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Synthesis and properties of quinoidal (di-)anionic coupled polymethine-oxonol dyes and their aromatic counterparts†

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We report the synthesis of a new family of chromophores based on benzothiazole, indandione, and tricyanofuran moieties, forming aromatic neutral precursors or negatively charged quinoidal counterparts. Upon deprotonation, these dyes undergo marked bathochromic shifts, leading to highly delocalized coupled polymethine structures, particularly in their moderatly antiaromatic dianionic forms. The evolution of optical properties was systematically studied through pH-dependent absorption measurements and supported by TD-DFT and CC2 calculations. Distinct behaviors were identified across the series: ESIPT in benzothiazole dyes, photoacidity in indandione derivatives, and spirocyclization in tricyanofurancontaining dyes. These findings provide insights into structure-property relationships in anionic polymethine systems.

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Introduction

Organic dyes capable of absorbing in the near-infrared (NIR) region are highly sought for modern applications, particularly in bio-imaging and molecular electronics. 1-5 Among NIR chromophores, polymethines are one of the most common families used for accessing such wavelengths, owing to their remarkable optical properties, which stem from the delocalization of an even number of π -electrons along an unsaturated chain composed of an odd number of sp²-hybridized carbon atoms. Symmetrical polymethines are typically synthesized by Knoevenagel condensations, in which two electron-donating (D) or electron-withdrawing (A) moieties react with an unsaturated dialdehyde. The resulting polymethine bridge enables charge delocalization between the terminal units, giving rise to different classes of polymethine dyes.⁶⁻⁹

Depending on the nature of the end groups, polymethines can be cationic (D- π -D, e.g. cyanines), neutral (D- π -A, e.g., merocyanines) or anionic (A- π -A, e.g., oxonols). While (mero) cyanines are well-documented in the literature, 6,7,10,11 and can readily achieve NIR absorption with straightforward

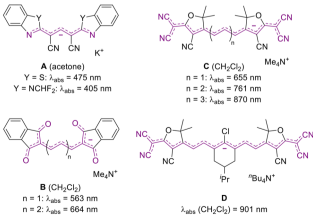


Fig. 1 Optical properties of previously reported anionic polymethines featuring the different end groups.

synthesis, 12-14 negatively charged polymethines remain relatively scarce. These dyes typically incorporate electron-withdrawing end groups such as benzothiazole (A),15 1,3-indandione (B),16-18 or tricyanofuran (C),19 which are among the most common building blocks in negatively charged polymethines (Fig. 1). Rigidified anionic polymethines have also dipyrromethene,20 designed introducing dioxaborine, 21,22 or a central cyclohexyl ring linking tricyanofurans (D), further redshifting their optical properties towards the NIR region.²³ Notably, negatively charged polymethine

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dyes have been explored in various applications, including light harvesting in solar cells, 24,25 nonlinear optics, 26-29 twophoton fluorescence bio-imaging, 30 sensing.31

In their heuristic 1966 study, Dähne and Leupold proposed the concept of coupled polymethines, in which two polymethine subunits are linked by one or more σ bonds. This coupling results in the hybridization of the key molecular orbitals, reducing the HOMO-LUMO gap, thereby inducing a bathochromic shift in the absorption spectrum.³² The validity of this principle was demonstrated in the same study through a comparison between the simple trimethine oxonol (Ox_3) , which exhibits an absorption maximum at 267 nm, and its coupled counterpart (DiOx₃), synthesized from 2,5-dihydroxybenzoquinone in basic media, which absorbs at 496 nm in aqueous sodium hydroxide (Fig. 2).32 Despite their intriguing optical properties, such coupled polymethines, and particularly negatively charged ones remain a relatively unexplored family of dyes.33

To date, a few dicationic coupled polymethines have been reported. These include the dicationic octamethyl-tetraaminobenzene studied by Staab and co-workers in 1986,34,35 acidochromic diaminobenzoquinonediiimine derivatives described by some of us, 36-38 as well as the more recently reported dicationic hexaaminophenazine, which absorbs in the NIR region.³⁹ Zwitterionic coupled polymethines, including a cationic and an anionic cyanine, have also been investigated, particularly derivatives of tetraazapentacene, 40,41 hexaazaanthracene, 42-44 or diaminobenzoquinonediimine incorporating electron-withdrawing groups on the nitrogen atoms, leading to intramolecular charge separation and unique solvatochromic properties. 45-47 Additional examples include zwitterionic aminobenzoguinonemonoinimine derivatives, 48,49 as well as the recently explored zwitterionic coupled heptamethine-oxonol systems (e.g., z-COPM, Fig. 2), which exhibited dual acidochromic behavior in both solution and thin films.⁵⁰

In contrast, dianionic coupled polymethines remain extremely rare. Beyond the previously mentioned DiOx3 in Dähne's seminal study, only a handful of examples have been described. Horner and Weber synthesized a derivative of 1,7-

2 Na⁺ 2 Na⁺ DiOx₃ DiOx₇ Ox₃ λ_{abs} (H₂O/NaOH) (MeOH/NaOH) λ_{abs} (H₂O/KOH) = 496 nm = 850 nm = 267 nmz-CoPM λ_{abs} (DMSO/DBU) = 885 nm

Fig. 2 Dianionic and zwitterionic coupled oxonol dyes and their key optical properties.

