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Synthesis, and the optical and electrochemical properties of a series of push–pull dyes based on the 4-(9-ethyl-9*H*-carbazol-3-yl)-4-phenylbuta-1,3-dienyl donor†

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A series of twelve dyes based on the 4-(9-ethyl-9*H*-carbazol-3-yl)-4-phenylbuta-1,3-dienyl donor were prepared with electron acceptors varying in their structures but also in their electron-withdrawing ability. For specificity, a butadienyl spacer was introduced between the donor and the acceptor to both lower the bandgap and furnish dyes with high molar extinction coefficients. The different dyes **A–N** were characterized using various techniques including UV-visible absorption and fluorescence spectroscopy, and cyclic voltammetry. All dyes showed an intense intramolecular charge transfer band located in the visible range. To further investigate the optical properties of the twelve dyes, their solvatochromism was investigated in twenty-three solvents of different natures, enabling linear correlations to be obtained on different polarity scales such as the Taft, Reichardt and Catalan scales. To support the experimental results, the optical properties were compared with those theoretically determined.

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A Introduction

During the past decades, organic dyes have attracted a great deal of interest from both the academic and industrial communities due to the numerous applications in which these structures can be used. Notably, organic dyes can be employed as singlet or triplet emitters for organic light-emitting diodes,^{1–3} as light-absorbing materials for organic photovoltaics (OPVs),^{4–6} as chromophores for non-linear optical applications,^{7–9} as dyes for various biological applications including cancer phototherapy or biological labelling,^{10,11} as photoinitiators of polymerization,^{12–20} and as photoredox catalysts for organic transformations or hydrogen production.^{21–24} Recently, significant efforts have been devoted to replace organic dyes by push–pull dyes in photopolymerization because the absorption spectra of push–pull dyes are easy to tune.^{25–32} For many of the abovementioned applications, the possibility to design metal-free dyes is especially attractive, addressing the toxicity issue often raised by the use of transition metal

complexes.^{33–36} For the design of organic molecules strongly absorbing in the visible range and displaying molar extinction coefficients on par with those of the transition metal complexes, push–pull dyes incorporating an electron donor connected to an electron acceptor by means of a π -conjugate spacer are the most obvious candidates and the most straightforward route to get molecules with high molar extinction coefficients.^{37–39} Indeed, the position of the intramolecular charge transfer (ICT) band can be easily tuned by modifying both the strength of the electron donating and the electron accepting abilities of the two groups attached at both ends of the π -conjugated spacer.^{40–43} This strategy is notably extremely useful to red-shift the ICT bands and thus reduce the HOMO–LUMO gap (where HOMO and LUMO respectively stands for highest occupied molecular orbital and lowest unoccupied molecular orbital).^{44–47} Additionally, the molar extinction coefficient can be drastically increased by elongating the π -conjugated spacer between the donor and the acceptor as a result of an improvement of the oscillator strength.^{47–49} However, elongation of the π -conjugation in these structures is not an easy task, as exemplified with the synthesis of 3-(9-alkyl-9*H*-carbazol-3-yl)acrylaldehydes. Indeed, for this aldehyde, only three syntheses are reported in the literature.^{50–54} In fact, carbazole is a remarkable electron donor exhibiting a relatively low oxidation potential, and good thermal stability and this polyaromatic structure can also be easily chemically modified.^{53,54} Carbazole is also a cheap compound

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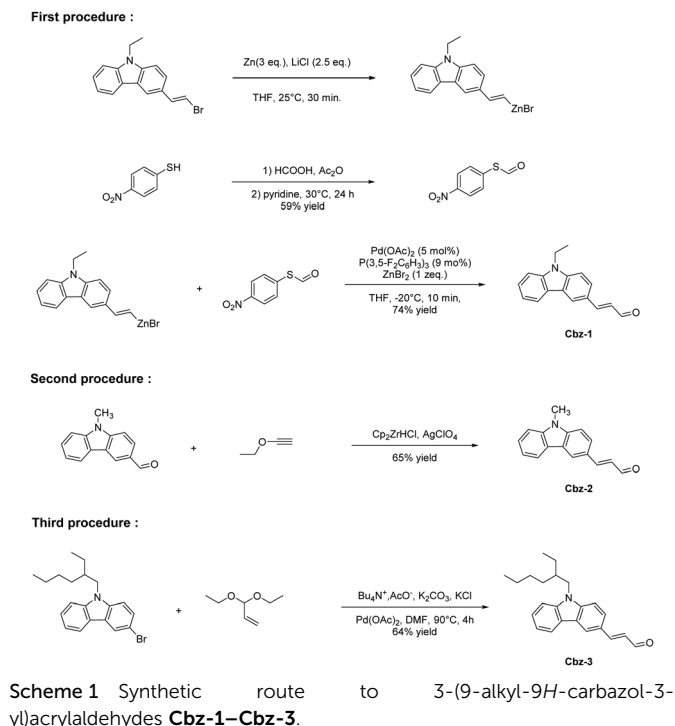
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and so this structure was extensively used in Organic Electronics, with applications ranging from fluorescence emitters to host materials for OLEDs.^{55–58}

While coming back to the synthesis of 3-(9-alkyl-9H-carbazol-3-yl)acrylaldehydes, the three procedures reported in the literature are based on metal-catalysed syntheses, namely the palladium-catalysed formylation of an alkenylzinc intermediate using *S*-(4-nitrophenyl)thioformate to introduce the aldehyde group for the synthesis of **Cbz-1**,⁵⁹ the hydrozirconation homologation method of Maeta and Suzuki for the synthesis of **Cbz-2**⁶⁰ or the Pd-catalysed coupling reaction of acrolein diethyl acetal on a brominated carbazole for the synthesis of **Cbz-3**^{61–63} (see Scheme 1). If large scale syntheses are required, these reactions are not adapted, requiring the use of multistep syntheses taking recourse to expensive catalysts, dry solvents and imposing a controlled atmosphere to proceed. Simpler synthetic methods are thus actively being researched. Recently, several studies were devoted to elaborate acrolein derivatives starting from benzophenone derivatives. This strategy was notably successfully applied to the synthesis of Michler's aldehyde starting from Michler's ketone.^{64,65} More recently, new benzophenone derivatives were also converted to their aldehyde analogues,^{66–71} furnishing dyes exhibiting a strong positive solvatochromism.^{72,73}

