

This is a repository copy of *Liquid-crystalline dual fluorescent dyes for white light emission*.

White Rose Research Online URL for this paper:

<https://eprints.whiterose.ac.uk/id/eprint/227255/>

Version: Published Version

Article:

Suleymanova, Alfiya F., Fortwengler, James R., Chen, Xinrui et al. (4 more authors) (2025)
Liquid-crystalline dual fluorescent dyes for white light emission. *Journal of Materials Chemistry C*. ISSN 2050-7534

<https://doi.org/10.1039/d5tc01393c>

Reuse

This article is distributed under the terms of the Creative Commons Attribution (CC BY) licence. This licence allows you to distribute, remix, tweak, and build upon the work, even commercially, as long as you credit the authors for the original work. More information and the full terms of the licence here:

<https://creativecommons.org/licenses/>

Takedown

If you consider content in White Rose Research Online to be in breach of UK law, please notify us by emailing eprints@whiterose.ac.uk including the URL of the record and the reason for the withdrawal request.



Cite this: DOI: 10.1039/d5tc01393c

Liquid-crystalline dual fluorescent dyes for white light emission†

Alfiya F. Suleymanova,^a James R. Fortwengler,^b Xinrui Chen,^d Rafal Czerwieniec,^c Yafei Wang,^{b,*d} Marsel Z. Shafikov^{b,*c} and Duncan W. Bruce^{b,*a}

Three isomeric phenanthrolines- **2,9-CzPhen**, **3,8-CzPhen** and **4,7-CzPhen** - are formed when the heterocyclic core is functionalised 2,9-, 3,8- or 4,7- with 3,6-bis-(3,4-dioctyloxybenzene)-9H-carbazole. All are liquid crystalline and show a single columnar hexagonal phase, although that of **2,9-CzPhen** has a very much higher clearing point (308 °C) compared to **3,8-CzPhen** and **4,7-CzPhen** (175 °C and 158.5 °C). This is rationalised through the sterically driven disposition of the carbazole groups on the phenanthroline ring alongside likely columnar packing motifs. Photophysical properties were investigated in toluene solution and in a polystyrene film. All compounds are luminescent (λ_{max} 440–455 nm) with decent (40%, **3,8-CzPhen**) to modest (**4,7-CzPhen**, 13% and **2,9-CzPhen**, 18%) solution fluorescence quantum yields. However, in a polystyrene film, **3,8-CzPhen** and **4,7-CzPhen** also emit at 520 nm and 570 nm, representing a still relatively rare example of anti-Kasha emission. The 520 nm emission of **4,7-CzPhen** shows a low-intensity decay component that involves a triplet state and is assigned to a thermally activated delayed fluorescence (TADF). At high emitter loadings in an OLED device, two emission frequencies result in the appearance of white light. Dual emission is understood through the relevant theory and considers rates of internal conversion following photoexcitation that populates the two emitting states – a carbazole-localised, singlet $^1[\pi_{\text{Cz}}\pi^*_{\text{Cz}}]$ state and a charge-transfer $^1[\pi_{\text{Cz}}\pi^*_{\text{phen}}]$ state. The rate and efficiency of internal conversion (IC) from the Cz-localised excited state to the charge-transfer state, involving electron transfer from the substituent(s) to 1,10-phenanthroline, appear relatively slow so that fluorescence from the higher-lying state competes. This is consistent with anti-Kasha behaviour as a result of a slow internal conversion between electronically decoupled states.

Received 2nd April 2025,
Accepted 6th May 2025

DOI: 10.1039/d5tc01393c

rsc.li/materials-c

Introduction

Tailoring new molecules to combine multiple physical properties can afford materials that allow new applications or improve those already in existence. For instance, designed to feature liquid-crystallinity, photoemissive materials based on multi-resonant TADF can be deposited with preferential horizontal orientation to have a bulk aligned transition dipole moment which, applied in organic light emitting diodes (OLEDs), has the potential to afford improved light outcoupling.^{1,2} Materials

combining white luminescence with mesogenic properties could therefore be beneficial for lighting application of OLEDs, for example as polarised white backlights for liquid crystal displays. Designing materials to show white emission, however, is not a trivial task. Covering the blue and red ranges of the spectrum typically requires dual emission, the occurrence of which in molecules is limited by Kasha's rule,³ which requires emission only from the lowest-energy excited state of a given multiplicity and relies on fast, efficient, non-radiative relaxation processes between the higher excited states. A rare exception is azulene^{4–8} where the two lowest excited singlet states are vibrationally decoupled due to the large energy gap. This reduces the efficiency of internal conversion so that dual fluorescence is observed. Known strategies for obtaining dual emissive materials are: (i) by designing molecules with two non-conjugated chromophores;⁹ (ii) combining the aggregate (excimer) emission with monomolecular emission^{10,11} and (iii) designing molecules with relatively slow intersystem crossing (ISC) so that both fluorescence and phosphorescence are seen.^{12–17} Recently, however, Li *et al.* showed¹⁸ that dual emissive behaviour is achievable in molecules with a

^a Department of Chemistry, University of York, Heslington, York YO10 5DD, UK.
E-mail: duncan.bruce@york.ac.uk; Tel: +44 1904 324085

^b Department of Chemistry, University of Southern California, Los Angeles, California, 90089, USA

^c Institut für Physikalische und Theoretische Chemie, Universität Regensburg, Universitätsstrasse 31, Regensburg, D-93053, Germany.
E-mail: shafikovf@gmail.com

^d School of Materials Science & Engineering, Changzhou University, Changzhou 213164, P. R. China. E-mail: qiji1830404@hotmail.com

† Electronic supplementary information (ESI) available. See DOI: <https://doi.org/10.1039/d5tc01393c>

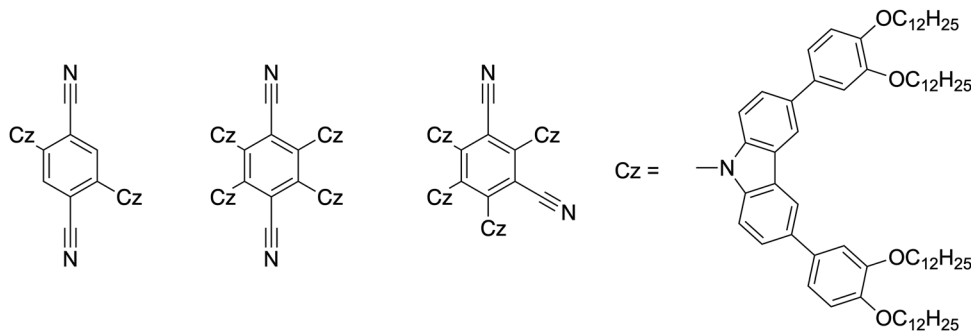


Fig. 1 Examples of LC TADF materials prepared with phthalonitriles as the acceptor and a functionalised carbazole as the donor.

donor-acceptor-donor (D-A-D) unit, where the excited states local to the donor (LE – local excitation) and the donor-acceptor charge-transfer (CT) excited states are decoupled electronically. This reduces the relaxation efficiency between the different states, affording dual emission that combines fluorescence from the donor-localised ^1LE state and thermally activated delayed fluorescence (TADF) from thermal equilibration of the singlet and triplet CT states. This design of dual emissive materials holds much promise for the development of white-emitting OLED materials, especially as TADF can utilise both singlet and triplet excitons formed in the emissive layer of the OLED.

In this regard, we have recently shown a working molecular design approach affording D-A-D type TADF molecules, which comprise terephthalonitrile or isophthalonitrile as the acceptor unit and alkoxyaryl-substituted carbazoles as the donor units and which are also liquid crystalline (Fig. 1).^{19,20} To develop this approach towards the design of dual emissive liquid-crystalline materials, 1,10-phenanthroline seemed an interesting and ubiquitous acceptor unit that offers the potential for functionalisation at different combinations of positions with donor units that may allow modulation of the internal conversion efficiency from the LE states of the donor units to the states of CT character. Due to its efficient electron-transporting properties,²¹ 1,10-phenanthroline and its derivatives are also well known in the field of TADF materials, although in most cases they play a role of ligands in Cu(i) and Ag(i) complexes.^{22–26} While there are several examples of pyrazino-phenanthrolines showing TADF behaviour, there is, however, only one such report of simply functionalised 1,10-phenanthrolines substituted in the 2,9-, 3,8- or 4,7 positions with an *N*-phenoxazine (Fig. 2).²⁷ In toluene solution, all three compounds exhibited emission *via* a TADF mechanism with efficiencies ranging from 45% (2,9-isomer) to 82% (3,8-isomer) and emission wavelengths in the range 535 to 554 nm. An OLED device was constructed based on the 3,8-isomer with an external quantum efficiency of $\approx 19\%$.

1,10-Phenanthroline may also be derivatised to generate liquid-crystalline materials and examples exist with 4,7-,²² 2,9-^{23,28} and 3,8-disubstitution.^{29,30} Given the potentially attractive device properties of 1,10-phenanthrolines and the possibility to generate emissive LC derivatives by using the derivatised

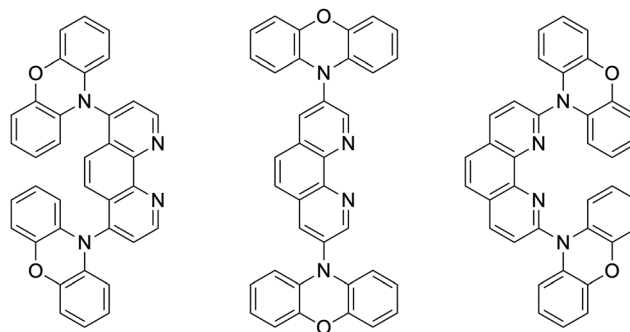


Fig. 2 Purely organic phenanthroline-based TADF compounds.

carbazoles (Scheme 1), we undertook the preparation and characterisation of three isomeric derivatives. The results of this study are now described.

Results and discussion

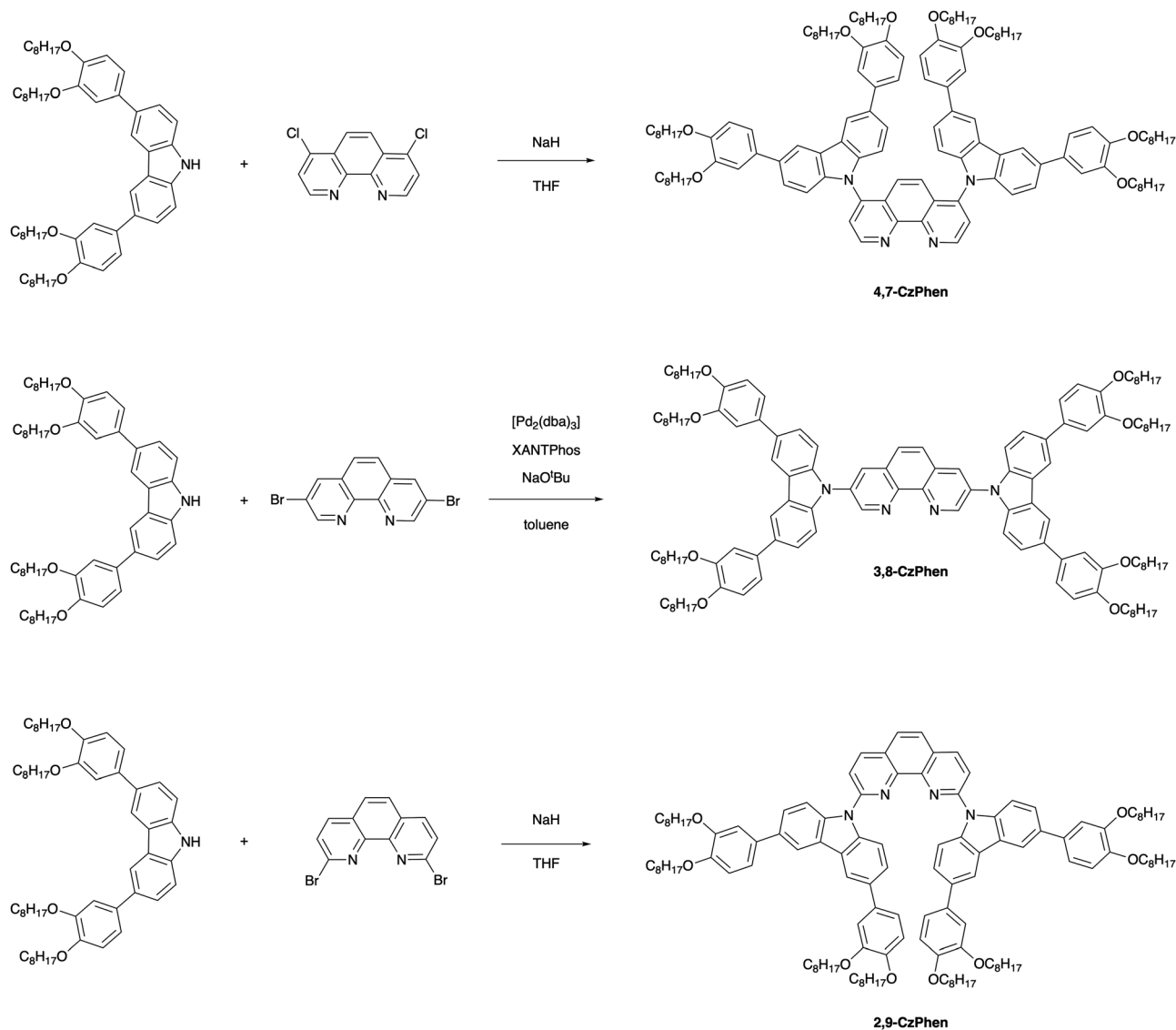
Synthesis

As shown in Scheme 1, the 4,7- and 2,9-disubstituted 1,10-phenanthroline derivatives (**4,7-CzPhen** and **2,9-CzPhen**) were obtained *via* nucleophilic substitution by reaction of dichloro- and dibromo-1,10-phenanthrolines with 3,6-bis-(3,4-dioctyloxybenzene)-9*H*-carbazole. However, due to the lower electrophilicity of 3- and 8-positions, the same protocol did not work for preparation of **3,8-CzPhen** which therefore was obtained using a Pd-catalysed Buchwald–Hartwig amination reaction of 3,8-dibromo-1,10-phenanthroline with the same carbazole.³¹ Unoptimised yields for the coupling reactions were between *ca.* 30% to 45%. The carbazole was prepared as described previously with details provided in the ESI.†²⁰ The structure and purity of the compounds were verified *via* ^1H and $^{13}\text{C}\{^1\text{H}\}$ NMR spectroscopy, elemental analysis and mass spectrometry (ESI†).