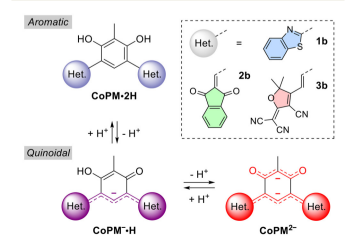
dihydoxy-naphthalene-2,6-dione, though its optical properties in basic media were not discussed in their work.⁵¹ Boldt reported a NIR-absorbing 2,6-dihydroxy-anthracene-1,5-dione dianion (DiOx₇, Fig. 2) in 1964,⁵² and Khodorkovsky observed a bis-indantetraone derivative in 1994.53 The structural resemblance between DiOx3 and z-CoPM motivated us to further investigate dianionic coupled polymethines using an analogous synthetic strategy. This approach aims to explore how increasing negative charge influences the stabilization, optical redshift, and photophysical properties of these systems, offering a meaningful comparison to more commonly studied zwitterionic and dicationic analogues.

In this work, we report the synthesis of novel mono- and bis-adducts of benzothiazole (1a, 1b), 1,3-indandione (2a, 2b), and tricyanofuran (3a, 3b) chromophores (Fig. 3). These dyes were successfully isolated and characterized, exhibiting dual acidochromic switching properties. Starting from neutral aromatic compounds (CoPM·2H), deprotonation leads to the formation of highly delocalized and redshifted anionic (CoPM⁻·H) and dianionic (CoPM²⁻) species with quinoidal sixmembered core. Additionally, the neutral dyes 1b and 2b were found to be emissive, with remarkably large Stokes shifts attributed to excited-state intramolecular proton transfer (ESIPT) in 1b and photoacid emission in 2b. Moreover, compound 3b revealed to undergo halochromic switching in basic medium, generating a spirocyclic isomer.

Results and discussion

Synthesis

The dialdehyde building block 4, synthesized through electrophilic formylation of 2-methylresorcinol,50 was used in condensation reactions with 2-aminothiophenol, 1,3-indandione or tricyanofuran 5 (Scheme 1). Initially, condensation of 4 with one equivalent of 2-aminothiophenol led to the precipitation of the pure mono-benzothiazole derivative 1a·2H in 70% yield. Under harsher conditions, i.e., boiling dimethylsulfoxide



Protonation states of the compounds investigated.

Scheme 1 Synthesis of chromophores 1–3 under their neutral or dianionic forms. Conditions: (i) 2-aminothiophenol, EtOH, 25 °C, 16 h; (ii) 2-aminothiophenol, DMSO, reflux, 30 min; (iii) 1,3-indandione, EtOH, pyridine, reflux, 16 h; (iv) 1,3-indandione, 2,6-lutidine, 95 °C, 16 h; (v) 5, MeOH, pyridine, reflux, 64 h; (vi) 5, 2,6-lutidine, 95 °C, 16 h.

(DMSO), the reaction afforded the bis-benzothiazole **1b-2H** with 81% yield.

Condensation of 4 with one equivalent of 1,3-indandione in boiling ethanol produced an orange precipitate identified as mono-condensed species 2a-2H, isolated in 53% yield after filtration. Its poor solubility suggested the need of a different solvent to facilitate a second condensation. Consequently, heating 4 with two equivalents of 1,3-indandione in 2,6-lutidine afforded the bis-indandione derivative 2b-2H with a rather modest 28% yield after precipitation in water.

Similarly, reacting 4 with one equivalent of tricyanofuran 5 at room temperature led to the precipitation of the pure monocondensed adduct 3a·2H with 33% yield. Attempts to obtain the corresponding bis-condensed product 3b·2H using two equivalents of 5 in boiling ethanol were unsuccessful, yielding only 3a·2H with a higher 66% yield. Switching to boiling methanol further improved the yield of 3a·2H to 84%. However, heating 4 with two equivalents of 5 in 2,6-lutidine at 95 °C successfully afforded the bis-condensed product 3b²⁻, isolated in 56% yield after precipitation in diethyl ether. The dianionic structure of 3b2- was confirmed by 1H NMR, which showed signals corresponding to two equivalents of 2,6-lutidinium cations (see Fig. S24, ESI†), and HRMS analysis in methanol, which detected both dianionic (z = 2) and anionic (z = 2) = 1) species, indicating partial protonation in this protic solvent (refer to Fig. S35, ESI†).

Notably, solubilizing compounds 1a·2H, 1b·2H, 2a·2H, 2b·2H, or 3a·2H in polar solvents (e.g., methanol), lead to remarkable color changes, prompting further investigation into the acidochromic and solvatochromic properties of these dyes and the impact of their protonation states on photophysical properties.

Photophysical properties

To investigate the optical properties of the synthesized chromophores, their electronic absorption spectra were recorded in DMSO. Acidification and basification were achieved using 0.1 M trifluoroacetic acid (TFA) or 1,8-diazabicyclo[5.4.0]undec-7-ene (DBU), respectively. The corresponding spectra (Fig. 4) reveal significant variations of the absorption properties depending on the solution conditions: starting from neutral species in acidic media, spontaneous deprotonation occurs in DMSO, which can be further promoted under basic conditions.