In light of these results, a hybrid carbazole-based benzophenone **Cbz-BP** in which the electron donating ability is reinforced by the presence of the carbazole moiety has been converted to its aldehyde analogue **Cbz-3**. The approach used to access this extended electron donor is unprecedented for carbazole derivatives. This aldehyde part allowed an affordable synthesis of 12 dyes **A–N** differing by the electron acceptors and containing two isolated isomers: **D/I** (see Fig. 1). It must be noticed that this strategy consisting in generating an extended carbazole-based electron

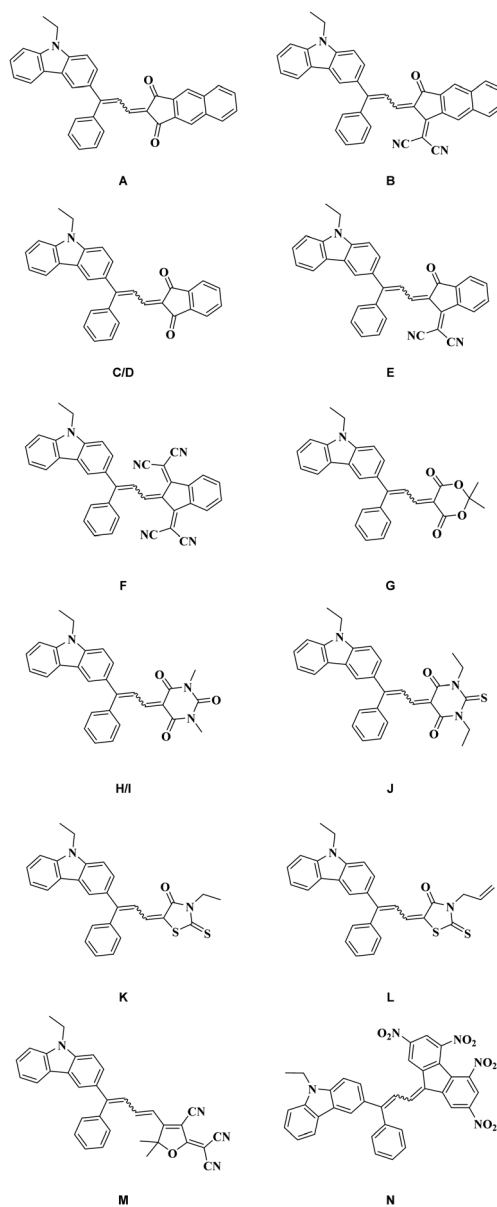


Fig. 1 Chemical structures of dyes **A–N**.

donor by preparing first a carbazole–benzophenone adduct later converted as an acrylaldehyde is unprecedented. To the best of our knowledge, no such 3,3-diphenylacrylaldehyde derivative has been previously reported in the literature. However, the conversion of a carbazole–benzophenone adduct as an alkene has recently been reported in the literature.⁷⁴ The different dyes were seen by a strong absorption extending over the visible range. These dyes were characterized using various techniques including UV-visible absorption and fluorescence spectroscopy and cyclic voltammetry. Theoretical calculations were also carried out to get a deeper insight into the electronic transitions involved in the optical transitions. Finally, solvatochromism of the 12 dyes was examined on several solvent polarities scales and linear correlations could be obtained using the Kamlet–Taft, Reichardt and Catalan empirical models.

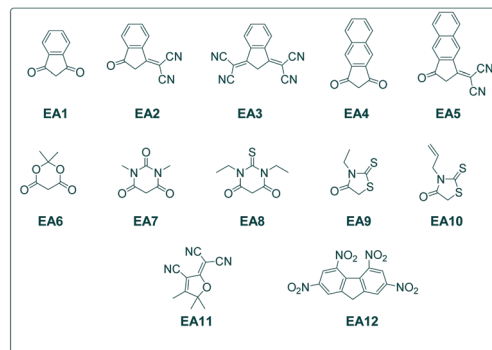


B Results and discussion

B1. Synthesis of the dyes

The extended aldehyde **Cbz-BP** has been synthesized in three steps starting from 9-ethyl-9H-carbazole. By performing a Friedel–Crafts reaction with benzoyl chloride in the presence of aluminum chloride, (9-ethyl-9H-carbazol-3-yl)(phenyl)methanone could be prepared in 84% yield. Addition of methylmagnesium iodide followed by a dehydration reaction furnished 9-ethyl-3-(1-phenylvinyl)-9H-carbazole in almost quantitative yield (see Scheme 2). Formylation of the alkene by a Vilsmeier–Haack reaction gave **Cbz-BP** as a mixture of *s-cis/s-trans* isomers that could not be separated *via* column chromatography. All attempts of iodine-catalyzed thermal *cis/trans* isomerization in toluene did not allow the modification of the ratio between isomers.⁷⁵ If no structural determination was carried out to identify which isomer was the main product, a 2 : 1 ratio could be found on the proton NMR spectrum of **Cbz-BP**. Interestingly, modification of the reaction time during the Vilsmeier–Haack reaction did not modify the ratio between isomers. Consequently, this mixture of isomers was subsequently used for the synthesis of the different dyes. Besides, in order to investigate the potential influence of the isomerization onto the optical properties, one of the two isomers could be obtained in almost pure form but in small quantity by crystallisation at $-20\text{ }^{\circ}\text{C}$ in a mixture of THF/pentane solvents.

