13.

45 C. Xu and W. W. Webb, *J. Opt. Soc. Am. B*, 1996, **13**, 481.

46 C. v. Bindhu, S. S. Harilal, G. K. Varier, R. C. Issac, V. P. N. Nampoori and C. P. G. Vallabhan, *J. Phys. D: Appl. Phys.*, 1996, **29**, 1074–1079.

47 M. G. Vivas, D. L. Silva, J. Malinge, M. Boujtita, R. Zalešny, W. Bartkowiak, H. Ågren, S. Canuto, L. de Boni, E. Ishow and C. R. Mendonca, *Sci. Rep.*, 2015, **4**, 4447.

48 C. Katan, F. Terenziani, O. Mongin, M. H. v. Werts, L. Porrès, T. Pons, J. Mertz, S. Tretiak and M. Blanchard-Desce, *J. Phys. Chem. A*, 2005, **109**, 3024–3037.

49 Y. Niko, S. Kawauchi and G. Konishi, *Chem.-Eur. J.*, 2013, **19**, 9760–9765.

50 C. Würth, M. Grabolle, J. Pauli, M. Spieles and U. Resch-Genger, *Nat. Protoc.*, 2013, **8**, 1535–1550.

51 J. Catalán, *J. Phys. Chem. B*, 2009, **113**, 5951–5960.