Liquid-crystalline properties

The thermal properties of the new compounds (Fig. 3) were studied by polarised optical microscopy, differential scanning calorimetry (DSC) and small-angle X-ray scattering (SAXS). Transition temperatures were first obtained by optical



Scheme 1 Synthesis of **4,7-CzPhen**, **3,8-CzPhen** and **2,9-CzPhen**.

microscopy before being confirmed using DSC, which then informed the temperature range over which X-ray were collected. None of the complexes was obtained as a crystalline solid on isolation from solution nor from cooling the sample on the microscope and so a supercooled mesophase was present at ambient temperature, even if visually the samples appeared solid-like. Initial identification of the phases by optical microscopy showed in each case textures characteristic of a columnar phase on cooling of the sample from the isotropic fluid (Fig. 3, top row) and, in the case of **2,9-CzPhen**, the texture alone allowed identification of the phase as Col_h. In fact, SAXS (Fig. 3, bottom row) showed that indeed all of the phases had the same symmetry, confirmed in each case by the observation of higher-order (11) and (20) reflections as well as (21) for both **4,7-CzPhen** and **2,9-CzPhen**. The *d*-spacings and hexagonal lattice parameters are found in Table 1.

Aspects of the thermal behaviour require further comment. Thus, while both **2,9-CzPhen** and **3,8-CzPhen** appear solid at

room temperature, under the polarising microscope it is evident that they are soft and deformable, consistent with the absence of a melting transition in the DSC trace (Fig. 3, middle row). **3,8-CzPhen** melts at *ca.* 175 °C and then the mesophase re-forms around 168 °C, with the enthalpies measured on heating and cooling being effectively the same. This general behaviour is mirrored in **2,9-CzPhen** with clearing around 308 °C with the mesophase re-forming around 300 °C, again with matching enthalpy change (the much higher temperatures found for **2,9-CzPhen** are discussed below). Both compounds show a hysteresis in the clearing temperature of 7–8 °C and, while this is much larger than would be expected in simple calamitic systems, it is not unduly large for a columnar system.

However, things are a little different with **4,7-CzPhen** and contemplation of the DSC trace shows two apparently anomalous yet related features. Thus, on heating there is an endothermic event with an onset temperature of 158.5 °C and then on cooling there is a much weaker exothermic event with



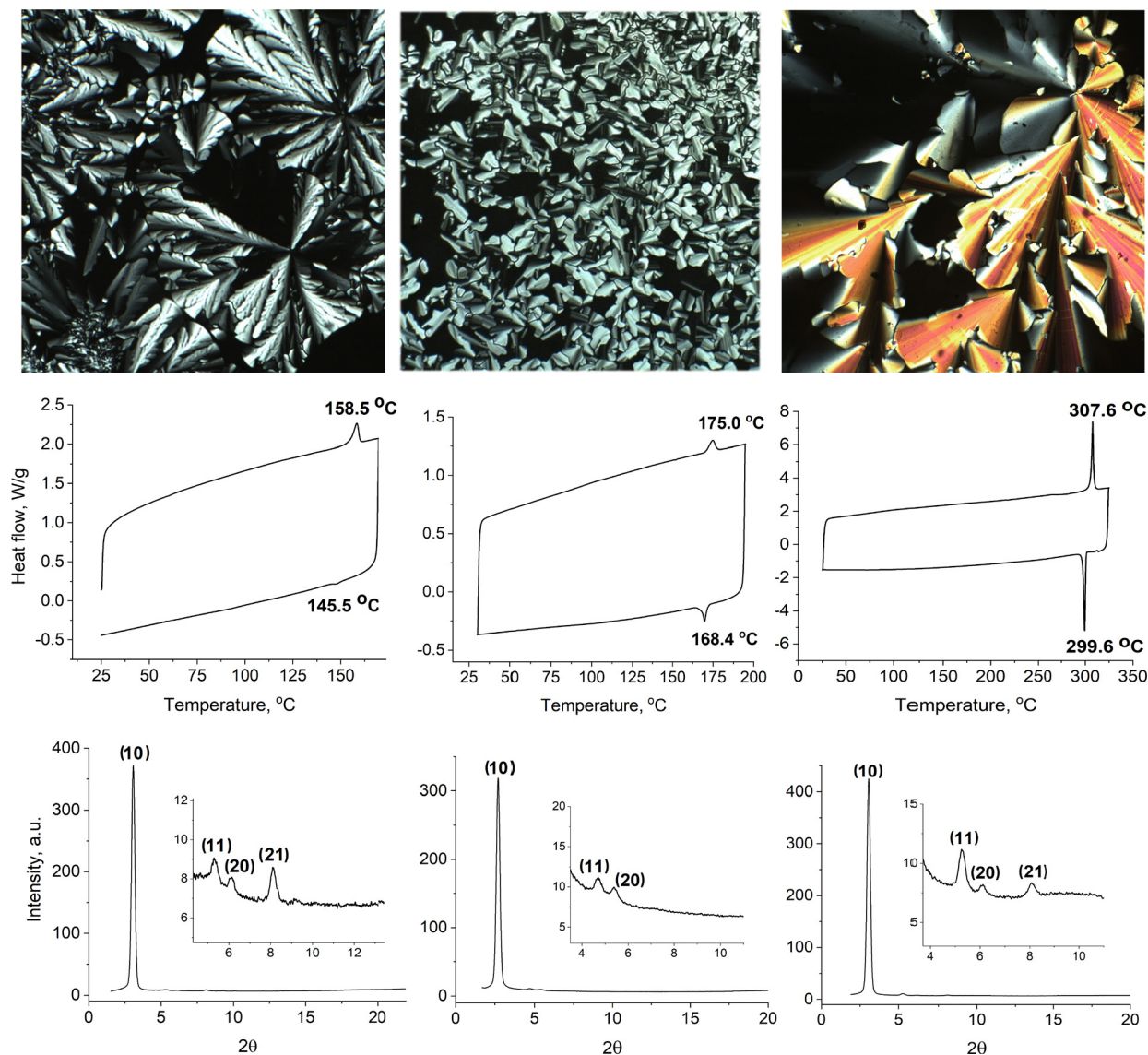


Fig. 3 Liquid crystal characterisation of the new phenanthrolines. Top row (left to right): Polarised optical microscopy (magnification of 200×): **4,7-CzPhen** at 83 °C on cooling; **3,8-CzPhen** at 156 °C on cooling; **2,9-CzPhen** at 304 °C on cooling. Middle row (left to right): DSC traces (at 10 K min⁻¹) **4,7-CzPhen**; **3,8-CzPhen**; **2,9-CzPhen**. Bottom row (left to right): SAXS data **4,7-CzPhen** at 124 °C on heating; **3,8-CzPhen** at 82 °C on heating; **2,9-CzPhen** at 209.6 °C on heating.

Table 1 Indexation of the SAXS reflections along with the d -spacings and the hexagonal lattice parameter, a ($= 2d/\sqrt{3}$)

Compound	Miller indices, (hk)	$d_{\text{obs}}/\text{\AA}$	$a/\text{\AA}$
4,7-CzPhen	10	28.7	33.1
	11	16.6	
	20	14.4	
	21	10.8	
3,8-CzPhen	10	32.5	37.5
	11	18.8	
	20	16.3	
	21	10.9	
2,9-CzPhen	10	28.7	33.1
	11	16.7	
	20	14.5	
	21	10.9	

an appreciably lower onset temperature of 145.5 °C. The difference in the two enthalpy changes is appreciable. When

viewed under the polarising microscope at room temperature, again the material is not crystalline and can be deformed, its pliability increasing with temperature. There is evident birefringence but as expected on heating a sample of this nature, there is no discernible texture. Clearing occurs at ≈ 158 °C and then whatever the rate of cooling (from 10 °C min⁻¹ to 0.1 °C min⁻¹), the texture does not re-appear before *ca.* 145 °C and, on re-heating, persists again to ≈ 158 °C. Obtaining an optical texture on cooling is not, however, straightforward and the one shown in Fig. 3 resulted from a combination of very slow cooling and isothermal annealing.

The degree of hysteresis in the clearing temperature and the difference in the two enthalpy changes is reminiscent of monotropic behaviour, but this is not consistent with the persistence of the mesophase texture to 158 °C on re-heating.



In addition, examination of the SAXS data (Fig. S7, ESI†), collected at 10 °C intervals on heating and cooling, shows behaviour that is 'symmetric' about the clearing temperature, inasmuch as the shape and position of the main (10) reflection at $2\theta \approx 2.7^\circ$ is almost exactly the same at the same temperature on heating and cooling. As such, it seems that the observed hysteresis is due to a slow re-constitution of the columnar structure after clearing, which leads to a broad and supercooled transition.

However, while the clearing point for **4,7-CzPhen** (158.5 °C) is not dissimilar to that of **3,8-CzPhen** (175 °C), both are very much lower than that observed for **2,9-CzPhen**, an observation that has been checked very carefully. In considering why this might be, it is instructive to contemplate how the compounds might self-organise in the columnar phase and this is helped by considering the 3D structures of the complexes, which have been calculated for the methoxy (as opposed to octyloxy)

derivatives as part of the quantum chemical study described below. The complete structures are found as Fig. S8 (ESI†), whereas the dimethoxyphenyl units have been removed from those in Fig. 4 for clarity.

Considering first **3,8-CzPhen** (Fig. 4a), then viewed along an axis containing the 3- and 8-carbons of the ring, it is evident that steric clashes with the 2-, 4-, 7- and 9-ring hydrogen atoms will twist the carbazole rings strongly out of plane ($\approx 53^\circ$), while viewed from above it is easy to see how two molecules might stack one upon the other in pairs being disposed at right angles (the out-of-plane twist of the carbazole groups would prevent a simple stacking arrangement of multiple molecules). In **4,7-CzPhen** (Fig. 4b), once more the carbazole units are also twisted well out of the plane of the phenanthroline ring ($\approx 60^\circ$ – steric clash with ring 3-, 5-, 6- and 8-hydrogen atoms) and in this case the possibility would exist for pairs of molecules to stack, although it is noted that the dipoles of the molecule (calculated