Finally, eleven of the twelve dyes of the series were prepared *via* a Knoevenagel reaction carried out under basic conditions. Except for dyes **B** and **E** for which diisopropylamine (DIPA) was used instead of piperidine, all dyes could be obtained with reaction yields ranging from 78 to 92% (see Scheme 3). In the case of dyes **B** and **E**, the choice of DIPA as the base was motivated by recent results reported in the literature mentioning a nucleophilic attack of secondary amines on the cyano groups of **EA5**, inducing a cyclization reaction and producing azafluorenone derivatives.^{76–80} However, piperidine could be used for the synthesis of dye **M**, no nucleophilic attack of amine being reported at present in the literature for **EA11**. Finally, dye **F** was prepared by using a specific procedure. Indeed, due to the

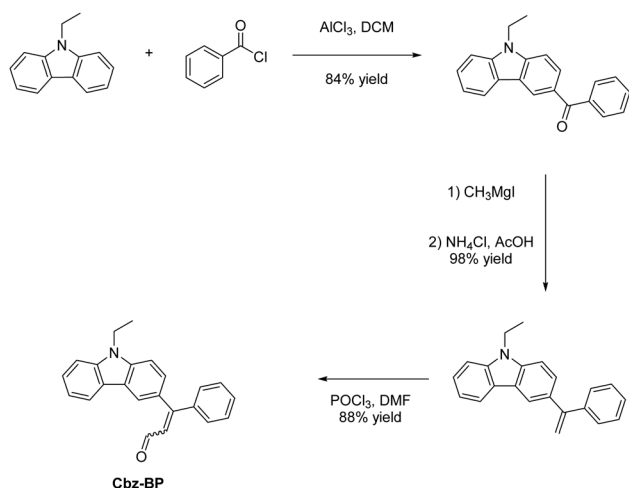


Scheme 3 Synthetic route to dyes A–N.

remarkable stability of the **EA3** anion in basic conditions, no condensation reaction can occur. To circumvent this problem, the condensation of **Cbz-BP** and **EA3** in acetic anhydride furnished dye **F** in 85% yield.

In the case of symmetric electron acceptors (**EA1**, **EA3**, **EA4**, **EA6**, **EA7**, **EA8** and **EA12**), the corresponding dyes (**A**, **C**, **F–J** and **N**) were obtained as a mixture of *s-cis* and *s-trans* isomers. Conversely, a more complex situation was found for dyes prepared with **EA2**, **EA5**, **EA9**, **EA10** and **EA11**, since in the complement of a *s-cis/s-trans* mixture, the different electron acceptors could adopt two different orientations so that a mixture of 4 isomers could be theoretically obtained in these cases (see Fig. 2). Besides, the steric hindrance induced by the use of asymmetric electron acceptors such as **EA2** and **EA5** certainly favour an orientation over the other. It has to be noticed that among the twelve electron acceptors issued in this study, **EA2**,⁸¹ **EA3**,⁸² **EA4**,⁸³ **EA5**,⁷⁶ **EA11**⁸⁴ and **EA12**⁸⁵ had to be prepared.

Interestingly, two isomers (**D** and **I**) could be isolated in pure form. Indeed, due to the marked insolubility of one of the isomers of dyes **I**, these could be easily separated from its mixture *via* precipitation. Conversely, in the case of dye **D** (which is a pure isomer of **C**), this latter was obtained by opposing the isolated isomer of **Cbz-BP** to **EA1** under basic conditions. Finally, due to the exceptional electron-withdrawing ability of **EA12**, a spontaneous deprotonation can occur in highly polar solvents such as DMF. Therefore, **N** could be prepared at room temperature in DMF, without using a base, and obtained as a precipitate upon addition of pentane



Scheme 2 Synthetic route to **Cbz-BP**.



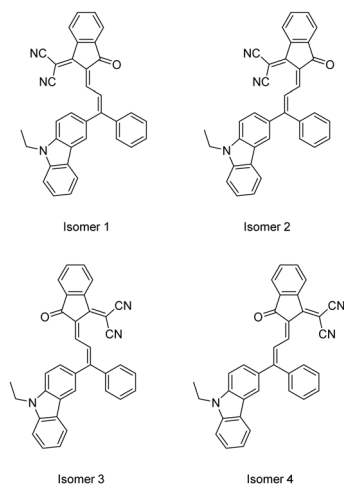


Fig. 2 The four possible isomers when asymmetric electron acceptors are used.

to the reaction media. Dye **N** was isolated in the pure form by filtration, in 87% yield.

B2. Optical properties

All dyes showed good solubility in chloroform so that a comparison of their optical properties could be established. A summary of the optical properties is provided in Table 1 and Fig. 3 and 4. As shown in the Fig. 3, a severe variation of the absorption maxima with the electron-withdrawing ability of dyes **A–N** could be detected. Thus, absorption maxima ranging from 475 nm (**K**) to 587 nm (**N**) and 610 nm (**B**) were found. Position of the ICT bands undergo a redshift following the order of the electron acceptor strength. Notably, comparison of the absorption maxima between **A** and **C** revealed the elongation of the indane-1,3-dione-based acceptor **EA4** to red-shift the absorption of **A** by *ca.* 30 nm compared to that determined for **C** comprising **EA1** as the acceptor. Interestingly, a superimposition of the absorption spectra of **A** and **J** could be

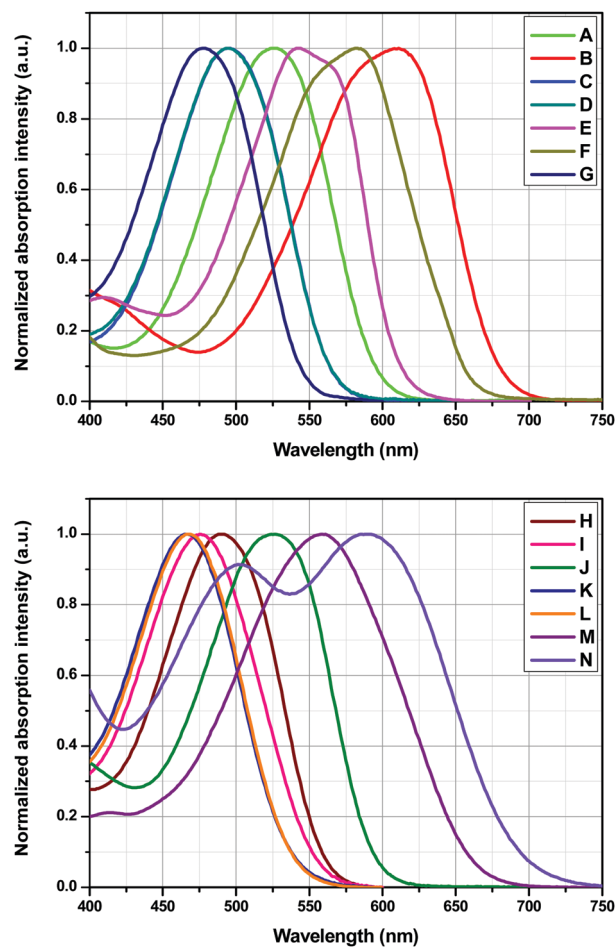


Fig. 3 Normalized UV-visible absorption spectra of dyes **A–N** in chloroform.