The absorption solvatochromism of the six dyes was examined in dichloromethane (DCM), methanol, and DMSO (Fig. S43-S45, ESI†). In these solvents, absorption profiles are primarily influenced by the acidic (e.g., DCM), or basic (e.g., DMSO) nature of the medium, which determines the predominance of either the neutral or deprotonated forms. To gain deeper insight into the protonation states of the species in solution, experimental absorption data were compared with the results of theoretical simulations. The computed spectra were obtained from vibrationally-resolved TD-DFT calculations using the cLR2 solvation model (DMSO),54 from which the wavelengths corresponding to the various peaks were extracted. Additional methodological details can be found in the ESI.† In general, at the selected level of theory, a blueshift with respect to experimental results of approximately 30 to 100 nm is expected for cyanine compounds when using TD-DFT.55 Improved estimates of absorption energies using CC2 corrections for the neutral and monoanionic species, are reported in Table S1 (ESI).† However, as outlined in the ESI,† such corrections could not be applied to the dianionic forms

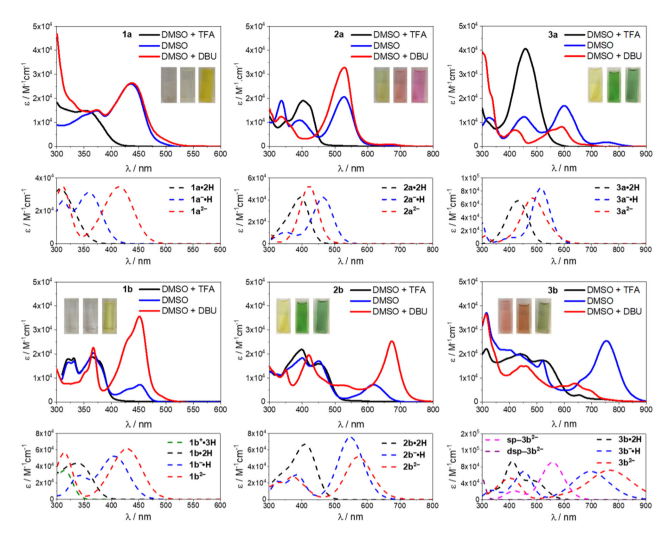


Fig. 4 UV-vis-NIR electronic absorption spectra of mono-condensed (top row) and bis-condensed dyes (bottom row) dissolved at $ca. 2 \times 10^{-5}$ M in acidic (0.1 M TFA, black line), neutral (blue line) or basic (0.1 M DBU, red line) DMSO (with corresponding pictures of the solutions presented in this order from left to right). Below each experimental UV-vis-NIR electronic absorption spectra are presented the corresponding theoretical vibrationally-resolved absorption computed for each relevant protonation state (dashed lines).

due to difficulties in reliably identifying the corresponding excited states. Theoretical calculations also provide a qualitative estimate of the molar absorptivity (ε) , which reflects the absorption intensity. Although the absolute intensities are significantly overestimated by theory in the present case, the relative trends between peaks are well reproduced. In short, the compiled data (Table 1) provide a basis for correlating experimental observations with theoretical predictions, aiding identifying the dominant protonation states in different conditions.

First, in the case of the neutral mono-condensed adduct of benzothiazole (1a·2H), the absorption maximum in acidic conditions appears in the UV region at 355 nm with a molar absorptivity of ca. $1.5 \times 10^4 \text{ M}^{-1} \text{ cm}^{-1}$. In neutral and basic DMSO, a bathochromic shift occurs towards the blue region at 435 nm, with a nearly doubled ε reaching $2.8 \times 10^4 \text{ M}^{-1} \text{ cm}^{-1}$. This shift likely corresponds to the formation of the deprotonated species (1a-H or 1a²⁻). Theoretical calculations predict a vertical transition at 413 nm for 1a²⁻, which is underestimated but generally aligns with the experimental data. In contrast, assigning the experimental 435 nm band to 1a-H would come with a 0.63 eV error from theory, deemed as large, even for TD-DFT.

Similarly, the mono-indandione derivative (2a·2H) follows a comparable trend, exhibiting a bathochromic shift from 402 nm in acidified DMSO to 527 nm in neutral and basic conditions. This experimental redshift upon deprotonation is also reproduced by theoretical calculations, which predict an absorption maximum at 463 nm for 2a-H. As the solution transitions from neutral to basic DMSO, ε^{527} increases from ca. $2.1 \times 10^4 \text{ M}^{-1} \text{ cm}^{-1}$ to $3.3 \times 10^4 \text{ M}^{-1} \text{ cm}^{-1}$, corresponding to the complete conversion to anion 2a-H. In such basic condition, the dianion species 2a²⁻ is not observed, since its formation should induce a small (non-observed) redshift compared to 2a·2H, according to theory.