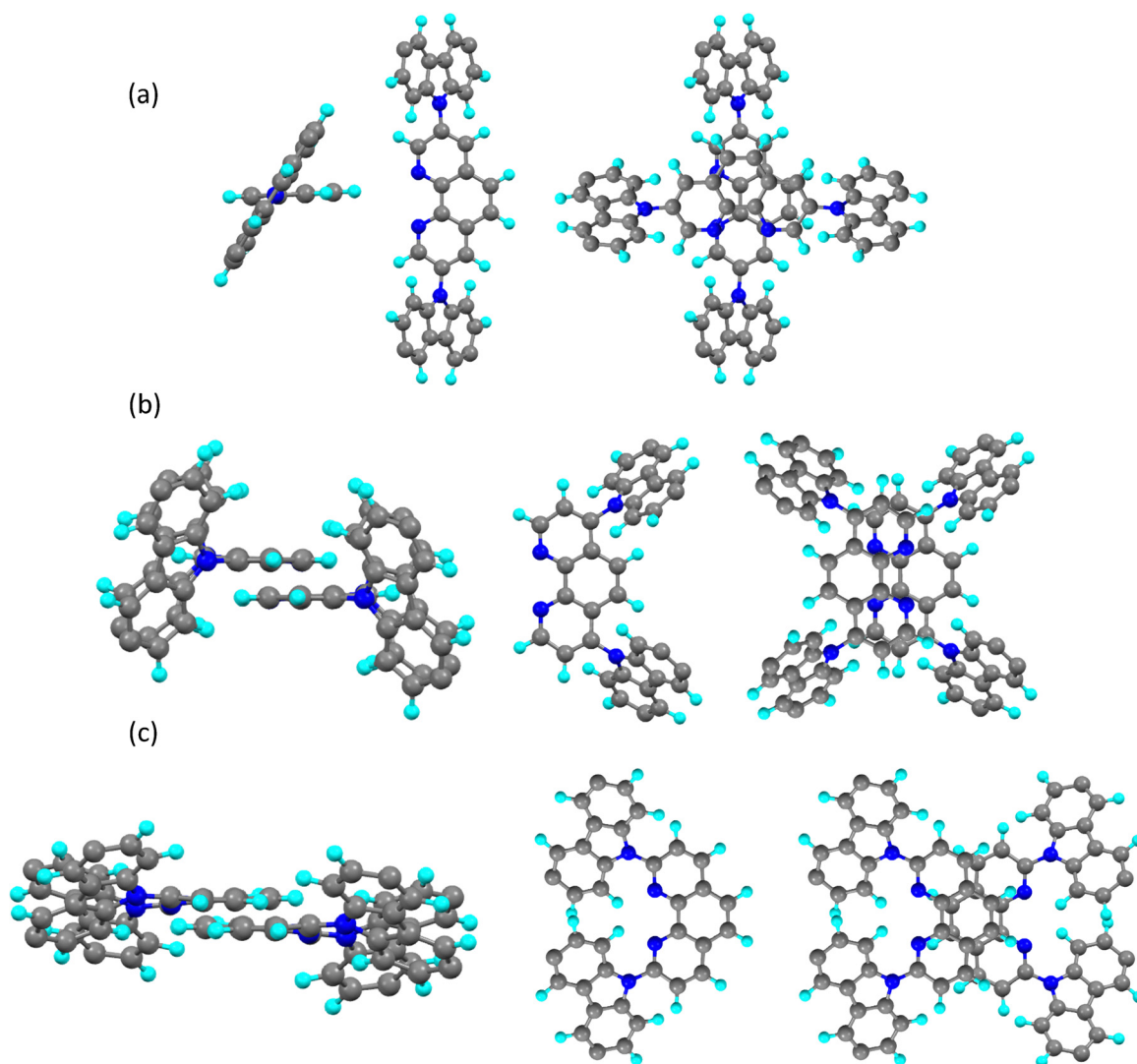


Fig. 4 Calculated structures (both dimethoxyphenyl groups removed from each structure for clarity) for (a) **3,8-CzPhen**, (b) **4,7-CzPhen** and (c) **2,9-CzPhen**. In all three cases, the representations (left to right) are: along the axis containing the 3- and 8-ring carbon atoms; perpendicular to the plane of the phenanthroline; proposed self-organisation of the molecules in the Col_h phase.



as 2.06 D as part of the study below) may in part compromise the stability of this arrangement. It is then reasonable on the basis of these two arrangements that the clearing points are not dissimilar, although the columnar phase of **3,8-CzPhen** would be expected (as observed) to be a little more stable given the slightly better packing available. However, in **2,9-CzPhen** (Fig. 4c), there is only a steric clash with the 3- and 8-ring hydrogen atoms and so the twisting out of plane of the carbazoles is very much less at *ca.* 32°, meaning that a much more compact and anisotropic arrangement of ring–ring dimers is possible and the back-to-back nature of the arrangement means that the very large dipole moments (4.32 D) will be antiparallel, which will stabilise the arrangement. The combination of these factors results in a very much more stable columnar phase.

Finally, it is noted that each of the arrangements shown in Fig. 4 will lead to a repeating unit of very similar lateral dimensions, consistent with the observed hexagonal lattice parameters, which are all rather similar.

Photophysical properties

The three novel compounds were examined for their photophysical properties and key data are collected in Table 2.

3,8-CzPhen. The absorption spectrum of **3,8-CzPhen** in toluene (Fig. 5a) shows an intense band ($\epsilon \approx 11 \times 10^4 \text{ M}^{-1} \text{ cm}^{-1}$) with the maximum at 290 nm, assigned to $\pi \rightarrow \pi^*$ transitions within the core carbazole unit and the 1,10-phenanthroline.^{32–37} A series of less intense bands ($\epsilon \approx 3 \times 10^4 \text{ M}^{-1} \text{ cm}^{-1}$) in the range 340–400 nm are assigned to the $\pi \rightarrow \pi^*$ transitions within the alkoxyaryl carbazole moieties^{19,20,35} and the π -system throughout the molecule. The weak, ($\epsilon \approx 870 \text{ M}^{-1} \text{ cm}^{-1}$) broad band centred at 485 nm is assigned to the charge-transfer transition(s) from the alkoxyaryl carbazole(s) to the 1,10-phenanthroline core.

The emission from a degassed sample ($\approx 10^{-5} \text{ M}$) measured at $T = 300 \text{ K}$ and centred at 440 nm (Fig. 5a) is intense with a quantum yield $\Phi_{\text{PL}} = 40\%$ and is to higher energy than the charge-transfer absorption band (*ca.* 485 nm). This indicates that the observed emission is associated with a state of local

character to one of the chromophores. 1,10-Phenanthroline has been reported to fluoresce at notably shorter wavelengths with the maximum in the range 360–380 nm depending on the solvent,^{33,37} whereas the emission spectrum taken for the precursor alkoxyaryl carbazole (Scheme 1) in toluene appears with a maximum at 397 nm (Fig. S9, ESI†). Therefore, in the light of the above evidence, we conclude that the emission of **3,8-CzPhen** in toluene is associated with transitions on the alkoxyaryl carbazole units (the largest chromophores in the molecule) affected by the central 1,10-phenanthroline unit, rather than arising from the alkoxyaryl carbazole-to-1,10-phenanthroline charge-transfer transition (*vide infra*). The emission at $T = 77 \text{ K}$ in toluene appears in the same spectral range with resolved vibrational progression of 1460 cm^{-1} (Fig. 5a). Such vibrational resolution of emission is indeed characteristic of excited states with local excitation character or, in other words, with a relatively small electron density redistribution within the molecule when compared to the ground state. The decay profile of this emission at $T = 300 \text{ K}$, obtained by time-correlated single photon counting (TCSPC), shows a fast decay component with a time constant of 4.7 ns (Fig. 5b), which is reproduced for both degassed and air-saturated samples. A nanosecond-scale decay time and the absence of sensitivity to molecular oxygen are characteristic of spin-allowed, radiative transitions with high oscillator strength and hence this emission is assigned to the fluorescence stemming from $\pi \rightarrow \pi^*$ character singlet state on alkoxyaryl carbazole unit(s) ($^1[\pi_{\text{Cz}}\pi^*_{\text{Cz}}]$).

Polystyrene (PS) has been chosen as a solid matrix that can suppress charge-separation processes due to its non-polar nature (similar to toluene), thus facilitating efficient emission excited states with local character in competition with relaxation to the lower-lying charge-transfer state. Thus, measured in a doped PS film at loadings much less than 1 wt%, two distinct emission bands are observed with maxima at 420 nm and 570 nm (Fig. 5c), and with a combined emission quantum yield of $\Phi_{\text{PL}}(\text{PS}) = 25\%$. The band centred at 420 nm is assigned as having the same origin as the emission observed in toluene solution, *e.g.* from a state local to an alkoxyaryl carbazole unit,

Table 2 A summary of key photophysical properties of **3,8-CzPhen**, **4,7-CzPhen** and **2,9-CzPhen** in dilute toluene solution ($\approx 10^{-5} \text{ M}$) and in doped polystyrene film ($\ll 1 \text{ wt}\%$) under a nitrogen atmosphere. The absorption spectra were measured under ambient conditions. The solution quantum yield and decay time values are for degassed samples

	3,8-CzPhen	4,7-CzPhen	2,9-CzPhen
Absorption			
$\lambda_{\text{max}}/\text{nm}$	485 (870), 390 (29 000), 290 (110 000)	385 (7500), 355 (14 700), 295 (110 000), 285 (107 000)	390 (30 300), 370 (32 300), 345 (34 400), 295 (102 000)
Photoluminescence in degassed toluene at 300 K			
$\lambda_{\text{max}}/\text{nm}$	440	455	450
$\phi_{\text{PL}}/\%$	40	13	18
τ/ns	4.7	7.5	4.8
Photoluminescence in polystyrene at 300 K			
$\lambda_{\text{max}}/\text{nm}$	420, 570	420, 570	425
$\phi_{\text{PL}}/\%$	25	25	13
τ/ns	2.5 (det. at 420 nm) 5.6 (det. at 570 nm)	5.9 (det. at 420 nm) 21 (det. at 515 nm, fast component)	3.4



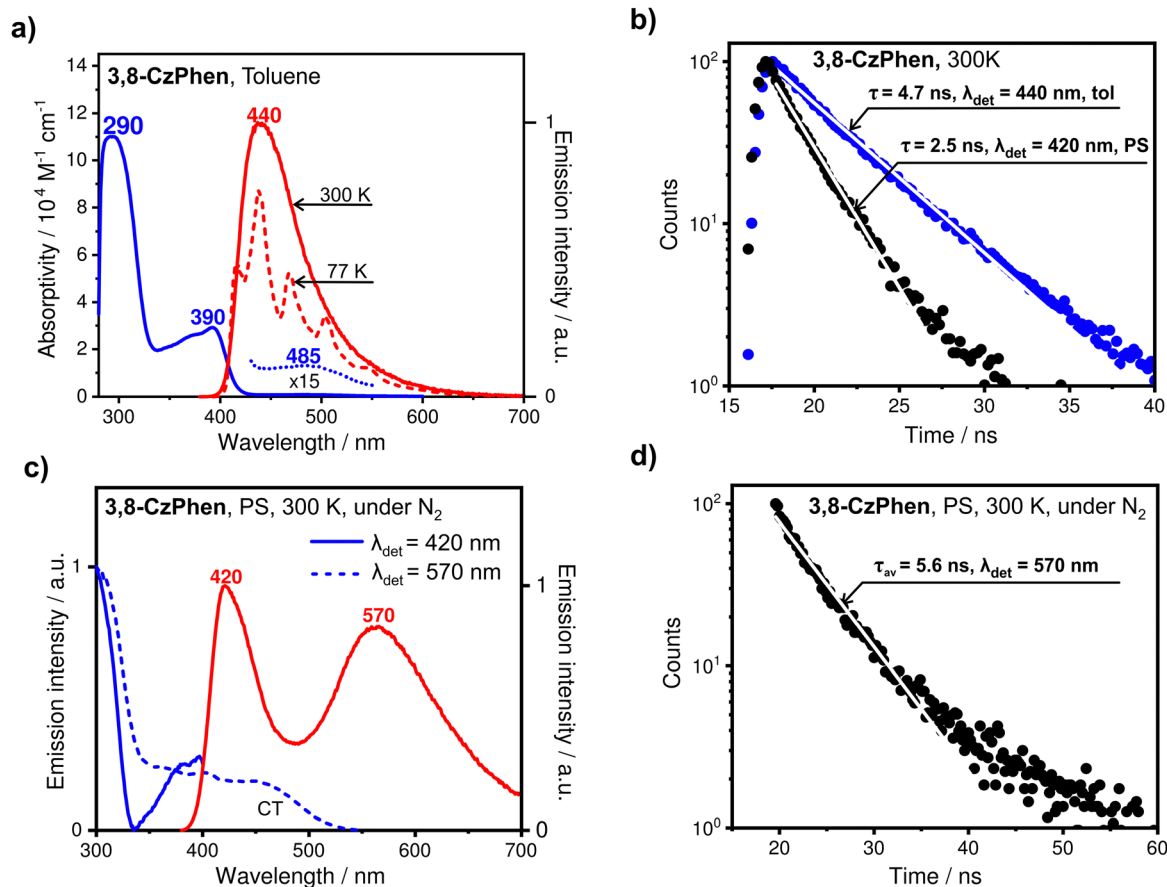


Fig. 5 (a) Absorption (blue) spectrum measured under ambient conditions and emission (red solid for 300 K and red dashed at 77 K) spectra of **3,8-CzPhen** in toluene ($\approx 10^{-5}$ M). (b) Emission decay profile of **3,8-CzPhen** sample in degassed in toluene ($\approx 10^{-5}$ M) measured with detection at 440 nm (blue dots) and in doped PS film ($\ll 1$ wt%) measured with detection at 420 nm, both at ambient temperature. (c) Emission spectrum (red) and excitation spectra of **3,8-CzPhen** doped in polystyrene ($\ll 1$ wt%) measured with detection at 420 nm (solid blue) and at 570 nm (dashed blue). (d) Emission decay profile of **3,8-CzPhen** doped in polystyrene ($\ll 1$ wt%) measured with detection at 570 nm under nitrogen atmosphere at ambient temperature.

but is blue-shifted owing to the different solvation by, and perhaps rigidity of, the polymer film, which limits the reorganisation-driven stabilisation of the emitting excited state. As a further supporting argument for this assignment, we note that the emission spectrum taken for the precursor alkoxyaryl carbazole (Scheme 1) doped in a PS film appears very similar with a slightly blue-shifted maximum at 408 nm (Fig. S9, ESI†). The emission decay profile of the 420 nm-centred emission obtained from TCSPC gives a decay time of 2.5 ns, which is shorter than that measured in degassed toluene solution (Fig. 5b) and may indicate a faster non-radiative relaxation of the emitting state ($^1[\pi_{Cz}\pi^*_{Cz}]$) to the charge-transfer state ($^1[\pi_{Cz}\pi^*_{Phen}]$) in the PS film medium.