clearly evidenced, demonstrating that the electron-withdrawing abilities of 1*H*-cyclopenta[*b*]naphthalene-1,3(2*H*)-dione **EA4** and 1,3-diethyl-2-thioxodihydropyrimidine-4,6(1*H*,5*H*)-dione **EA8** were

Table 1 Optical characteristics of the different compounds in chloroform with the values theoretically determined in CH₂Cl₂. HOMO–LUMO gaps (ΔE) are also shown in this table

Compounds	A	B	C	D	E	F	G
λ_{exp} (nm)	525	610	495	495	575	583	479
λ_{theo} (nm)	495	559	475	—	542	554	470
ϵ_{exp} (M ^{−1} cm ^{−1})	42 600	57 750	36 400	35 900	40 300	37 700	35 400
ϵ_{theo} (M ^{−1} cm ^{−1})	81 300	62 600	66 000	—	63 300	93 000	46 700
ΔE_{exp} (eV)	2.36	2.03	2.50	2.50	2.15	2.12	2.59
ΔE_{theo} (eV)	2.51	2.22	2.61	—	2.29	2.22	2.64

Compounds	H	I	J	K	L	M	N
λ_{exp} (nm)	490	475	526	467	467	558	501, 587
λ_{theo} (nm)	477	—	506	487	487	550	517, 656
ϵ_{exp} (M ^{−1} cm ^{−1})	45 100	43 200	61 600	49 600	47 300	55 100	51 300, 56 000
ϵ_{theo} (M ^{−1} cm ^{−1})	54 400	—	68 600	72 600	72 700	96 200	70 800, 8900
ΔE_{exp} (eV)	2.53	2.61	2.36	2.65	2.65	2.22	2.11
ΔE_{theo} (eV)	2.60	—	2.22	2.55	2.55	2.25	1.89



comparable, because the two dyes possessed the same electron donor.

Among the different electron acceptors, the most interesting one is undoubtedly **EA12**, which enables the design of a dye (**N**) exhibiting two ICT bands located at 510 and 587 nm respectively. For comparison, **EA5** which is the strongest electron acceptor of the series could only furnish a dye (**B**) with an absorption maximum at 610 nm, and therefore only shifted by about 20 nm compared to dye **N**. Based on previous reports devoted for tetranitrofluorene-based dyes, the presence of the two intense ICT bands detected in the UV-visible absorption spectrum of **N** can be assigned to a significant intramolecular charge transfer occurring in the ground state.⁸⁶ In particular, due to the presence of four electron withdrawing nitro groups onto TNF, numerous mesomeric forms can be written, differing by the length of the π -conjugated spacer between the nitro and the amino groups (see Fig. 4).

As anticipated, the highest molar extinction coefficients could be found for the four dyes bearing the strongest electron acceptors (**EA5**, **EA8**, **EA11** and **EA12**), namely dyes **B**, **J**, **M** and **N**. By increasing the electron acceptor strength, the oscillator strength is logically increased, enhancing the molar extinction coefficients (see Fig. S1, ESI†).

A comparison with the simulated UV-visible absorption spectra in dichloromethane revealed the calculations to give absorption maxima to be in perfect agreement with the experimental one (see Fig. S1 and S2, ESI†). Finally, considering that 2 isomers (**D** and **I**) could be separated from their respective mixtures (**C** and **H**), a comparison between the absorption spectra of the isolated isomer and the mixture could be established (see Fig. 5). Interestingly, if the absorption spectrum of **C** perfectly superimposed that of **D**, a completely different behavior was found for **H** and **I**, the ICT band of **I** being blue-shifted by 14 nm (476 nm vs. 490 nm for **H**). Considering that the isomer **I** is less conjugated than its analogue, on the basis of the UV-visible absorption spectra, it can be confidently proposed that **I** is the *s-cis* isomer of **H**.

Additionally, in this configuration, the *s-cis* configuration imposes an internal torsion inside the molecules due to the steric hindrance generated by the donor and the acceptor so that a reduction of the conjugation between the two partners is

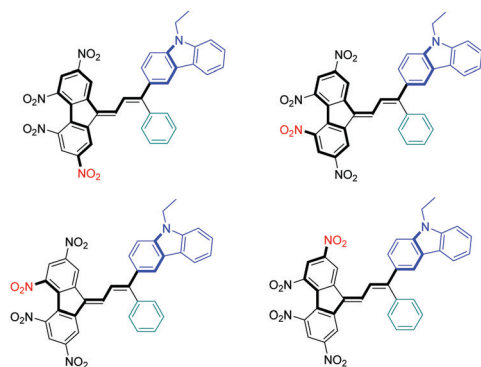


Fig. 4 Different mesomeric forms supporting the presence of two ICT bands for the TNF-based dye **N**.

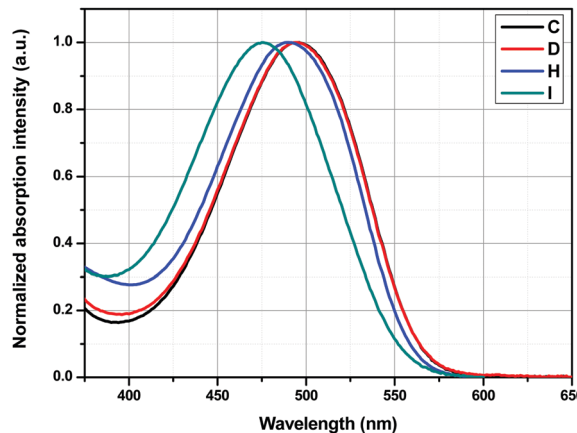


Fig. 5 Comparison of the UV-visible absorption spectra of **C/D** and **H/I** in chloroform.

logically observed. Considering that severe variations of the absorption maxima were found by varying the electron-withdrawing groups, the experimental behavior observed for dyes **A–N** was rationalized by theoretical calculations.