In contrast, the mono-condensed tricyanofuran adduct (3a·2H) displays a distinct behavior. The protonated form 3a·2H absorbs at 459 nm, but in neutral DMSO, deprotonation

Table 1 Experimental electronic absorption and the theoretical absorption for given species in DMSO estimated from spectra in Fig. 4, with NICS values as estimates of the relative aromaticity of the central ring of these systems

Experime	ntal		Theoretical			
Dyes	Conditions ^a	$\lambda_{\max}^{\exp} [nm] (\varepsilon [M^{-1} cm^{-1}])$	Species	$\lambda_{\rm abs}^{\rm theo\ \it b}$ [nm] (ε [M ⁻¹ cm ⁻¹])	NICS(0) [ppm]	
1a	DMSO/TFA	355 (14 700)	1a∙2H	306 (33 400)	-7.16	
		282 (28 000)		264 (44 800)		
	DMSO	436 (26 000)	1a⁻∙H	$358 (31 \ 100)^c$	-2.48^{c}	
		374 (14 900)		$311(26200)^{c}$		
	DMSO/DBU	438 (26 400)	1a ²⁻	413 (34 700)	0.33	
		374 (15 200)		310 (34 900)		
1b	DMSO/TFA	378 (sh, 17 000)	1b ⁺ ⋅3H	313 (35 100)	-6.48	
		364 (18 800)	1b•2H	337 (43 900)	-6.87	
		331 (18 300)		274 (46 400)		
	DMSO	452 (7200)	1b¯·H	404 (52 400)	-2.21	
		430 (sh, 5400)		342 (29 200)		
		367 (20 500)		,		
	DMSO/DBU	451 (35 600)	$1b^{2-}$	428 (61 500)	-0.29	
		430 (sh, 26 400)		313 (56 400)		
		367 (22 400)		,		
2a	DMSO/TFA	402 (19 000)	2a-2H	395(42 500)	-5.49	
		350 (sh, 10 700)				
	DMSO	527 (20 600)	2a⁻-H	463 (42 400)	0.06	
		391 (10 900)		349 (10 500)		
		336 (18 900)		()		
	DMSO/DBU	529 (33 000)	$2a^{2-}$	420 (52 500)	3.60	
		336 (12 300)		296 (11 200)		
		(-2 333)		228 (3400)		
2b	DMSO/TFA	452 (16 000)	2 b ⋅2H	409 (67 000)	-3.80	
-~		398 (21 900)		340 (32 100)		
	DMSO	617 (7400)	2b [−] ·H	546 (75 500)	1.80	
		452 (17 100)		383 (29 900)		
		400 (18 200)		328 (22 300)		
	DMSO/DBU	674 (25 400)	$2b^{2-}$	571 (51 800)	4.80	
		534 (7000)		371 (27 900)	-100	
		421 (19 700)		336 (20 766)		
3a	DMSO/TFA	459 (40 800)	3a-2H	$429 (65 600)^c$	-7.40^{c}	
ou	21.100,1111	288 (21 200)	ou 211	123 (66 666)	7.120	
	DMSO	755 (1700)	3a⁻-H	$512 (84 600)^c$	-1.17^{c}	
	21.150	600 (17 000)	ou 11	$297 (32 600)^c$	111,	
		453 (12 400)		237 (02 000)		
	DMSO/DBU	595 (8300)	$3a^{2-}$	484 (69 700) ^c	2.96^{c}	
	DI IOO/DBC	420 (7200)	ou.	101 (03 7 00)	2.50	
3b	DMSO/TFA	523 (17 300)	3 b ⋅2H	$479 (47 000)^c$	-6.20	
36	DINISO/ IIII	441 (20 000)	00 211	$409 (93 300)^c$	0.20	
		111 (20 000)		$349 (25 200)^c$		
	DMSO	754 (25 500)	3b [−] ·H	697 (68 700) ^c	0.50	
	Diviso	660 (sh, 9100)	0 0 11	$451 (68 800)^c$	0.00	
		521 (17 000)		$335 (23 400)^c$		
	DMSO/DBU	690 (sh, 5000)	$3b^{2-}$	765 (71 800) ^c	4.40	
	DIVIDO/DB0	638 (7800)	36	$403 (52 300)^c$	4.40	
		591 (6800)		100 (02 000)		
		331 (0000)	sn-3h ²⁻	556 (91 100) ^c	-1.37	
			sp-3b ²⁻ dsp-3b ²⁻	$417 (21700)^c$	1.07	
			dsn_3h2-	296 (55 100)	-7.65	
			usp—sv	250 (33 100)	-7.03	

^a Pure DMSO or mixture of DMSO + 0.1 M TFA or DBU. ^b Theoretical absorption maxima and absorptivity coefficients determined in methanol. ^c Computed on the most stable isomer. sh: shoulder.

presumably leads to $3a^-\cdot H$, which shows a redshifted absorption at 600 nm ($\varepsilon^{600}=1.7\times10^4~M^{-1}~cm^{-1}$), accompanied with a weaker low-intensity band at 755 nm ($\varepsilon^{755}=0.2\times10^4~M^{-1}~cm^{-1}$). In basic DMSO, a blueshifted band at 595 nm emerges, suggesting the formation of the dianionic species $3a^{2-}$. This is corroborated by the calculations, which also predict a noticeable blueshift (from 512 to 484 nm) when going from the anionic species $3a^{-\cdot}H$ to the dianionic species $3a^{2-}$.