The longer-wavelength emission band centred at 570 nm is relatively broad, indicating significant electron density redistribution and, consequently, stronger geometric reorganisation between the emitting and ground states. It is, therefore, assigned to a CT state arising from the transition from an alkoxyaryl carbazole to the 1,10-phenanthroline core, mentioned above as weakly absorbing at *ca.* 485 nm. It is noted that three control experiments, in which the dopant

concentration in the film was consequently halved starting from $c \ll 1$ wt%, did not change the spectral ratio and characteristics of this emission ruling out the involvement of excimers which also would not show up on the absorption spectrum. The TCSPC-measured decay profile of this emission gives a short lifetime of only 5.6 ns (Fig. 5d), indicative of the singlet multiplicity of the emitting state (1CT or $^1[\pi_{Cz}\pi^*_{Phen}]$) as well as, judged from the low intensity of the corresponding absorption band that suggests slow, radiative decay of the CT state as well as its fast and efficient non-radiative decay to the ground state.

The excitation spectra measured for the two emission bands in the polystyrene film at 420 nm and 570 nm, have very similar spectral features (Fig. 5c), evidenced by their features in overlapping range (300–400 nm), which further supports the view that they indeed belong to the same emitting species. The excitation spectrum obtained for the 570 nm emission in addition reproduces the charge-transfer band observed in the absorption spectrum, which, however, now appears with a higher intensity ratio with respect to the shorter wavelength bands. This is because the corresponding transitions evidently



populate the state emitting at 570 nm more efficiently than the higher-energy transitions local to the chromophores. The observation of this CT band in the excitation spectrum further supports the assignment of the emission centred on 570 nm to a charge-transfer $^1[\pi_{\text{Cz}}\pi^*_{\text{phen}}]$ state. Importantly, the excitation spectrum measured with detection at 420 nm has relatively low intensity at *ca.* 335 nm compared both to the absorption spectrum and to the excitation spectrum taken with detection at 570 nm. This shows that the excited states absorbing at *ca.* 335 nm relax relatively efficiently to the charge-transfer state that emits at 570 nm and are more decoupled (weak electronic coupling and/or vibrational overlap with) from the state emitting at 420 nm. The energy of the state populated at 335 nm in the absorption spectrum agrees very closely with the energy of the lowest excited singlet state of 1,10-phenanthroline reported in handbooks³⁶ and with the position of the 0-0 transition in its fluorescence spectrum.^{33,34} This (local) excited state of 1,10-phenanthroline ($^1[\pi_{\text{phen}}\pi^*_{\text{phen}}]$) would then indeed be decoupled from the $^1[\pi_{\text{Cz}}\pi^*_{\text{Cz}}]$ states of the alkoxyaryl carbazole units (which emit at 420 nm) by the wavefunctions of both unpaired particles, the hole and the excited electron, which

then could indeed strongly limit the internal conversion (by Dexter energy transfer) efficiency between them (*vide infra*).

4,7-CzPhen. Measured in toluene and in common with **3,8-CzPhen**, the absorption spectrum of **4,7-CzPhen** shows an intense band at 290–300 nm due to the local $\pi \rightarrow \pi^*$ transitions of the carbazole cores and the 1,10-phenanthroline. Other bands in the range 340–400 nm are once more assigned to the transitions within the alkoxyaryl carbazole units (Fig. 6a) and appear appreciably less intense compared to those in **3,8-CzPhen**. This may be due to the relatively higher steric hindrance of the carbazole substituents caused by the C–H hydrogen atoms in positions 5- and 6- in the 1,10-phenanthroline (Fig. 4 and Fig. S8, ESI[†]), keeping the two substituents more twisted with respect to the heterocyclic core and thus limiting conjugation. Also, in contrast to **3,8-CzPhen**, no charge-transfer transition from the carbazole moieties to the 1,10-phenanthroline core is detected in the absorption spectrum, indicating much lower oscillator strengths of the corresponding charge-transfer transitions in this case. This might be due to the fact that the carbazole substituents in **4,7-CzPhen** are not aligned along a single axis as in **3,8-CzPhen**, so that transition dipole moments

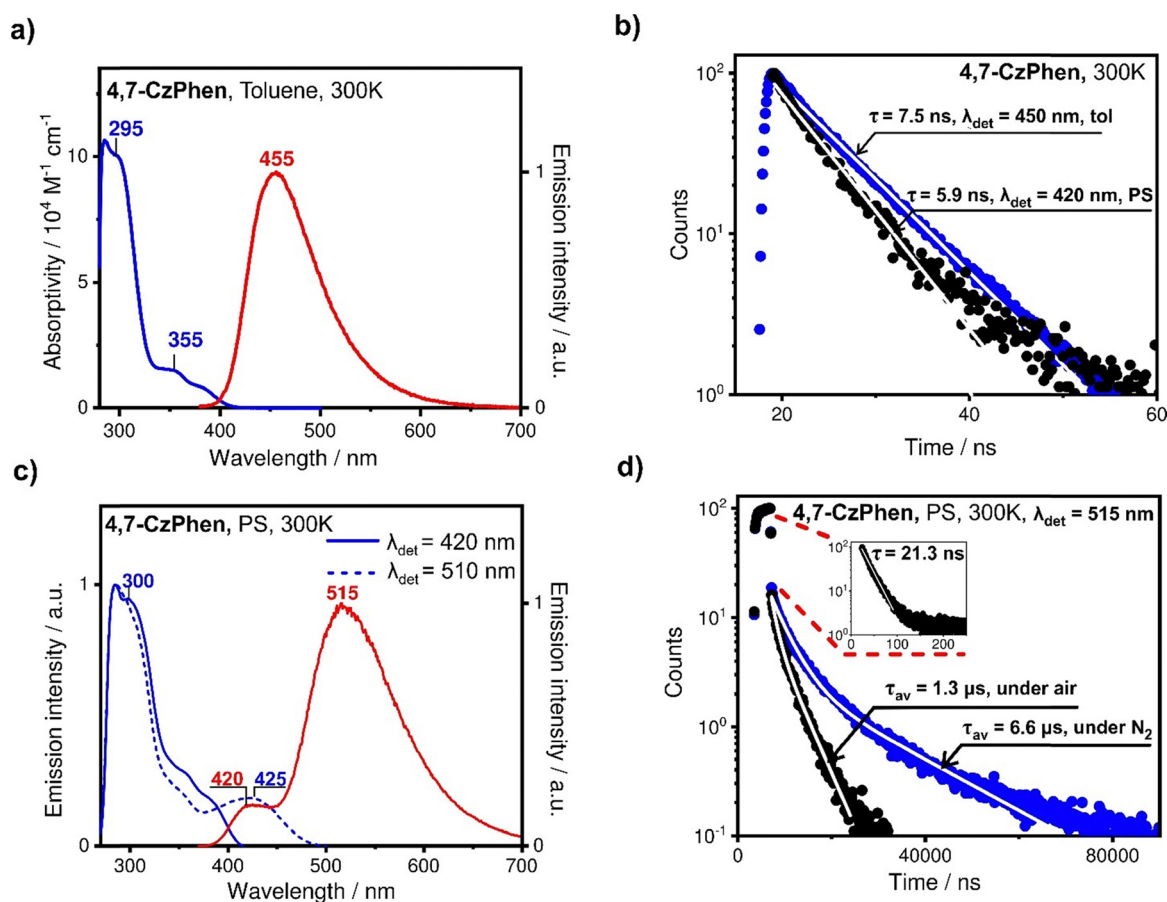


Fig. 6 (a) Absorption (blue) and emission (red) spectra of **4,7-CzPhen** in toluene ($\approx 10^{-5}$ M) under ambient conditions. (b) Emission decay profiles of **4,7-CzPhen** in degassed toluene ($\approx 10^{-5}$ M) taken with detection at 455 nm under ambient temperature (blue dots) and in doped PS film ($\ll 1$ wt%) black dots measured with detection at 420 nm under ambient conditions. (c) Emission spectrum (red) and excitation spectra of **4,7-CzPhen** doped in polystyrene ($\ll 1$ wt%) measured with detection at 420 nm (solid blue) and at 510 nm (dashed blue). (d) Emission decay profiles of **4,7-CzPhen** doped in polystyrene ($\ll 1$ wt%) at ambient temperature under air (black dots) and under nitrogen atmosphere (blue dots) measured with detection at 515 nm.



of the two sides cannot sum up as efficiently, as well as the fact that they are more twisted with respect to 1,10-phenanthroline. The combination of these factors could well act to further diminish the transition dipole moment of these charge-transfer transitions.

The emission spectrum of **4,7-CzPhen** in toluene under ambient conditions is centred at 455 nm and is characterised by a TCSPC-measured decay time of 7.5 ns, assigned to fluorescence stemming from a state local to the alkoxyaryl carbazole moieties ($^1[\pi_{Cz}\pi^*_{Cz}]$). In a doped polystyrene film, emission is observed from two distinct bands centred at 420 nm and 515 nm (Fig. 6c), the former being assigned to the same emission observed in toluene although blue shifted in the PS film. The emission decay time of $\tau = 5.9$ ns of the band at 420 nm is close to that in toluene solution. The broad emission band centred on 515 nm is assigned to charge-transfer state(s) arising from transitions from the alkoxyaryl carbazole moiety to the 1,10-phenanthroline ($[\pi_{Cz}\pi^*_{phen}]$) and the decay curve consists of two components (Fig. 6d). The fast component, assigned to prompt fluorescence from the $^1[\pi_{Cz}\pi^*_{phen}]$ CT state, accounts for about 95% of the overall intensity and has a decay time of 21 ns (Fig. 6d inset), while the slow component shows non-monoexponential behaviour and is oxygen sensitive. The biexponential fit of this slow component gives the intensity-averaged decay time of 1.3 μ s in air-saturated conditions, increasing to 6.6 μ s under a nitrogen atmosphere (Fig. 6d). The sensitivity of the slow component to oxygen and the microsecond decay time implies involvement of a triplet state in the emission mechanism. This is most likely to be thermally activated delayed fluorescence (TADF), as was reported for analogous *N*-phenoxazine substituted phenanthrolines.²⁷ Here, due to the weak exchange interaction in the charge-transfer state, which defines the energy separation of singlet and triplet states of the same orbital origin,³⁸ the $^1[\pi_{Cz}\pi^*_{phen}]$ singlet and $^1[\pi_{Cz}\pi^*_{phen}]$ triplet states appear energetically close to degenerate and can be equilibrated *via* ISC and thermally activated reverse intersystem crossing (RISC) processes. It is noted that, typical for organic compounds, ISC and RISC processes between the singlet and triplet state are probably too slow for fast thermal equilibration, so that on the TCSPC curve, TADF appears as a separate decay component with a small, almost negligible, share of the intensity amounting to only 5% of the total measured intensity in deoxygenated conditions. We note that in principle, this slow component of the decay curve could also arise from phosphorescence from a lower-lying triplet state, although in this scenario a much longer decay time would be expected and it would be also observed for **3,8-CzPhen**, so this is regarded as unlikely. It is noted that the relatively long decay time of the fast component and the appearance of a slow component involving a triplet state, is indicative of relatively slow and inefficient non-radiative decay of the S_1 ($^1[\pi_{Cz}\pi^*_{phen}]$) state and perhaps T_1 ($^3[\pi_{Cz}\pi^*_{phen}]$) state of **4,7-CzPhen** to the ground state in a PS film as compared to **3,8-CzPhen** compound. This may be related to the higher energy of the CT states ($^1[\pi_{Cz}\pi^*_{phen}]$ and $^3[\pi_{Cz}\pi^*_{phen}]$) of **4,7-CzPhen** resulting in their relatively weaker vibrational overlap

with the ground state. Indeed, emission from the $^1[\pi_{Cz}\pi^*_{phen}]$ state of **4,7-CzPhen** is centred at 515 nm and is blue shifted compared to that of **3,8-CzPhen** (570 nm), which may also be due to the more twisted geometry and consequently limited conjugation in **4,7-CzPhen**, thus resulting in a relatively destabilised CT state. In common with the observations for **3,8-CzPhen**, the excitation spectra of the two emission bands at 420 nm and 515 nm, are found similar in the range where each is measured (Fig. 6c), again showing that both originate from the same species. It is noted that the excitation spectrum of the emission centred at 420 nm does not reduce at *ca.* 335 nm to the extent found in **3,8-CzPhen**, which might be due to the overlap of the excitation transition of the lowest local singlet state of 1,10-phenanthroline with bands of other transitions involving the alkoxyaryl carbazole moieties or faster Dexter energy transfer between local states of alkoxyaryl carbazole moieties and of 1,10-phenanthroline, so that internal conversion between them can be relatively efficient. Finally, the excitation spectrum obtained with detection at 515 nm reveals the charge-transfer transition band, centred at *ca.* 425 nm (Fig. 6c), which was not detectable in the absorption spectrum.