B3. Theoretical calculations

The energy levels of the highest occupied molecular orbital (HOMO) and the lowest unoccupied molecular orbital (LUMO) were determined by theoretical calculations and a representative set of the electronic distribution is provided in Fig. 6. As evidenced in Fig. 6, a HOMO energy level specifically localized on the carbazole group and a LUMO energy level comprising both the acceptor and the lateral phenyl ring could be found for all dyes. While examining the energy levels of both the HOMOs and LUMOs of all dyes, almost no variation of the HOMO level was found, consistent with a localization of the HOMO onto the carbazole moiety (see Table 2). Conversely, major differences could be found for the positions of the LUMO levels, resulting from their localization onto the acceptor moieties. As a result of this, the variation of the HOMO–LUMO gap only originates from the variation of the positions of the LUMO levels. More precisely, by improving the electron-accepting ability, the LUMO level is stabilized, decreasing the HOMO–LUMO gap.

B4. Solvatochromism

Push–pull dyes are compounds that are highly sensitive to their environment (polarity, and polarizability of the solvents) and position of the ICT bands can drastically vary with the solvent effects.^{87,88} Generally, for push–pull dyes, both HOMO and LUMO energy levels are precisely localized onto two different parts of the molecule. In the present case, a positive solvatochromism is typically observed, evidencing an important charge redistribution upon photoexcitation and a dipole moment larger in the excited state than in the ground state. This behavior is typically found for push–pull dyes exhibiting a relatively small dipole moment in the ground state, which is achievable by separating the donor from the acceptor by an elongated spacer.^{89–93} To investigate the solvatochromic



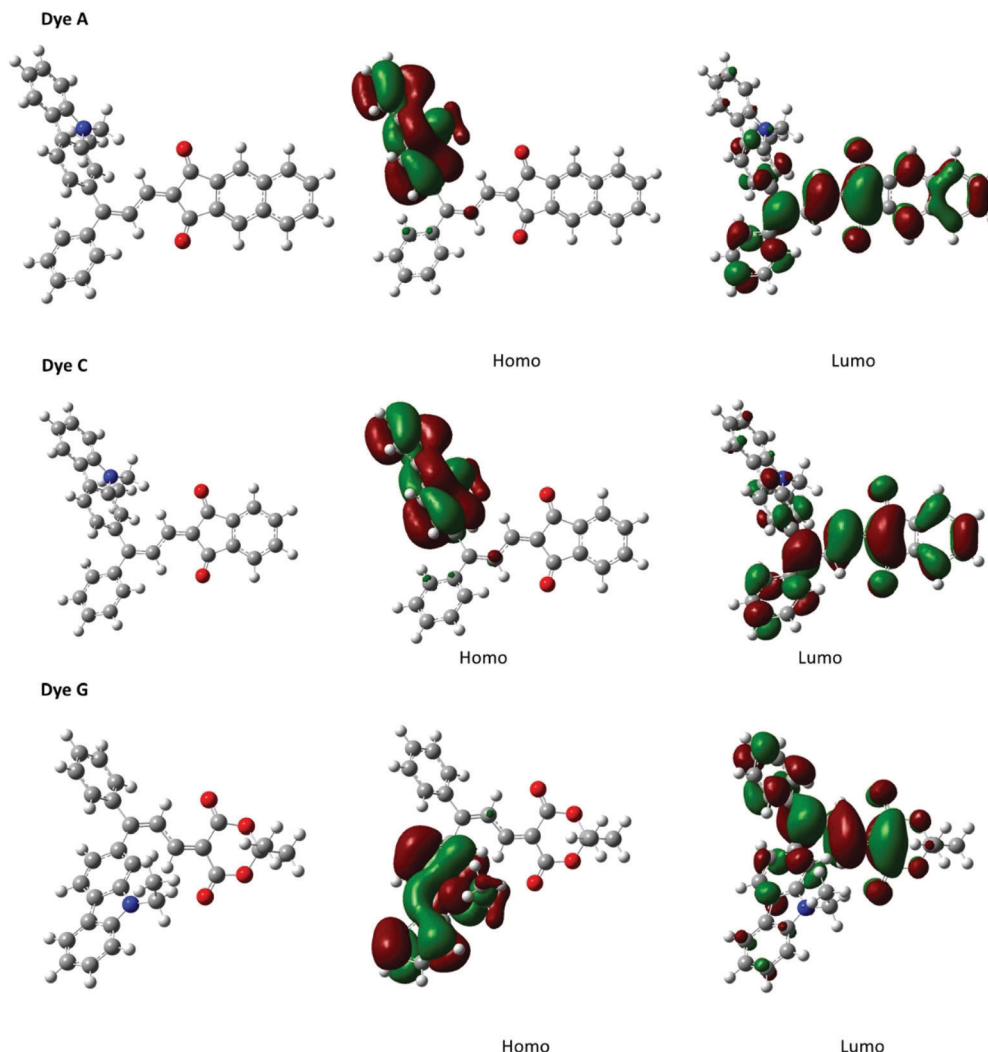


Fig. 6 Optimized geometries and HOMO/LUMO electronic distributions of dyes **A**, **C** and **G**.

Table 2 Theoretical positions of the HOMO and LUMO energy levels

Compounds	A	B	C/D	E	F	G
HOMO (eV)	−5.55	−5.48	−5.56	−5.71	−5.83	−5.69
LUMO (eV)	−2.72	−3.17	−2.64	−3.06	−3.27	−2.78

Compounds	H/I	J	K	L	M	N
HOMO (eV)	−5.64	−5.71	−5.66	−5.67	−5.95	−5.90
LUMO (eV)	−2.79	−2.99	−2.72	−2.72	−3.33	−3.91

properties of dyes, numerous solvent polarity scales have been developed over the years and the Kamlet–Taft's,⁹⁴ Dimroth–Reichardt's,⁹⁵ Lippert–Mataga's,⁹⁶ Catalan's,^{97,98} Kawski–Chamma–Viallet's,⁹⁹ McRae's,¹⁰⁰ Suppan's,¹⁰¹ and Bakhshiev's¹⁰² scales can be cited as the most popular ones. Investigations of the solvatochromic properties of dyes **A–N** were carried out in 23 solvents of different polarities and changes in the longest-wavelength absorption maxima λ_{max} for the 14 dyes are summarized in Table 3. First, the intramolecular nature of

the charge transfer detected for all dyes was verified by realizing successive dilutions and linear plots could be obtained by measuring the absorbance *vs.* the dye concentration. This point being verified, the resulting data were evaluated by means of the different abovementioned empirical models. Interestingly, remarkable correlations were obtained on the Kamlet–Taft and the Catalan scales as well as, to a lesser extent, on the Reichardt scale. In particular, for the Catalan, several parameters have been developed such as the solvent polarizability (SP), the solvent dipolarity (SdP) and the solvent polarity/polarizability (SPP) parameters.^{103–105} In this work, the best correlations were obtained with the Catalan SPP parameters taking into account the polarity/polarizability of the solvents. This is also this interaction, which is considered by the two other solvatochromic scales, namely the Kamlet–Taft and the Reichardt models. Therefore, the solvent–solute interactions are mainly governed by the polarity/polarizability of the solvents, which was confirmed by three polarity scales based on these interactions. Among all dyes, the best correlations were obtained for dyes **B** or **K**, as shown in the Fig. 7.