The study of bis-condensed adducts provides striking results, with their symmetrical structures favoring the formation of highly delocalized charged species, and with the establishment of coupled polymethine systems inducing pronounced spectral shifts. The bis-benzothiazole derivative (1b·2H) appears colorless in acidic solution, with absorption in the UV region at 365 nm. Titration of 1b with TFA did not evidenced the formation of the cationic species $1b^+\cdot 3H$

(Fig. S46†), for which an absorption maximum is predicted at 313 nm. In neutral DMSO, a band emerges at ca. 450 nm, intensifying in basic conditions to reach $\varepsilon^{451} = 3.6 \times 10^4 \text{ M}^{-1} \text{ cm}^{-1}$. As observed for the mono-benzothiazole analogue (1a·2H), this evolution suggests the formation of the deprotonated species (1b⁻·H or 1b²) in neutral and basic solutions, with theoretical calculations predicting absorption maxima at 404 and 428 nm for 1b⁻·H and 1b², respectively. It is noteworthy that the experimental wavelengths are comparable to the anionic unrigidified benzothiazole-based analogue **A**, which presents an absorption maximum at 475 nm (Fig. 1). ¹⁵

For the bis-indandione derivative 2b·2H in acidic solution, the absorption spectrum exhibits two maxima at 452 (ε = 1.60 $\times 10^4 \text{ M}^{-1} \text{ cm}^{-1}$) and 396 nm ($\varepsilon = 2.19 \times 10^4 \text{ M}^{-1} \text{ cm}^{-1}$). In neutral DMSO, these bands persist, but an additional transition emerges at 617 nm, which we attribute to the presence of deprotonated species 2b-H, in agreement with theoretical predictions, anticipating a transition at 546 nm for this form. In basic conditions, an intense bathochromic shift occurs, with a new band at 674 nm ($\varepsilon = 2.5 \times 10^4 \text{ M}^{-1} \text{ cm}^{-1}$), a wavelength comparable to that of the bis-indandione anionic pentamethine analogue B, which absorbs at 664 nm (Fig. 1).17 Theoretical calculations suggest that further deprotonation leads to increased redshifts, supporting the assignment of this band to the dianionic species (2b²⁻). Interestingly, in basified methanol, the absorption is blueshifted to 622 nm (Fig. S44, ESI†), likely due to the hydrogen bonding between the solvent and the multiple oxygen atoms of the dye promoting the monoanionic species 2b-H.

Finally, the bis-tricyanofuran derivative 3b·2H shows two broad absorption bands at 523 ($\varepsilon = 1.73 \times 10^4 \text{ M}^{-1} \text{ cm}^{-1}$) and 441 nm ($\varepsilon = 20.0 \times 10^4 \text{ M}^{-1} \text{ cm}^{-1}$) in acidic conditions. In neutral DMSO, a strong band develops in the NIR range, peaking at 754 nm ($\varepsilon = 2.6 \times 10^4 \text{ M}^{-1} \text{ cm}^{-1}$). This transition likely corresponds to either the anionic species 3b-H or its dianionic counterpart 3b2-, for which theoretical absorption maxima are predicted ca. 700 and 760 nm, respectively. The absorption wavelength (754 nm) is notably smaller than the 900 nm absorption maximum of the anionic heptamethine analogue D (Fig. 1). This difference is likely due to the destabilizing effect of the oxonol subunit on the negatively charged polymethine system, combined with the s-cis conformation adopted by the heptamethine bridge, which may hinder linear charge delocalization. In basic conditions, a noticeable blueshift of absorption occurs for 3b²⁻, a phenomenon discussed in detail in a subsequent section.

Computational study

A conformational analysis of the possible isomers and tautomers of dyes **1b**, **2b**, and **3b**, including their corresponding neutral, mono- and dianionic forms, was carried out using DFT calculations to identify the most thermodynamically stable structures at room temperature (see Fig. S50, ESI†). In few cases, the analysis revealed multiple conformers with relative Gibbs free energies (ΔG) within 2.0 kcal mol⁻¹ of the most stable geometry; these alternatives structures are reported in

Fig. S51–S58, ESI.† For all neutral species of 1 and 2 series and their corresponding monoanionic counterparts, the analysis of the tautomers indicates that the hydrogen atoms are preferentially bonded to the oxygen atoms of the central six-membered ring instead of the benzothiazole and indandione moieties.

Both **1b·2H** and **2b·2H** exhibit a single, stable, and planar structure in which the phenolic hydrogen atoms engage in intramolecular hydrogen bonding, either with the nitrogen atoms of the benzothiazole (**1b·2H**) or with the oxygen atoms of the **1,3-indandione** fragments (**2b·2H**), as shown in Fig. S50 (ESI).† Under basic conditions, deprotonation disrupts one of these hydrogen bonds, leading to a **180°** rotation of the corresponding functional group in **2b**-·**H**, whereas **1b**-·**H** retains its initial geometry. In the dianionic forms, the loss of both hydrogen bonds induces a rotation of both peripheral units in **1b**²⁻ and **2b**²⁻.