Interestingly, the emission quantum yield of **4,7-CzPhen** increases from 13% in degassed toluene to 50% in a PS film under nitrogen, in contrast to the case of **3,8-CzPhen** where the opposite trend was found. Apparently in **4,7-CzPhen**, the PS film suppresses the non-radiative decay processes from the lower-energy charge-transfer state(s) to the ground state much more efficiently and the increase in overall emission quantum yield is associated with drastically increased efficiency of emission from the CT $^1[\pi_{Cz}\pi^*_{phen}]$ state. Apart from the effect of rigid media of the PS film, the higher energy of the CT state in **4,7-CzPhen** compared to that in **3,8-CzPhen**, may result in a slower and less efficient non-radiative decay to the ground state in accordance with the energy gap law.³⁹ The lower efficiency of the non-radiative decay is also reflected in a significantly longer decay time of the lower-energy emission of **4,7-CzPhen** in PS (515 nm, 21 ns), as compared to **3,8-CzPhen** (570 nm, 5.6 ns) (compare Fig. 5b and 6b). Notably, the decay time of the emission stemming from the $^1[\pi_{Cz}\pi^*_{Cz}]$ state of **4,7-CzPhen**, local to alkoxyaryl carbazole units (420 nm), does not decrease as significantly from a toluene solution to a PS film as in the case of **3,8-CzPhen**, indicating that there is less reduction in emission efficiency of the $^1[\pi_{Cz}\pi^*_{Cz}]$ state from internal conversion.

2,9-CzPhen. In toluene solution at room temperature, this isomer shows photophysical properties reminiscent of both **3,8-CzPhen** and **4,7-CzPhen**. Its absorption bands measured in toluene (Fig. 7a) are similar to, and interpreted in the same way as, those of **3,8-CzPhen**, while in common with **4,7-CzPhen**, the absorption spectrum features no detectable charge-transfer transitions from the alkoxyaryl carbazoles to the phenanthroline core. Under ambient conditions, the same toluene solutions exhibit an emission centred at 450 (Fig. 7a) with a quantum yield of 18% and a decay time of 4.8 ns, which is blue shifted to 425 nm with a quantum yield of 13% and a decay time of 3.4 ns in a doped PS film (Fig. 7b and c). In



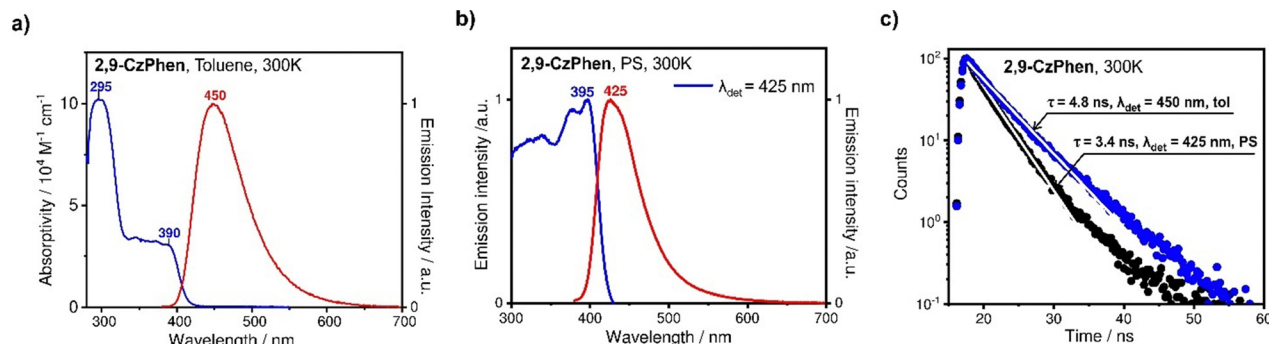


Fig. 7 (a) Absorption (blue) and emission (red) spectra of **2,9-CzPhen** in toluene ($\approx 10^{-5}$ M) under ambient conditions. (b) Emission (red) and excitation (blue) spectra of **2,9-CzPhen** doped in PS film ($\ll 1$ wt%). (c) Emission decay profiles of **2,9-CzPhen** in toluene ($\approx 10^{-5}$ M) at ambient temperature (blue dots) and as doped in PS film (black dots) ($\ll 1$ wt%).

common with the other isomers, this emission is assigned to a singlet state localised on the alkoxyaryl carbazole. Notably however, **2,9-CzPhen** does not show additional emission assignable to $^1[\pi_{Cz}\pi^*_{phen}]$ CT states in a PS film, as apparently its less twisted geometry (compared to **4,7-CzPhen** and **3,8-CzPhen**) pushes the lowest $^1[\pi_{Cz}\pi^*_{phen}]$ state to even lower energies so that its non-radiative decay to the ground state remains very efficient even in the PS film.

Quantum chemical calculations

Electronic structures of the three materials were examined with quantum chemical calculations using a DFT approach. To simplify the calculations, the alkoxy chains were truncated to methoxy groups and the corresponding structures are denoted as **2,9-CzPhen'**, **3,8-CzPhen'**, **4,7-CzPhen'**. Calculations were carried out using Gaussian G09 code⁴⁰ with the M11L^{41,42} DFT functional, the def2-SVP basis set^{41,43} and the C-PCM solvation model⁴⁴ using parameters for toluene. The molecules were optimised in their ground-state electronic configuration and the given excited state energies represent vertical excitation energies with respect to the ground state. Iso-surface contour plots for the frontier orbitals are collected in Fig. 8, while Cartesian coordinates of the atomic positions for the optimised geometries are given in the ESI†

3,8-CzPhen'. The S_1 state (2.18 eV (568 nm), $f = 0.0045$) is of HOMO \rightarrow LUMO origin and has $^1[\pi_{Cz}\pi^*_{phen}]$ character, while the T_1 state (2.15 eV, 577 nm), also HOMO \rightarrow LUMO in origin ($^3[\pi_{Cz}\pi^*_{phen}]$), is calculated as the only excited state lying below the S_1 state. Since these two states have the same orbital origin, $S_1 \rightarrow T_1$ ISC can indeed be very slow and inefficient as predicted by the El-Sayed's rule.⁴⁵ The decent oscillator strength calculated for the S_1 state, which is consistent with a state of strong CT character, should make the $S_1 \rightarrow S_0$ fluorescence of **3,8-CzPhen** relatively efficient, which indeed is seen from relatively prominent CT absorption band (485 nm) with the corresponding fluorescence becoming observable centred at 570 nm in a PS film. The lowest energy excited singlet state of $^1[\pi_{Cz}\pi^*_{Cz}]$ character, to which the 420 nm emission of **3,8-CzPhen'** in PS film is assigned, is calculated as S_{10} at this geometry with vertical excitation energy of 2.97 eV (417 nm, $f = 0.0011$) and has

HOMO \rightarrow LUMO+2 (72%) and HOMO-1 \rightarrow LUMO+2 (26%) origin.

4,7-CzPhen'. The S_1 state (2.182 eV (568 nm), $f = 0.0243$) of **4,7-CzPhen'** is, as expected, calculated to be HOMO \rightarrow LUMO in origin and, according to orbital localisations (Fig. 9), has $^1[\pi_{Cz}\pi^*_{phen}]$ character. The triplet state T_1 (2.167 eV (572 nm)) is also HOMO \rightarrow LUMO in origin with $^3[\pi_{Cz}\pi^*_{phen}]$ character. It is notable that the energy of the S_1 state is slightly underestimated compared to the spectroscopic data obtained in a PS film. The calculated energy gap $\Delta E(S_1-T_1) = 0.015$ eV (15 meV, 121 cm^{-1}) is very small. Interestingly, the second triplet state T_2 (2.181 eV (567 nm)) of HOMO-1 \rightarrow LUMO origin is calculated to be located between states S_1 and T_1 . Thus, with states S_1 and T_2 having different orbital origin and with a very small energy gap between them, ISC $S_1 \rightarrow T_2$ could indeed be relatively efficient. This agrees well with the observation of a microsecond decay component, although with a very small intensity share (*ca.* 5%), of the 515 nm emission of **4,7-CzPhen** in a PS film, discussed above and interpreted as TADF.

2,9-CzPhen'. The lowest excited singlet state, S_1 (2.29 eV (540 nm), $f = 0.0000$) is calculated to have the HOMO \rightarrow LUMO transition origin. Thus, according to the orbital localisations (Fig. 8), the S_1 state has charge-transfer character from the alkoxyaryl carbazoles to the 1,10-phenanthroline, $^1[\pi_{Cz}\pi^*_{phen}]$. The T_1 state, also being HOMO \rightarrow LUMO in origin, is calculated at 2.23 eV and thus is 60 meV (480 cm^{-1}) below the S_1 state. Calculations also suggest that T_2 (2.25 eV, HOMO \rightarrow LUMO+1) and T_3 (2.28 eV, HOMO-1 \rightarrow LUMO) states exist between S_1 and T_1 . The presence of nine states (*i.e.* three triplets) lying close in energy and below the S_1 state could indeed result in a fast, efficient non-radiative decay of the S_1 state *via* ISC pathways to those lower triplet states and then further to the ground state. The efficiency of this non-radiative decay is further enhanced by poor competition from the $S_1 \rightarrow S_0$ fluorescence, calculated to have a vanishingly small oscillator strength. Perhaps as a result of this combination of properties, **2,9-CzPhen** shows no fluorescence assignable to $^1[\pi_{Cz}\pi^*_{phen}]$ state ($S_1 \rightarrow S_0$) even in a PS matrix. Notably, the lowest-energy excited singlet state with assignable $^1[\pi_{Cz}\pi^*_{Cz}]$ character (to which the observed fluorescence of **2,9-CzPhen** is



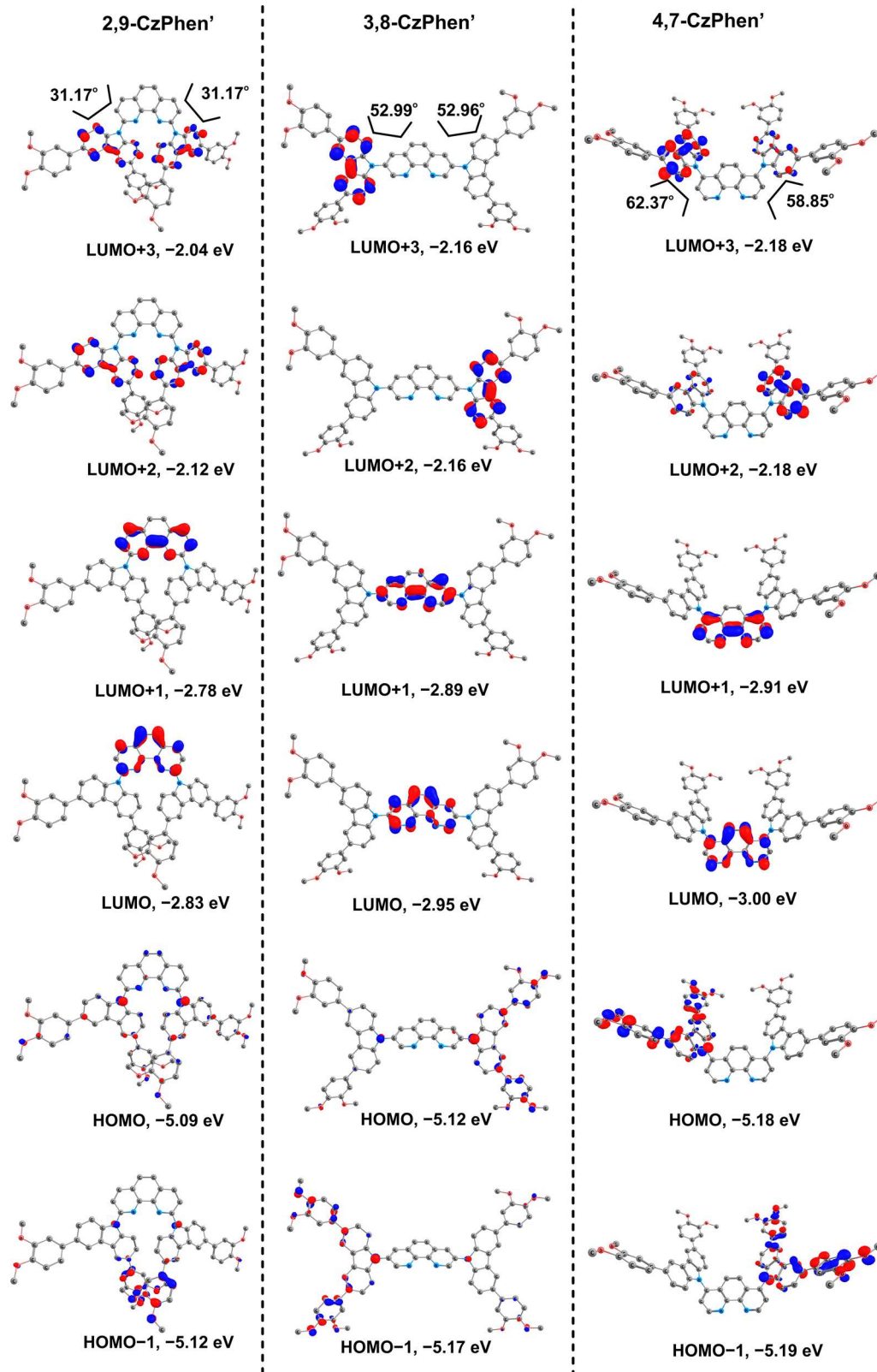


Fig. 8 Iso-surface contour plots (iso-value = 0.05) of the frontier orbitals of 2,9-CzPhen', 3,8-CzPhen', 4,7-CzPhen', each in its optimised ground state geometry. Hydrogen atoms are omitted for clarity.