Table 3 Summary of the optical properties of compounds **A–H** in solvents of various natures

Compounds	A^a	B^a	C^a	D^a	E^a	F^a	G^a	H^a	I^a	J^a	K^a	L^a	M^a	N^a
Acetone	510	585	456	456	553	546	456	466	462	460	456	457	534	562
Acetonitrile	508	579	489	476	550	545	461	466	463	504	455	456	526	561
AcOEt	502	581	479	477	552	551	457	465	458	499	455	455	525	561
Anisole	516	591	491	489	562	573	471	474	466	508	464	464	549	582
Butanol	525	604	498	497	569	nd ^b	475	486	474	521	461	462	550	568
Chloroform	526	610	495	495	542	583	478	489	475	524	466	468	559	588
Cyclohexane	488	563	466	466	564	543	446	459	450	488	445	445	520	nd ^b
1,2-Dichloroethane	515	592	488	488	564	576	468	479	470	516	462	465	551	586
Dichloromethane	517	597	489	489	548	577	471	480	469	514	463	464	553	586
Diethyl carbonate	499	575	474	475	547	546	456	462	459	496	452	452	521	553
Diethyl ether	496	572	472	469	548	547	451	462	455	497	453	451	528	nd ^b
Diglyme	508	588	483	483	559	574	463	471	466	511	461	461	545	569
1,4-Dioxane	503	573	477	477	548	546	458	467	459	501	454	456	525	559
Dimethylacetamide	516	595	488	491	564	577	470	472	471	518	466	466	542	568
Dimethylformamide	517	596	489	488	560	576	469	472	468	512	463	465	539	577
DMSO	524	598	493	495	565	580	474	476	471	516	468	468	541	581
Ethanol	522	596	492	491	563	nd ^b	475	480	466	517	458	459	548	nd ^b
Heptane	485	560	464	464	539	539	443	457	446	485	442	442	514	nd ^b
Nitrobenzene	528	617	500	502	579	587	479	488	481	528	473	473	568	562
THF	507	583	482	478	556	563	460	467	460	503	458	458	533	567
Toluene	509	581	483	483	551	554	464	471	465	506	459	460	527	567
Triethylamine	—	nd ^b	470	470	nd ^b	nd ^b	455	467	455	nd ^b	450	450	522	nd ^b
<i>p</i> -Xylene	508	572	479	479	553	550	462	468	464	504	458	459	528	569

^a Position of the ICT bands are given in nm. ^b nd: not determined.

For all dyes, negative slope lines were found for the linear regressions, irrespective of the polarity scales, indicative of positive solvatochromism. Considering that the absolute value of the slopes can provide information of the sensitivity of the dyes to the solvent polarity and thus of the charge redistribution upon excitation, a summary of the different negative slope lines is provided in Fig. 8. Interestingly, in the series of 12 dyes, three different behaviors could be detected. Thus, only dyes **E**, **F**, **H**

and **N** differ from the others for which similar slopes were found for both the Kamlet–Taft or the Catalan plots. Comparison between **H** and **I** is particularly interesting since **I** is a pure isomer isolated from **H**. Besides, a completely different behavior with regard to the solvent polarity was found, **H** exhibiting a lower sensitivity than **I**. These results confirm that **I** is certainly the *s-cis* isomer, and that the internal torsion induced by the steric hindrance between the donor and the acceptor are

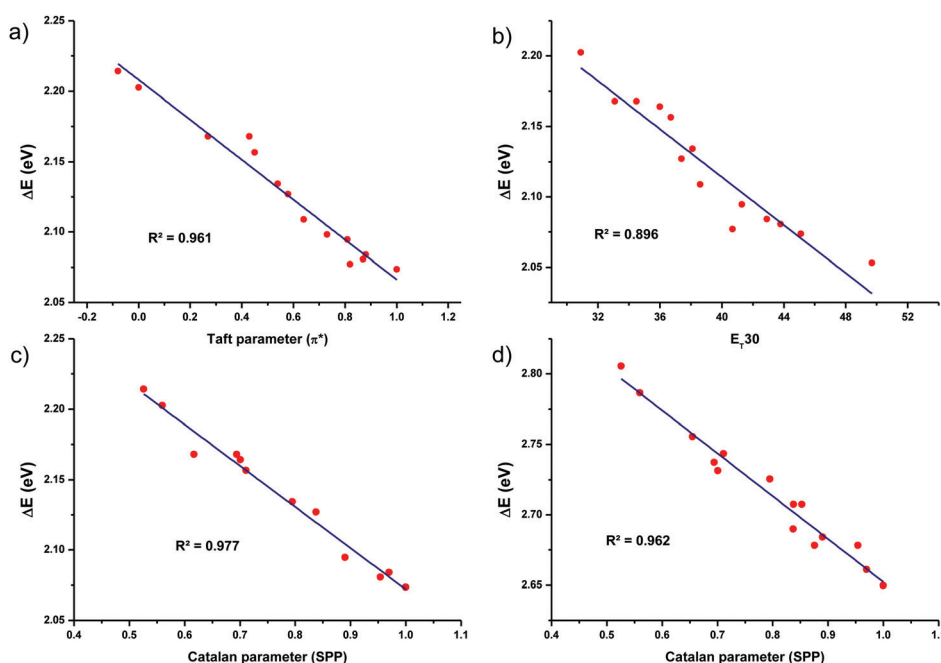


Fig. 7 Scatterplots with overlaid linear regression plots for dye **B** with (a) the Kamlet–Taft graph (b) the Reichardt graph (c) the Catalan graph and for dye **K** (d) the Catalan graph.