In contrast, the structure of **3b·2H** displays notable conformational flexibility at room temperature, with eight conformers lying within 1.22 kcal mol^{-1} of the global minimum. These arise from rotations of the two tricyanofuran moieties around the C–C bonds of the polymethine chain connecting them to the central core (see Fig. S55, ESI†). Although these rotations have minimal impact on the nature of the principal electronic transitions, they cause minor shifts in the calculated absorption maxima. To account for this, the relevant absorption wavelengths are reported as room-temperature Boltzmann-weighted averages. A similar degree of conformational freedom is observed for both the monoanionic and dianionic forms of **3b**, which respectively exhibit seven and four stable conformers within a 2 kcal $\text{mol}^{-1} \Delta G$ window (see Fig. S42 and S43, ESI†).

To assess the aromatic or quinoidal nature of the central core of the chromophores depending on their protonation state, nucleus independent chemical shifts (NICS) were computed (Table 1). As anticipated for the symmetrical dyes, the neutral forms exhibit negative NICS(0) values of -6.87, -3.80 and -6.20 ppm for $1b\cdot 2H$, $2b\cdot 2H$, and $3b\cdot 2H$, respectively, indicating a predominantly aromatic character. Upon single deprotonation, the NICS increase significantly for $1b^-\cdot H$, $2b^-\cdot H$, $3b^-\cdot H$, reaching -2.21, 1.80 and 0.50 ppm, respectively, consistent with a progressive loss of aromaticity in favor of a more quinoidal structure. In the fully deprotonated dianionic forms, this trend continues: $1b^{2-}$ exhibits a nearly non-aromatic character (NICS = -0.29 ppm), whereas both $2b^{2-}$ and $3b^{2-}$ display positive NICS of 4.80 and 4.20 ppm, respectively, hinting at a moderately antiaromatic character of the central ring.

Theoretical calculations predict a progressive bathochromic shift in the absorption energies of all three symmetrical dyes upon sequential deprotonation, from the neutral to anionic and then to the dianionic forms. This trend is consistent with the increasing negative charge, which promotes a quinoidal electronic structure, enhancing conjugation along the two coupled polymethine backbones (Fig. 3). In these species, the two polymethine subunits are connected *via* single bonds, facilitating charge delocalization across the entire system. This delocalization gives rise to cyanine-like electronic transitions, in line with the coupling principle discussed earlier.

Research Article

Excited-state calculations attribute the long-wavelength absorption bands of the dyes to HOMO → LUMO transitions, each characterized by high oscillator strengths (see Tables S1, ESI†). Electron density difference (EDD) maps between the ground (S₀) and first excited (S₁) states for neutral 1b, 2b, 3b, and their corresponding anionic and dianionic forms provide further insight into the nature of these electronic excitations (Fig. 5). In the neutral species, the EDD maps reveal pronounced intramolecular charge transfer (ICT) predominantly from the phenolic functions towards the electron-withdrawing moieties. The density redistribution is centered on the chromophoric core and extends along the substituted polymethine chain, with partial involvement of the oxygen atoms attached to the upper methine bridges. In particular, for 3b·2H, the extended conjugation and the stronger electron-withdrawing groups enhance delocalization, resulting in a more extensive ICT character.

In the anionic forms, the EDD maps for the lowest excited state indicate a broader charge delocalization across the molecules compared to the neutral species. However, the presence of a negative charge on the oxygen of the oxonol subunit imparts a quinoidal character on the central core, which localizes the donor group involved in the charge transfer primarily around the oxygen and diminishes the contribution from other parts of the molecule. In the dianionic species, the aromatic character is lost, and the EDD becomes predominantly localized on the lower polymethine chain with only minor contributions from the oxygen atoms of the oxonol subunit.

Spirocyclization of dye 3

Unexpectedly, when $3b^{2-}$ is dissolved in basic DMSO, a significant hypsochromic effect occurs, shifting the absorption to the 600-700 nm range. This suggests that the dianionic species undergoes structural reorganization, likely evolving into a lessconjugated spirocyclic chromophore sp-3b2- and dsp-3b2-, depicted in Scheme 2. Similar behavior has been previously observed in simpler tricyanofuran-based systems. 56-60 Theoretical calculations reveal that the cyclization of one tricyanofuran group leads to two stable isomers, depending on whether the second tricyanofuran unit is rotated 0° or 180° relative to the central ring plane. These sp-3b2- isomers are 7.1 and 8.4 kcal mol^{-1} more stable than the open form $3b^{2-}$, respectively (see Fig. S58, ESI†). Dual cyclization involving both tricyanofuran units leads to the putative dsp-3b²⁻ struc-

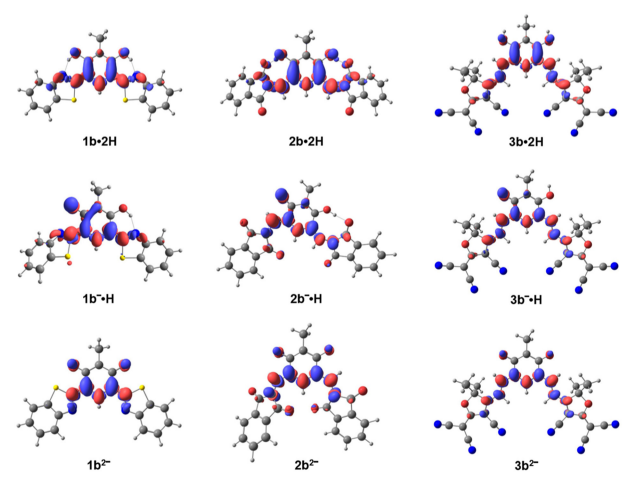


Fig. 5 Electronic density differences (EDD) between the ground (S₀) and the first excited state (S₁) for molecules 1b, 2b and 3b (most stable isomers) under different protonation states. Regions in red indicate an increase in electronic density from S₀ to S₁, while regions in blue indicate a decrease. The isosurfaces are visualized with an isovalue of 0.003.