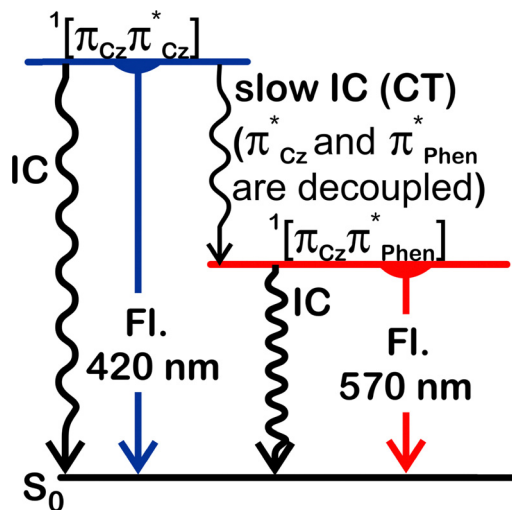


Fig. 9 Simplified Jablonski diagram summarising luminescent properties of **3,8-CzPhen** in a PS film (Fl. = fluorescence). A slightly more detailed diagram is found in the ESI† (Fig. S10).

assigned) was calculated to be a S_{10} state (3.00 eV (412 nm), $f = 0.0011$) of $\text{HOMO} \rightarrow \text{LUMO}+2$ (82%) and $\text{HOMO}-1 \rightarrow \text{LUMO}+3$ (16%) origin.

It is noted that this computational result is specific to the level of theory employed and that the actual character and properties of the $^1[\pi_{\text{Cz}}\pi^*_{\text{Cz}}]$ emitting state of **2,9-CzPhen** will be modified by the relaxation and the specific conditions of the experiment.

Discussion

The photophysics of the three isomeric compounds are rich and also somewhat involved. Therefore, in order to understand the experimental observations and contextualise them in terms of the results of the quantum chemical calculations, it is helpful to appreciate the theory underlying the different processes involved. The discussion that follows is predicated on the properties of **3,8-CzPhen**, from which the behaviour of the other isomers may be understood and relates to the Jablonski diagram in Fig. 9, which shows the different emission and deactivation pathways.

The appearance of two spectrally well-separated fluorescence emissions in a rigid polymer medium, where excited-state molecular geometry reorganisation is relatively limited, is indicative of a limited rate of internal conversion (IC) of the higher-energy emitting state. Indeed, the fact that the absorption band assigned to charge transfer from the alkoxyaryl carbazoles to the phenanthroline has almost vanishing intensity in **3,8-CzPhen** (485 nm) and is so weak that it is not detected in **4,7-CzPhen** and **2,9-CzPhen**, points to the fact that the two units are electronically well decoupled in all three isomers. Note that the CT transition $S_0 \rightarrow ^1[\pi_{\text{Cz}}\pi^*_{\text{phen}}]$ in **4,7-CzPhen** has been detected only in the excitation spectrum taken for 515 nm emission in PS film. Consequently, the rate of internal conversion processes requiring electron transfer(s)

between the chromophores, such as $^1[\pi_{\text{Cz}}\pi^*_{\text{Cz}}] \rightarrow ^1[\pi_{\text{Cz}}\pi^*_{\text{phen}}]$, can indeed be strongly diminished in agreement with the observation of dual fluorescence, $^1[\pi_{\text{Cz}}\pi^*_{\text{Cz}}] \rightarrow S_0$ and $^1[\pi_{\text{Cz}}\pi^*_{\text{phen}}] \rightarrow S_0$, in the PS film (see also ref. 18).

The internal conversion between local state of 1,10-phenanthroline and a local state of an alkoxyaryl carbazole unit – $^1[\pi_{\text{phen}}\pi^*_{\text{phen}}] \rightarrow ^1[\pi_{\text{Cz}}\pi^*_{\text{Cz}}]$ – requires that not just one but two electrons are exchanged between the two decoupled chromophores (Dexter energy transfer) and therefore that the rate of such process can also be very low. This is especially true in a relatively rigid and π -stacked medium within a PS film, where both vibration and the reorganisation of the molecular geometry are restricted. Thus, the reduction in intensity of the peak at ca. 335 nm to near zero in the excitation spectrum obtained for **3,8-CzPhen** in a PS film with detection at 420 nm, further supports the earlier assignment of the 420 nm-centred emission as the $^1[\pi_{\text{Cz}}\pi^*_{\text{Cz}}]$ states of the alkoxyaryl carbazole units. For comparison, in order that the $^1\pi_{\text{Cz}}\pi^*_{\text{phen}}$ CT state of **3,8-CzPhen**, which emits at 570 nm, is populated from the $^1[\pi_{\text{phen}}\pi^*_{\text{phen}}]$ state of 1,10-phenanthroline requires a transfer of a single electron between the two decoupled chromophores and that internal conversion of $^1[\pi_{\text{phen}}\pi^*_{\text{phen}}] \rightarrow ^1[\pi_{\text{Cz}}\pi^*_{\text{phen}}]$ is more efficient than $^1[\pi_{\text{phen}}\pi^*_{\text{phen}}] \rightarrow ^1[\pi_{\text{Cz}}\pi^*_{\text{Cz}}]$. This is indeed seen from comparison of excitation spectra taken using detection at 420 nm and 570 nm (Fig. 6c).

The electronic decoupling of the $^1[\pi_{\text{Cz}}\pi^*_{\text{Cz}}]$ and $^1[\pi_{\text{Cz}}\pi^*_{\text{phen}}]$ states is probably the result not only of two chromophores being twisted with respect to one other, but also a contribution from orbital topology. Thus, consider the lowest unoccupied orbitals of the alkoxyaryl carbazoles (π^*_{Cz}) and those of phenanthroline (π^*_{phen}) – the orbitals whose mutual overlap would mediate the electron transfer in the internal conversion from the $^1[\pi_{\text{Cz}}\pi^*_{\text{Cz}}]$ state to the CT $^1[\pi_{\text{Cz}}\pi^*_{\text{phen}}]$ state(s). The lowest unoccupied orbitals of the alkoxyaryl carbazole units (LUMO+2 and LUMO+3 orbitals in Fig. 8) have a node at the carbazole nitrogen atom – the linking point with the phenanthroline core – and so do those on the phenanthroline (LUMO and LUMO+1). Thus the lowest unoccupied orbitals of the two chromophores are isolated and do not overlap. Since this is found in the calculations for all three compounds, it is evidently not affected by differences in the twist angle between the alkoxyaryl carbazole and phenanthroline chromophores. This indicates that the vibronic coupling of the electronic wavefunctions of the two states ($^1[\pi_{\text{Cz}}\pi^*_{\text{Cz}}]$ and $^1[\pi_{\text{Cz}}\pi^*_{\text{phen}}]$), which arise from vibrations along the dihedral angles between the chromophores and could in principle greatly enhance the internal conversion rate by charge transfer compared to that predicted by a Franck-Condon approximation, cannot be significant.

While the appearance of the 570 nm-centred emission of **3,8-CzPhen** in a PS film indicates lower efficiency of the non-radiative decay of the CT $^1\pi_{\text{Cz}}\pi^*_{\text{phen}}$ state due in part to the rigidity of the medium, the observed quantum yield of 25% is appreciably lower than the 40% found in toluene solution. This would be consistent with increased efficiency of internal conversion to the lower-energy, emitting $^1[\pi_{\text{Cz}}\pi^*_{\text{phen}}]$ state with concomitant reduction in the emission efficiency of the



higher-energy $^1[\pi_{Cz}\pi^*_{Cz}]$ state. To understand why this might be, the processes that define the quantum yield of a system of two emitting states is analysed. Thus, the emission quantum yield for fluorescence (Φ_{PL}) from an individual excited state is defined as the ratio of its rate of fluorescence (k_r) to the sum of all decay rates radiative and non-radiative with $\sum k_{nr}$ – representing the sum of the rates of all non-radiative decay paths so that $\Phi_{PL} = k_r/(k_r + \sum k_{nr})$. Hence, with the assumption that the non-radiative decay of the $^1[\pi_{Cz}\pi^*_{Cz}]$ state consists both of a direct, non-radiative decay pathway to the ground state and of internal conversion leading ultimately to population in the lower energy emitting $^1[\pi_{Cz}\pi^*_{phen}]$ state, the overall quantum yield $\Phi_{PL(tot)}$ of the system of these two states, initially excited to the $^1[\pi_{Cz}\pi^*_{Cz}]$ state, can be approximated as shown in eqn (1):

$$\Phi_{PL(tot)} = (1 - \theta_{IC})(k_r/(k_r + k_{nr})) + \theta_{IC}(k'_r/(k'_r + \sum k'_{nr})) \quad (1)$$

where k_r and k_{nr} are the rates of fluorescence and of direct non-radiative decay to the ground state, respectively, of the higher-energy emitting state $^1[\pi_{Cz}\pi^*_{Cz}]$; k'_r and $\sum k'_{nr}$ are the rates of fluorescence and sum of all non-radiative decay rates, respectively, of the lower-energy $^1[\pi_{Cz}\pi^*_{phen}]$ emitting state. θ_{IC} is the efficiency of the internal $^1[\pi_{Cz}\pi^*_{Cz}] \rightarrow ^1[\pi_{Cz}\pi^*_{phen}]$ conversion, which is in turn defined as $\theta_{IC} = k_{IC}/(k_{IC} + k_r + k_{nr})$, with k_{IC} being the rate of this internal conversion.

Indeed, the rate of fluorescence from this lower-energy, $^1[\pi_{Cz}\pi^*_{phen}]$ emitting state of CT character is expected to be much smaller in comparison to that from the higher-energy $^1[\pi_{Cz}\pi^*_{Cz}]$ emitting state, as can be judged from the relevant transition intensities in the absorption spectrum. This is due both to poorer overlap of the excited-state and ground-state wavefunctions due to spatial separation of the hole and electron, and a smaller value for Einstein's spontaneous emission coefficient,⁴⁶ which is proportional to the cube of the emission frequency. In addition, the rate of non-radiative relaxation directly to the ground state is expected to be higher for the lower-energy $^1[\pi_{Cz}\pi^*_{phen}]$ state, due to the stronger vibrational overlap of the lower energy state with the ground state, as expressed through the energy-gap law.³⁹ The CT character of the $^1[\pi_{Cz}\pi^*_{phen}]$ state will further enhance its vibrational overlap with the ground state due to the more extensive geometry reorganisation, when compared to that in the local character states. This is consequent on the significant redistribution of electron density in the charge-transfer state, such as $^1[\pi_{Cz}\pi^*_{phen}]$, that is manifest by the broad emission band. Hence this further increases the non-radiative decay rate (k_{nr}) of the $^1[\pi_{Cz}\pi^*_{phen}]$ state to the ground state. Further, in addition to the direct decay to the ground state, the $^1[\pi_{Cz}\pi^*_{phen}]$ state can be energetically close to the triplet $^3[\pi_{Cz}\pi^*_{phen}]$ state due to the small exchange energy between the states of CT character, which may add an additional non-radiative decay pathway *via* intersystem crossing. Therefore, for **3,8-CzPhen** with $^1[\pi_{Cz}\pi^*_{Cz}]$ and $^1[\pi_{Cz}\pi^*_{phen}]$ states, it is expected that in eqn (1), $k_r \gg k'_r$ and $k_{nr} \ll \sum k'_{nr}$. Then the overall quantum yield of the system is expected to be greatest when the internal

conversion from $^1[\pi_{Cz}\pi^*_{Cz}] \rightarrow ^1[\pi_{Cz}\pi^*_{phen}]$ shows negligible efficiency, *i.e.* $\theta_{IC} \approx 0$ in eqn (1). Thus, as θ_{IC} increases, the overall emission quantum yield of the system is expected to decrease. Indeed, the observed decrease in the decay time of the higher-energy $^1[\pi_{Cz}\pi^*_{Cz}]$ emitting state from 4.7 ns in a toluene solution to 2.5 ns in a PS film (Fig. 6b and d) might be associated with an increase in the rate of internal conversion (greater k_{IC}) and greater efficiency (greater θ_{IC}) for $^1[\pi_{Cz}\pi^*_{Cz}] \rightarrow ^1[\pi_{Cz}\pi^*_{phen}]$ in the PS film. This could be due to the rigid medium of the PS film affecting the relaxed geometries and energies of the two excited states resulting in better vibrational overlap between them. As a result, the loss of quantum yield from the higher-energy $^1[\pi_{Cz}\pi^*_{Cz}]$ emitting state due to the increase of the internal conversion efficiency in the PS film, is not fully compensated by the quantum yield of the lower-energy emitting state. Consequently, the total fluorescence quantum yield of the system is lower compared to that in toluene solution.