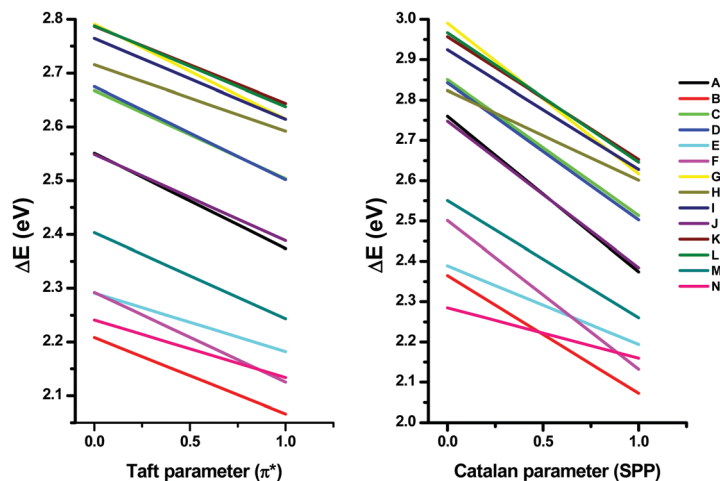


Fig. 8 Variation of the positions of the charge transfer band with the Kamlet–Taft and Catalan empirical parameters for dyes A–N.

drastically modified by the polarity of the solvents. Conversely, C and its pure isomer D show similar slopes in both plots. Based on the NMR experiments, similarity of behaviors between C and D arises from the fact that D is mostly composed of C, resulting in a minor contribution of the second isomer on the optical properties. A lower sensitivity to the solvent polarity is also found for dyes E and N what is counter-intuitive considering that these two dyes possess strong electron acceptors. Finally, the most negative slopes were determined for dye F bearing EA3 as the acceptor. This pronounced sensitivity to the solvent polarity can be assigned to the presence of the numerous cyano groups on EA3, favoring a major redistribution of the electronic density upon excitation when highly polar solvents are used.

B5. Photoluminescence spectroscopy

The emission properties of all dyes were examined in chloroform as the solvent. Since almost all compounds were fluorescent, and were therefore not sensitive to oxygen, all experiments were carried out in air. Most of the dyes were photoluminescent so that a Stokes shift could be determined for most of the dyes. In addition, as shown in Fig. 9, by determining the crossing point between the UV-visible and the fluorescence spectra of

each dye, energy of the first singlet excited state (E_{S1}) could be determined. All data are summarized in Table 4.

Carbazoles are known to be fluorescent compounds which explains why the different dyes were emissive.^{106–108} Only dye N comprising a TNF moiety as the acceptor was not emissive. The lack of photoluminescence of compound N can be attributed to the presence of the nitro groups, well-known to efficiently quench the photoluminescence of luminescent compounds.^{109,110} For all dyes, Stokes shift ranging from 50 nm for compound B to 97 nm for compound M could be determined. The largest Stokes shift was obtained for dye M comprising the 2-dicyanomethylidene-3-cyano-4,5,5-trimethyl-2,5-dihydrofuran (TCF) group as the acceptor.^{111,112} These large Stokes shifts are indicative of a significant electronic redistribution between the ground state and the excited state.¹¹³ Additionally, the fluorescence of the indane compounds (comprising EA1–EA5 as the electron acceptors) seems to be highly dependent of the number of the cyano functions. Indeed, the Stokes shift of F was only 61 nm while a Stokes shift of 68 nm and 78/77 nm could be respectively determined for compounds E and C/D. In fact, a reduction of the Stokes shift with a redshift of the absorption maxima was observed, consistent with a decrease of the gap between the occupied and unoccupied orbitals. The same trend was observed while comparing the Stokes shifts between B and E. Indeed, due to the improved electron-withdrawing ability of EA2 compared to EA5, a reduction of the Stokes shift was detected, going from E (68 nm) to B (50 nm).

B6. Electrochemical properties

All push–pull dyes were investigated by cyclic voltammetry (CV) in deaerated acetonitrile solutions. The selected set of voltammograms is shown in Fig. 9 and all CV curves are given in the ESI.† The redox potentials of all compounds are summarized in the Table 5 in which the redox potentials are given against the half wave oxidation potential of the ferrocene/ferrocenium cation couple.

All the synthesized compounds exhibited irreversible single-electron oxidation and single-electron reduction processes. Concerning the oxidation, all the push–pull molecules were

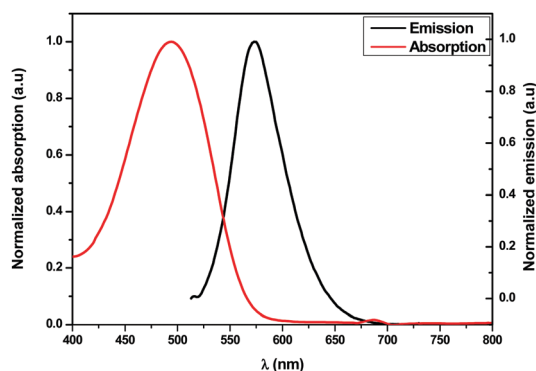


Fig. 9 Normalized absorption and emission intercept curves of dye D in chloroform.



Table 4 Fluorescence properties of the different compounds recorded in chloroform

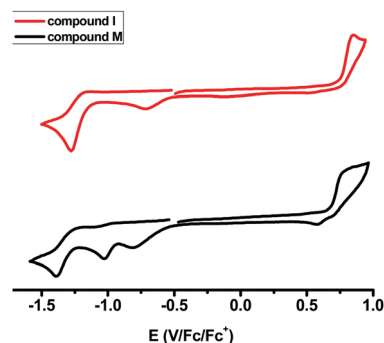
	A	B	C	D	E	F	G
λ_{ex} (nm)	526	610	495	495	575	583	479
λ_{em} (nm)	612	660	573	572	643	644	562
Stokes shift (cm^{-1})	116 279	200 000	128 205	129 870	147 059	163 934	120 482
E_{S1} (eV)	2.13	1.92	2.28	2.28	2.01	2.01	2.34

	H	I	J	K	L	M	N
λ_{ex} (nm)	490	475	526	467	467	558	587
λ_{em} (nm)	570	570	600	555	550	655	—
Stokes shift (nm)	125 000	105 263	135 135	113 636	120 482	103 093	—
E_{S1} (eV)	2.30	2.31	2.18	2.41	2.43	1.99	—

designed from the same donor moiety, namely the **Cbz-BP**, which explains that comparable oxidation potential are detected for all dyes, the oxidation of the carbazole group being detected around 750 mV per Fc per Fc^+ . An oxidation process lowered by *ca.* 100 mV was also observed for the compounds **L** and **M** and reduction of the oxidation potential can be explained by an oxidation process occurring on the amino group of the rhodanine acceptors.