Scheme 2 Spirocyclization of 3b occurring in basic conditions.

ture, which is even more stable, with an energy gain of 13.6 kcal mol⁻¹ compared to the open form. These data indicate that the spirocyclic forms are thermodynamically favored. However, their presence in solution may be limited by a high activation barrier associated with the cyclization process from sp-3b²⁻ to dsp-3b²⁻. The sp-3b²⁻ isomer exhibits a bright electronic transition at 556 nm (see Table 1), whereas for dsp-3b²⁻, the lowest-energy transition is predicted around 300 nm, consistent with a loss of extended π -conjugation. Interestingly, the spirocyclic forms sp-3b²⁻ and dsp-3b²⁻ show contrasting NICS values of -1.37 and -7.65 ppm, reflecting their quinoidal and aromatic central cores.

The formation of sp-3b²⁻ was also supported by ¹H NMR spectroscopy upon addition of Cs₂CO₃, which revealed the appearance of new doublet signals corresponding to the heptamethine protons (Fig. S29, ESI†). These consist of two distinct pairs of doublets, with coupling constants (^3J) of 10.0 Hz and 16.1 Hz, attributed to the cis and trans configurations of the protons on the closed and open sides of the molecule, respectively.

Emission properties

The emission properties of the six chromophores (1a,b, 2a,b, and 3a,b) were investigated in DMSO, with TFA or DBU added as needed to generate either the neutral or charged species. Fluorescence was observed for the bis-benzothiazole dye (1b) and bis-indandione dye (2b) only, whose emission properties are summarized in Table 2.

The neutral dye 1b·2H exhibits a yellow fluorescence in acidic DMSO, peaking at 546 nm with a low quantum yield of 0.2% and a notably large Stokes shift of 9100 cm⁻¹ (Fig. 6). The origin of this emission was confirmed by its excitation spectrum, which closely matches the absorption profile of **1b·2H** (Fig. S47, ESI†). Benzothiazole-substituted *o*-phenols are known to undergo excited-state intramolecular proton transfer (ESIPT), leading to significant Stokes shifts. 61-65 Thus, we hypothesized that 1b-2H undergoes ESIPT and computed its emission from both the enol (EE-1b-2H, local state excitation) and keto forms (KE-1b-2H, mono-ESIPT process). These excited-state calculations indicate a favorable ESIPT, with a ΔG of -0.27 eV between the excited enol and keto tautomers (Fig. 7). Additionally, the predicted fluorescence maximum for KE-1b-2H at 523 nm aligns well with the experimental value $(\lambda_{\rm em}^{\rm exp} = 546 \text{ nm})$, whereas the emission from the enol tautomer (EE-1b·2H) theoretically expected at 404 nm does not fit the experiments.

The possibility of double proton transfer was also investigated, and calculations revealed a favorable ΔG of -0.29 eV for the transition from the excited (mono) keto form (KE-1b·2H) to the doubly proton-transferred form (KK-1b·2H). This species is predicted to emit at 860 nm with a negligible oscillator strength, indicating an emission in the NIR range with an intensity (radiative rate) too low to be detected with our current setup. Note that the presence of a second ESIPT process is consistent with the very low quantum yield observed (0.2%).

Upon deprotonation in DMSO in the presence of DBU, dye 1b displays a blue emission at 498 nm, with a significantly reduced Stokes shift of 1900 cm⁻¹ and a markedly higher fluorescence quantum yield of 11%. This emission aligns well with theoretical predictions, which estimate fluorescence maxima

Table 2 Emission properties of compounds 1b and 2b. Theoretical estimates of energies are obtained through the cLR² model, while the data in parenthesis correspond to the emission energies corrected with CC2 approach (if available) as explained in the computational details

Experimental								Theoretical	
Dyes	Conditions ^a	$\lambda_{\max}^{b} [nm]$	λ _{em} [nm]	Stokes shift [cm ⁻¹]	Φ [%]	T[ns]	Species	λ _{em} ^{theo} [nm]	
1b	DMSO/TFA	365	546	9100	0.2	3.2	EE−1b·2H KE−1b·2H	404 486 (526)	
	DMSO/DBU	454	498	1900	11	2.3	1b [−] ·H 1b ^{2−}	415 481	
2b	DMSO/TFA	460	704	7500	2	1.1	2 b ·2H	623	
	DMSO	633	705	1600	2	1.0	2b ⁻ ∙H 2b ²⁻	577 (681) 596	

^a Pure DMSO or mixture of DMSO + 0.1 M TFA or DBU. ^b Absorption value extracted from the maximum of the recorded excitation spectra (Fig. S47, ESI†).