An important implication of the appearance of the two spectrally well-separated fluorescence emissions, where the higher-energy fluorescence from the $^1[\pi_{Cz}\pi^*_{Cz}]$ state has to compete with the internal conversion to the lower-energy, electronically decoupled $^1[\pi_{Cz}\pi^*_{phen}]$ state, is that the $^1[\pi_{Cz}\pi^*_{Cz}]$ emitting state in its own relaxed geometry is probably not the lowest excited state of singlet multiplicity. This would correspond to the textbook definition of the rare example where Kasha's rule³ is not followed (so-called anti-Kasha emission). Indeed, the three compounds exhibit fluorescence stemming from the $^1[\pi_{Cz}\pi^*_{Cz}]$ state in the relatively rigid medium of a PS film. This behaviour includes **4,7-CzPhen** and **2,9-CzPhen** where, due to the relatively greater steric hinderance in the former and due to the π - π overlap of the alkoxyaryl carbazole substituent in the latter, geometric reorganisations in the excited state are restricted compared to the case of **3,8-CzPhen**. Moreover, **3,8-CzPhen** also shows $^1[\pi_{Cz}\pi^*_{Cz}]$ fluorescence in the medium of frozen toluene, where excited-state geometry reorganisations are also diminished. These evidence that the $^1[\pi_{Cz}\pi^*_{Cz}]$ state emits while remaining in the Marcus inverted region with respect to the lower energy $^1[\pi_{Cz}\pi^*_{phen}]$ charge-transfer state and that its emission is indeed anti-Kasha in nature. The slow internal conversion of $^1[\pi_{Cz}\pi^*_{Cz}]$ to the charge-transfer state, affording dual fluorescence, is then interpreted as the result of both electronic decoupling of the states, as discussed above, as well as diminished vibrational overlap of states due to large energy gap in the Marcus inverted region (energy gap law).

Device performance

While dominated by singlet emission and not capable of efficient utilisation of triplet excitons (statistically 75% of all excitons), the observation of dual fluorescence (white emission) in two of the compounds prompted the fabrication of solution-processed OLED devices to explore their electroluminescent (EL) properties. The configuration used was: ITO/poly(3,4-ethylenedioxythiophene):poly(styrene sulfonate) (PEDOT:PSS) (40 nm)/emitter layer (50 nm)/1,3,5-tris(3-pyridyl-3-phenyl)benzene



(TmPyPB) (45 nm)/LiF (0.5 nm)/Al (120 nm) (Fig. 10a). In this device structure, PEDOT:PSS and LiF are used as the hole-injection layer and the electron-injection layer, respectively while TmPyPB is used as the electron-transport layer. Lithium fluoride (LiF) and aluminum (Al) were utilised as electron-injection layer and cathode, respectively. The host matrix of the emitting layer is a blend of poly(*N*-vinylcarbazole) (PVK) and 1,3-bis-(5-(4-*tert*-butylphenyl)-1,3,4-oxadiazol-2-yl)benzene (OXD-7), used in a 7:3 ratio. The dopant concentrations used were 1 and 3 wt%, with the optimised emitter concentration being established as 1 wt%. The relevant EL data are summarised in Table S4 (ESI†).

As shown in Fig. 10b, all devices emit in the blue region of the spectrum with emission peaks in the range 440–470 nm, assignable to $^1[\pi_{Cz}\pi^*_{Cz}]$ state emission. The CIE coordinates are correspondingly found (0.25 ± 0.02 , 0.29 ± 0.03) (Table S4, ESI†). The EL spectra demonstrate that there is a complete energy transfer from the host matrix to the dopant, with both electron and hole ending up on the alkoxyaryl carbazole chromophore, and reasonable turn-on voltages of between 6.0 and 8.0 V are observed for the devices.

When the emitter concentration was increased to 3 wt%, both 2,9-CzPhen and 3,8-CzPhen showed a decreased device

performance, probably due to concentration quenching and so all data were collected using 1 wt% of emitter. Thus, consistent with the highest PLQY, the 3,8-CzPhen-based device is most efficient with an EQE_{\max} of 3.01%, a maximum brightness (L_{\max}) of 145.6 cd m^{-2} and a current efficiency (CE_{\max}) of 4.34 cd A^{-1} (Table S4, ESI† and Fig. 10). Devices prepared using 4,7-CzPhen and 2,9-CzPhen showed $EQE_{\max} = 0.32$ and 0.60%, respectively.

The EL spectra of the devices prepared using 3,8-CzPhen do not show dual emission which is likely due to a poor population and/or efficient non-radiative decay of the charge-transfer $^1[\pi_{Cz}\pi^*_{phen}]$ state in the host employed, although a shoulder is seen at *ca.* 570 nm, whose intensity increases with increasing drive voltage (Fig. S11, ESI†).

The devices employing 3,8-CzPhen were then prepared using much higher emitter loadings of 20 wt% and 40 wt%, both emitted white light with the EL spectra featuring an additional, red-shifted band centred at *ca.* 570 nm (Fig. S11, ESI†). This new band, assigned as the $^1[\pi_{Cz}\pi^*_{phen}]$ charge-transfer state, intensifies relative to the blue band with increasing drive voltage. We believe this effect arises as in the matrix of the emitting layer at low loadings of 3,8-CzPhen, the charge-transfer

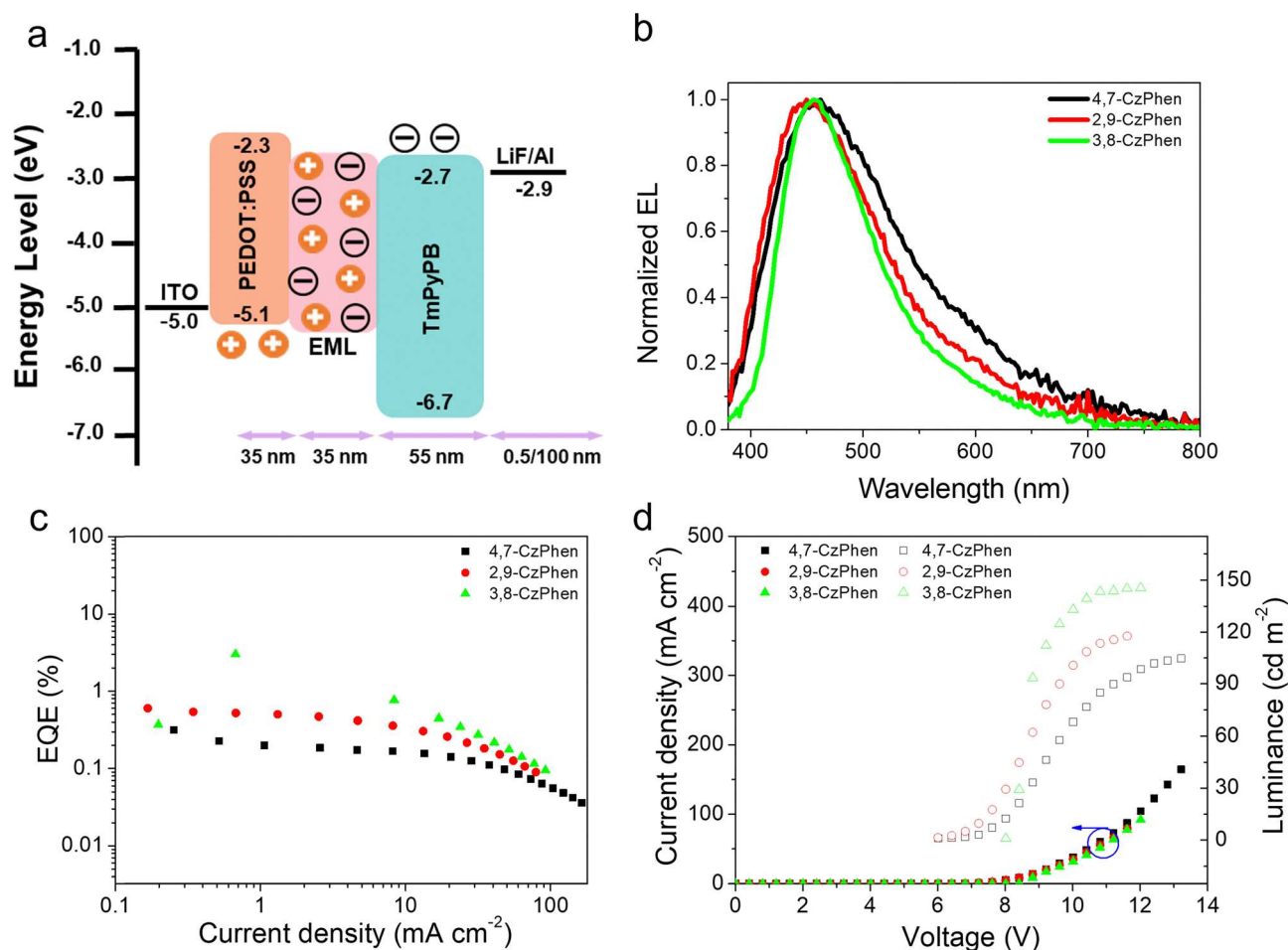


Fig. 10 Device performance at 1 wt% loading of emitter. (a) Schematic of the energy levels and device structure; (b) EL spectra; (c) the EQE-current density curves; (d) the current density-voltage-luminance curves.



$^1[\pi_{Cz}\pi^*_{phen}]$ state is not populated efficiently by internal conversion from the higher-lying state localized on alkoxy-aryl carbazoles ($^1[\pi_{Cz}\pi^*_{Cz}]$) whose sole emission is observed in the devices with lower doping levels. However, with much higher loadings, the phenanthroline core may start acting as the electron transport material, leading to direct population of the $^1[\pi_{Cz}\pi^*_{phen}]$ charge-transfer state without the need for internal conversion from the higher-lying $^1[\pi_{Cz}\pi^*_{Cz}]$ state. This is supported by the fact that photoluminescence of **3,8-CzPhen** when doped at 20% into the host does not show signs of emission from the $^1[\pi_{Cz}\pi^*_{phen}]$ state (Fig. S13, ESI†). The sensitivity of the longer-wavelength band to the drive voltage of the device may then indicate the more efficient charge separation (electron transfer) needed to overcome the exciton binding energy, which is in turn necessary to populate the charge-transfer states from the local state of the alkoxyaryl carbazole substituents.

Conclusion

Despite the non-planar nature of the molecules, which arises from sterically induced twisting of the carbazole unit out of the plane of the central phenanthroline unit, all of the new compounds are liquid crystalline, each showing a single Col_h mesophase. However, while the mesophase stabilities of **4,7-CzPhen** and **3,8-CzPhen** are rather similar, the Col_h phase of **2,9-CzPhen** is very much more stable, clearing some 130 °C higher (≈ 308 °C) than that of **3,8-CzPhen**. From the calculated structures, it is possible to predict how the compounds are likely to self organise in the columnar structure and such an analysis is able to rationalise both the similarities between the clearing points of **4,7-CzPhen** and **3,8-CzPhen** and also the difference with **2,9-CzPhen**.

All three compounds – **2,9-CzPhen**, **3,8-CzPhen** and **4,7-CzPhen** – show fluorescence in toluene solution centred at *ca.* 450 nm arising from a singlet state localised on the alkoxyaryl carbazole moieties ($^1[\pi_{Cz}\pi^*_{Cz}]$). In the doped PS film this emission shift to *ca.* 420 nm in case of **3,8-CzPhen** and **4,7-CzPhen** and is also accompanied by additional fluorescence bands centred at 570 nm and 515 nm, respectively, assigned to the charge-transfer singlet state ($^1[\pi_{Cz}\pi^*_{phen}]$). Such unusual dual fluorescent behaviour is rationalised by the electronic decoupling of the $^1[\pi_{Cz}\pi^*_{Cz}]$ and $^1[\pi_{Cz}\pi^*_{phen}]$ states, brought primarily by spatially isolated π^*_{Cz} and π^*_{phen} orbitals, where the wavefunction of the excited electron is well separated between the two states. This drastically limits the rate and efficiency of internal conversion between the two states and thus allows rare anti-Kasha emission from a higher-energy excited singlet state. Only for **4,7-CzPhen** in a PS matrix could a very small share of the emission intensity (*ca.* 5% of emission from the $^1[\pi_{Cz}\pi^*_{phen}]$ state) be identified as arising with the participation of a triplet state, being assigned to TADF. This is associated with (i) the presence of two CT triplet states, T₁ and T₂, in very close energetic proximity to the S₁ state ($^1[\pi_{Cz}\pi^*_{phen}]$) so that ISC and RISC processes can be relatively fast, and (ii) relatively high energy of the lowest singlet and triplet CT states limiting the efficiency of their non-radiative decay rate to the ground state, manifested by much longer decay time of

$^1[\pi_{Cz}\pi^*_{phen}]$ prompt fluorescence (21 ns) compared to that of other two materials. Both photophysical parameters are understood to arise from the relatively twisted geometry of the alkoxyaryl carbazole substituents with respect to the phenanthroline, diminishing $\Delta E(S_1-T_1)$ and giving higher energies of S₁ and T₁ as well as a hindered geometry, so restricting reorganisation in the CT excited state. It is concluded, therefore, that molecular design strategies involving substitutions that would position the alkoxyaryl carbazoles at a larger angle with respect to the phenanthroline core and hinder geometry changes in the CT character excited state would be beneficial for the improvement of the TADF properties of these liquid-crystalline, white-luminescent materials. Additionally, complexation of the phenanthroline with transition metals could help enhance the spin-orbit coupling effect and thus afford more efficient ISC and RISC processes involved in TADF.