Regarding the reduction process, most of the molecules are affected by a monoelectronic reduction located on the central double bond, slightly impacted by the acceptor moiety. Indeed, the stronger the electron withdrawing group, the stronger the cathodic shift. A relevant example of this can be obtained by comparing the reduction potentials of **M** and **I** (See Fig. 10 and Table 5). In this case, **EA11** is more accepting than **EA7**, explaining the 200 mV cathodic shift occurring during the reduction of the double bond.

Nonetheless, a reduction at a smaller cathodic potential was observed for **N**, presumably resulting from the reduction of the

**Fig. 10** Cyclic voltammograms of **I** (up) and **M** (down) in acetonitrile solutions (10^{-3} M) using tetrabutylammonium perchlorate (0.1 M) as the supporting electrolyte. Scan rate: 100 mV s^{-1} .

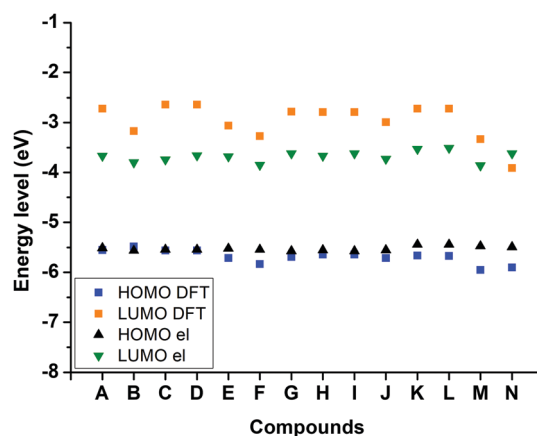
nitro groups prior to the central double bond. Reduction of the nitro groups is frequently observed in the case of poly(nitrofluorene) structures, explaining the difference of 200–500 mV observed between the reduction peak of this compound and that of the other dyes.^{114,115}

Finally, by using the ferrocene (Fc) ionization potential value as a standard and the equation established by Pommerehne *et al.*, energy levels of the highest occupied molecular orbital (HOMO) and the lowest unoccupied molecular orbital (LUMO) could be estimated from the redox potentials (see Table 5).¹¹⁶

Table 5 Outline of the electrochemical properties of dyes **A–N** compared with DFT calculations. E_{ox} correspond to all the oxidation potentials and E_{red} the reduction potentials. ΔE_{el} corresponds to the difference between the HOMO and the LUMO levels, electrochemically obtained. Finally, ΔE_{th} is the difference between the HOMO and LUMO levels, theoretically obtained. Fc/Fc^+ was used as internal reference standard

Compounds	$E_{\text{ox onset}}$ (V per Fc)	$E_{\text{red onset}}$ (V per Fc)	E_{HOMO} (eV)	E_{LUMO} (eV)	ΔE_{el}^a (eV)	ΔE_{th}^b (eV)
A	0.71	−1.13	−5.51	−3.67	1.84	2.83
B	0.76	−1.00	−5.56	−3.80	1.76	2.31
C	0.74	−1.06	−5.54	−3.74	1.80	2.92
D	0.74	−1.14	−5.54	−3.66	1.88	2.92
E	0.72	−1.12	−5.52	−3.68	1.84	2.65
F	0.74	−0.95	−5.54	−3.85	1.69	2.56
G	0.77	−1.18	−5.57	−3.62	1.95	2.91
H	0.75	−1.14	−5.55	−3.67	1.93	2.85
I	0.77	−1.18	−5.57	−3.62	1.95	2.85
J	0.75	−1.07	−5.55	−3.73	1.82	2.72
K	0.63	−1.27	−5.44	−3.53	1.91	2.94
L	0.63	−1.29	−5.44	−3.51	1.93	2.95
M	0.67	−0.94	−5.47	−3.86	1.61	2.62
N	0.69	−1.18	−5.49	−3.62	1.87	1.99

^a Onset of the oxidation and reduction potentials *versus* ferrocene (E_{onset} vs. Fc). E_{HOMO} and E_{LUMO} were determined from the formulae: $E_{\text{HOMO}} = -4.8 - E_{\text{ox}}$ vs. Fc and $E_{\text{LUMO}} = -4.8 - E_{\text{red}}$ vs. Fc.¹¹³ ^b E_{HOMO} and E_{LUMO} are given in eV.

**Fig. 11** Comparison between frontier orbital energy levels obtained from cyclic voltammetry and DFT calculations.

Thanks to the determination of the HOMO and LUMO energy levels by electrochemistry, a comparison could be made between the experimental and the theoretical values of the two energy levels (see Fig. 11). If a clear difference can be noticed between the LUMO_{DFT} and the LUMO_{el}, conversely, a good adequation between the theoretical and the experimental energy level of the HOMO orbital was found, as shown in the Fig. 11. Even if the position of the LUMO energy levels was overestimated by theoretical calculations, similar variations were found between the LUMO_{DFT} and the LUMO_{el}. This difference can be attributed to the fact that influence of the solvent on the LUMO energy level is not fully considered in the calculations.

C Conclusions

To conclude, a series of 12 push-pull dyes differing by the electron-acceptors have been designed and synthesized. Interestingly, the carbazole-based electron donor reported in this work has been synthesized for the first time according to a multistep synthesis. All dyes prepared using this electron donor proved to be soluble in most of the common organic solvents, even if electron acceptors such as (thio)barbituric acids and tetranitrofluorene derivatives well known to produce low soluble dyes have been used. By modifying the electron acceptors, a set of dyes absorbing over the whole visible range have been obtained. For all dyes, a positive solvatochromism could be determined, evidencing that the excited state of the dye is more polar than the ground state. If promising results have been obtained with this electron donor, the asymmetry of this elongated aldehyde is at the origin of the presence of numerous isomers. As a result of this, complicated NMR results were obtained. Future work will consist in developing a more symmetric structure bearing two carbazole moieties, to first simplify the NMR assignment, but also to improve the electron-donating ability of the donor.

Conflicts of interest

There are no conflicts to declare.

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