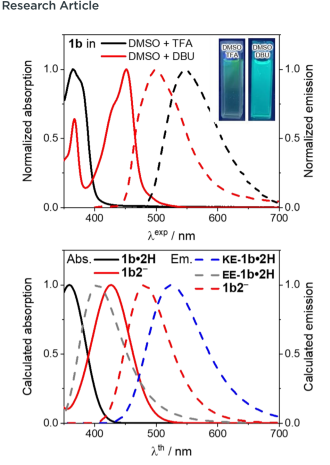


Fig. 6 Experimental (top) and theoretical (bottom) absorption (plain lines) and emission (dotted lines) spectra of 1b.

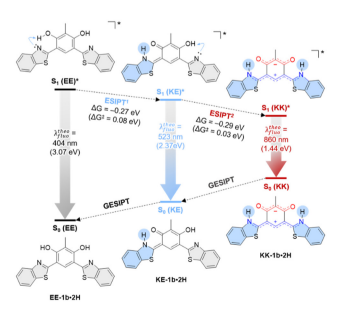


Fig. 7 Computed differences of free energies (ΔG) between tautomers of 1b-2H (1st and 2nd ESIPT) at the first excited state and computed emissions

at 415 nm for the monoanionic species (1b⁻⋅H) and 481 nm for the dianionic species (1b²⁻) with a large oscillator strength of 1.09 and 0.60, respectively.

Regarding bis-indandione derivative 2b·2H, fluorescence is observed in the red-NIR region, peaking at 704 nm with a rather large Stokes shift of 7500 cm⁻¹ (Fig. 8). Surprisingly, the deprotonated species 2b-H exhibits an almost identical emission profile, with comparable fluorescence quantum yield of 2% and lifetime of ca. 1 ns (Fig. S48, ESI†). The excitation spectra of both 2b·2H and 2b-H closely match their respective absorption profiles (Fig. S47, ESI†), demonstrating that both species relax to the same emissive state.

This observation suggests that 2b·2H behaves as a photoacid (Fig. 9); upon excitation, its acidity is significantly enhanced, resulting in a proton release and the formation of the excited anionic species 2b-H, which subsequently relaxes via red light emission. This behavior is reminiscent of known ESIPT fluorophores, such as benzazole-based dyes, which act as photoacids in neutral DMSO.64 Theoretical calculations support this interpretation, indicating that the 2b·2H form undergoes a barrierless ESIPT process, as the excited-state geometry yields a spontaneous proton transfer, likely preceding the proton dissociation described above. The emission spectrum of the keto tautomers closely resembles those of the

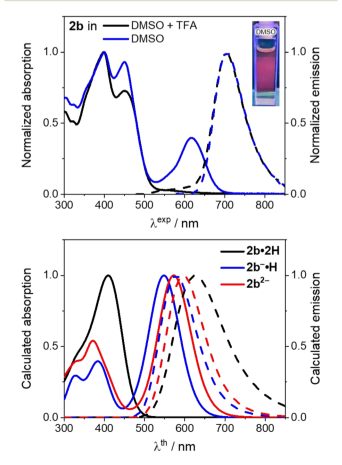


Fig. 8 Experimental (top) and theoretical (bottom) absorption (plain lines) and emission (dotted lines) spectra of 2b.

Proposed photoacidic process occurring upon irradiation of 2b-2H.

anionic and dianionic species, with all three peaks appearing between ca. 600-700 nm (see Fig. 9 and Table 2). However, because these spectral differences lie within the typical error bar of TD-DFT calculations, the experimental emission near 700 nm cannot be unambiguously assigned to one specific structure.

Conclusions

In this study, we reported the single-step synthesis of six chromophores built around a tetra-substituted six-membered ring incorporating benzothiazole, indandione, or tricyanofuran moieties. The dyes containing indandione and tricyanofuran moieties exhibit visible coloration in solution, attributed to their tendency to spontaneously deprotonate and form anionic species. We systematically explored their optical properties, with a particular focus on their absorption, emission, and protonation-dependent behavior. Experimental and theoretical investigations revealed that deprotonation of the neutral symmetrical dyes, initially featuring aromatic cores, induces pronounced bathochromic shifts, stemming from the formation of highly delocalized oxonol-polymethine structures with a quinoidal six-membered core. Notably, the bis-tricyanofuran derivative (3b) reached NIR absorption in solution and underwent further transformation via spirocyclization under basic conditions, yielding a distinct optical signature.

Fluorescence studies revealed that the bis-benzothiazole (1b) and bis-indandione (2b) derivatives exhibit emission. The neutral bis-benzothiazole dye undergoes an ESIPT process, resulting in an exceptionally large Stokes shift, which was further validated through theoretical modeling. Additionally, the bis-indandione dye exhibits a photoacidic behavior, undergoing excited-state deprotonation to generate an emissive anionic species. These findings provide deeper insights into the complex electronic properties of coupled polymethines and anionic dyes while contributing to the development of functional chromophores with tunable absorption and wellfluorescence properties suited defined specific applications.

Author contributions

Conceptualization: S. P.; investigations: B. M., C. N., S. P.; D. J. supervision: S. P., D. J., O. S.; writing - original draft: B. M., S. P., C. N.; writing - review & editing: B. M., C. N., D. J., O. S., S. P.

Conflicts of interest

There are no conflicts to declare.

Data availability

The data supporting this article have been included as part of the ESI.†

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