The use of **3,8-CzPhen** as the emitter in an OLED device gave an EQE of *ca.* 3% which, while low by many published standards, actually represents a very efficient device in this instance. Thus, exciton multiplicity is defined purely statistically by the spins of injected electrons and holes, over which we have no control, resulting in 25% singlets and 75% triplets. The materials are then able to use only these (25%) singlet excitons and so the EQE is then understood in the context of a PLQY of 40% and a maximum outcoupling efficiency on the order of 30%. At high emitter loadings of 20 or 40 wt%, dual-emission, white-emitting devices are realised arising from the electron-transporting properties of the phenanthroline unit of the emitter materials, although the EQE suffers from quenching effects and the inability of the emitter to utilise triplet excitons. However, and as noted previously,^{47,48} the non-LC nature of the host materials and the loadings of LC emitter used means that LC properties are not expressed in the device, which is a facet that requires further attention.

However, while it is demonstrated here that dual emission affording white luminescence is readily achievable in a PS film, the choice of a suitable OLED host to transfer it to the device turned out not to be a trivial task. Comparison of photo- and electro-luminescence spectra of **3,8-CzPhen** in PVK/OXD-7 shows that the main problem is the slow and inefficient population of the CT state by internal conversion from the higher-lying states local to the carbazoles. As such, CT emission intensity is low/vanishing at low loadings and appears significant only at high concentrations where the phenanthroline unit of the emitter start acting as an electron-transport material.

Nonetheless, the observation of both matrix- and device-based dual emission giving rise to white light is a rare^{18,49,50} and very significant finding and it is concluded that host blends that allow direct population of local states and CT states need to be tried and developed, an approach requiring emitter materials with better-developed TADF properties.

Author contributions

AFS, MZS and DWB conceived the project. AFS undertook all of the synthetic work and characterised the liquid crystal



behaviour with DWB. Photophysical characterisation and the quantum chemical calculations were undertaken by MZS and JRF. XC made the OLED devices. Liquid crystal behaviour was analysed and interpreted by DWB, photophysical data were analysed and interpreted by MZS and YW analysed and interpreted the device data. AFS, MZS and YW wrote the first drafts of the manuscript, which they then helped refine with DWB. DWB and RC secured funding for the work.

Data availability

The data supporting this article: (i) have been included as part of the ESI† and (ii) are available at <https://doi.org/10.15124/e2cc8792-1bc6-43f2-914f-47b40a609b08>.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

The authors would like to thank the University of York (AFS), the Universität Regensburg (MZS), the NSF (GRFP Grant DGE-1842487 to JRF) and the German Research Foundation (DFG) (Project No. 389797483) for financial support, Dr Stephen Cowling (York) for assistance with the polarised optical microscopy and Xiaoyi Lai for assistance with some photoluminescence measurements.

References

- 1 D. Chen, F. Tenopala-Carmona, J. A. Knöller, A. Mischok, D. Hall, S. Madayanad Suresh, T. Matulaitis, Y. Olivier, P. Nacke, F. Gießelmann, S. Laschat, M. C. Gather and E. Zysman-Colman, *Angew. Chem., Int. Ed.*, 2023, **62**, e202218911.
- 2 C. Mayr, M. Taneda, C. Adachi and W. Brütting, *Org. Electron.*, 2014, **15**, 3031–3037.
- 3 M. Kasha, *Discuss. Faraday Soc.*, 1950, **9**, 14–19.
- 4 D. Klemp and B. Nickel, *Chem. Phys. Lett.*, 1986, **130**, 493–497.
- 5 G. D. Gillispie and E. C. Lim, *J. Chem. Phys.*, 1978, **68**, 4578–4586.
- 6 G. Eber, F. Grüneis, S. Schneider and F. Dörr, *Chem. Phys. Lett.*, 1974, **29**, 397–404.
- 7 D. Huppert, J. Jortner and P. M. Rentzepis, *Chem. Phys. Lett.*, 1972, **13**, 225–228.
- 8 M. Beer and H. C. Longuet-Higgins, *J. Chem. Phys.*, 1955, **23**, 1390–1391.
- 9 T. Yoshihara, S. Murayama and S. Tobita, *Sensors*, 2015, **15**, 13503–13521.
- 10 C. Si, T. Wang, Y. Xu, D. Lin, D. Sun and E. Zysman-Colman, *Nat. Commun.*, 2024, **15**, 7439.
- 11 M. Z. Shafikov, A. F. Suleymanova, D. N. Kozhevnikov and B. König, *Inorg. Chem.*, 2017, **56**, 4885–4897.
- 12 M. Shafikov, A. Suleymanova, R. J. Kutta, A. Gorski, A. Kowalczyk, M. Gapińska, K. Kowalski and R. Czerwieniec, *J. Mater. Chem. C*, 2022, **10**, 5636–5647.
- 13 M. Shafikov, A. Suleymanova, R. J. Kutta, F. Brandl, A. Gorski and R. Czerwieniec, *J. Mater. Chem. C*, 2021, **9**, 5808–5818.
- 14 M. Z. Shafikov, D. N. Kozhevnikov, M. Bodensteiner, F. Brandl and R. Czerwieniec, *Inorg. Chem.*, 2016, **55**, 7457–7466.
- 15 D. N. Kozhevnikov, V. N. Kozhevnikov, M. Z. Shafikov, A. M. Prokhorov, D. W. Bruce and J. A. Williams, *Inorg. Chem.*, 2011, **50**, 3804–3815.
- 16 T. Hofbeck, Y. C. Lam, M. Kalbáč, S. Zális, A. Vlček and H. Yersin, *Inorg. Chem.*, 2016, **55**, 2441–2449.
- 17 Y. Liu, H. Guo and J. Zhao, *Chem. Commun.*, 2011, **47**, 11471–11473.
- 18 X. Li, G. Baryshnikov, C. Deng, X. Bao, B. Wu, Y. Zhou, H. Ågren and L. Zhu, *Nat. Commun.*, 2019, **10**, 731.
- 19 A. Suleymanova, M. Shafikov, X. Chen, Y. Wang, R. Czerwieniec and D. W. Bruce, *Phys. Chem. Chem. Phys.*, 2022, **24**, 22115–22121.
- 20 A. Suleymanova, M. Shafikov, A. C. Whitwood, R. Czerwieniec and D. W. Bruce, *J. Mater. Chem. C*, 2021, **9**, 6528–6535.
- 21 S. Naka, H. Okada, H. Onnagawa and T. Tsutsui, *Appl. Phys. Lett.*, 2000, **76**, 197–199.
- 22 H. Yersin, R. Czerwieniec, M. Z. Shafikov and A. F. Suleymanova, *ChemPhysChem*, 2017, **18**, 3508–3535.
- 23 M. Z. Shafikov, A. F. Suleymanova, R. Czerwieniec and H. Yersin, *Inorg. Chem.*, 2017, **56**, 13274–13285.
- 24 M. Z. Shafikov, A. F. Suleymanova, R. Czerwieniec and H. Yersin, *Chem. Mater.*, 2017, **29**, 1708–1715.
- 25 M. Z. Shafikov, A. F. Suleymanova, A. Schinabeck and H. Yersin, *J. Phys. Chem. Lett.*, 2018, **9**, 702–709.
- 26 C. E. Housecroft and E. C. Constable, *J. Mater. Chem. C*, 2022, **10**, 4456–4482.
- 27 K. Wu, T. Zhang, L. Zhan, C. Zhong, S. Gong, Z.-H. Lu and C. Yang, *Adv. Opt. Mater.*, 2016, **4**, 1558–1566.
- 28 H. Norouzi-Arasi, W. Pisula, A. Mavrinskiy, X. Feng and K. Müllen, *Chem. – Asian J.*, 2011, **6**, 367–371.
- 29 S. J. P. Bousquet and D. W. Bruce, *J. Mater. Chem.*, 2001, **11**, 1769–1771.
- 30 T. Cardinaels, J. Ramaekers, P. Nockemann, K. Driesen, K. Van Hecke, L. Van Meervelt, G. Wang, S. De Feyter, E. F. Iglesias, D. Guillon, B. Donnio, K. Binnemans and D. W. Bruce, *Soft Matter*, 2008, **4**, 2172–2185.
- 31 H. Suzuki, T. Kanbara and T. Yamamoto, *Inorg. Chim. Acta*, 2004, **357**, 4335–4340.
- 32 K. M. Knötig, D. Gust, T. Lenzer and K. Oum, *J. Photochem.*, 2024, **4**, 163–178.
- 33 B. N. Bandyopadhyay and A. Harriman, *J. Chem. Soc., Faraday Trans.*, 1977, **73**, 663–674.
- 34 G. Accorsi, A. Listorti, K. Yoosaf and N. Armaroli, *Chem. Soc. Rev.*, 2009, **38**, 1690–1700.
- 35 N. Armaroli, L. De Cola, V. Balzani, J.-P. Sauvage, C. O. Dietrich-Buchecker and J.-M. Kern, *J. Chem. Soc., Faraday Trans.*, 1992, **88**, 553–556.



- 36 M. Montalti, A. Credi, L. Prodi and M. T. Gandolfi, *Handbook of Photochemistry*, CRC Press, 3rd edn, 2006.
- 37 I. Berlman, *Handbook of Fluorescence Spectra of Aromatic Molecules*, Academic, 2nd edn, 1971.
- 38 A. Szabo and N. S. Ostlund, *Modern Quantum Chemistry: Introduction to Advanced Electronic Structure Theory*, Dover Publications, 1989.
- 39 R. Englman and J. Jortner, *Mol. Phys.*, 1970, **18**, 145–164.
- 40 M. J. Frisch, G. W. Trucks, H. B. Schlegel, G. E. Scuseria, M. A. Robb, J. R. Cheeseman, G. Scalmani, V. Barone, B. Mennucci, G. A. Petersson, H. Nakatsuji, M. Caricato, X. Li, H. P. Hratchian, A. F. Izmaylov, J. Bloino, G. Zheng, J. L. Sonnenberg, M. Hada, M. Ehara, K. Toyota, R. Fukuda, J. Hasegawa, M. Ishida, T. Nakajima, Y. Honda, O. Kitao, H. Nakai, T. Vreven, J. A. Montgomery Jr., J. E. Peralta, F. Ogliaro, M. J. Bearpark, J. Heyd, E. N. Brothers, K. N. Kudin, V. N. Staroverov, R. Kobayashi, J. Normand, K. Raghavachari, A. P. Rendell, J. C. Burant, S. S. Iyengar, J. Tomasi, M. Cossi, N. Rega, N. J. Millam, M. Klene, J. E. Knox, J. B. Cross, V. Bakken, C. Adamo, J. Jaramillo, R. Gomperts, R. E. Stratmann, O. Yazyev, A. J. Austin, R. Cammi, C. Pomelli, J. W. Ochterski, R. L. Martin, K. Morokuma, V. G. Zakrzewski, G. A. Voth, P. Salvador, J. J. Dannenberg, S. Dapprich, A. D. Daniels, Ö. Farkas, J. B. Foresman, J. V. Ortiz, J. Cioslowski and D. J. Fox, *Gaussian 09*, Gaussian, Inc., Wallingford, CT, USA, 2009.
- 41 R. Peverati and D. G. Truhlar, *J. Phys. Chem. Lett.*, 2012, **3**, 117–124.
- 42 R. Peverati and D. G. Truhlar, *Phys. Chem. Chem. Phys.*, 2012, **14**, 11363–11370.
- 43 F. Weigend and R. Ahlrichs, *Phys. Chem. Chem. Phys.*, 2005, **7**, 3297–3305.
- 44 M. Cossi, N. Rega, G. Scalmani and V. Barone, *J. Comput. Chem.*, 2003, **24**, 669–681.
- 45 M. A. El-Sayed, *J. Chem. Phys.*, 1963, **38**, 2834–2838.
- 46 R. C. Hilborn, *Am. J. Phys.*, 1982, **50**, 982–986.
- 47 Y. Hong, J. W. Lam and B. Z. Tang, *Chem. Commun.*, 2009, 4332–4353.
- 48 R. R. Parker, D. Liu, X. Yu, A. C. Whitwood, W. Zhu, J. A. G. Williams, Y. Wang, J. M. Lynam and D. W. Bruce, *J. Mater. Chem. C*, 2021, **9**, 1287–1302.
- 49 S. Tang, Y. Tsuchiya, J. Wang, C. Adachi and L. Edman, *Nat. Commun.*, 2025, **16**, 653.
- 50 A. Maggiore, Y. Qu, R. Guillot, P. Pander, M. Vasylieva, P. Data, F. B. Dias, P. Audebert, G. Clavier and F. Miomandre, *J. Phys. Chem. B*, 2022, **126**, 2740–2753.

