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Modular synthesis, host–guest complexation and solvation-controlled relaxation of nanohoops with donor–acceptor structures†

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Carbon nanohoops with donor–acceptor (D–A) structures are attractive electronic materials and biological fluorophores, but their synthesis is usually challenging. Moreover, the preparation of D–A nanohoop fluorophores exhibiting high fluorescence quantum yields beyond 500 nm remains a key challenge. This study presents a modular synthetic approach based on an efficient metal-free cyclocondensation reaction that readily produced nine congeners with D–A or donor–acceptor–donor' (D–A–D') structures, one of which is water-soluble. The tailored molecular design of nanohoops enabled a systematic and detailed study of their host–guest complexation with fullerene, optical properties, and charge transfer (CT) dynamics using X-ray crystallography, fluorescence titration, steady and ultrafast transient absorption spectroscopy, and theoretical calculations. The findings revealed intriguing physical properties associated with D–A motifs, such as tight binding with fullerene, moderate fluorescence quantum yields (37–67%) beyond 540 nm, and unique solvation-controlled CT relaxation of D–A–D' nanohoops, where two CT states (D–A and A–D') can be effectively tuned by solvation, resulting in dramatically changed relaxation pathways in different solvents.

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Introduction

Incorporating electronic donor (D) and acceptor (A) units into the molecular backbone endows organic materials with attractive features, such as the redistribution of frontier orbitals,¹ solvato-fluorochromism,² and captodative effect.³ These features are highly important for material applications in the fields of organic electronics,⁴ photovoltaics⁵ and biology.⁶ In contrast to conjugated polymers with linearly aligned D–A motifs,⁷ the advent of cyclo-paraphenylenes (CPP),⁸ also known as carbon nanohoops, has made it possible to confine the D–A units into a cyclic geometry with quasi-infinite conjugation,⁹ bringing about exciting new features in association with the unique geometric and electronic structures of nanohoops.¹⁰ Fig. 1a shows the two main D–A nanohoops reported so far. The first type has one acceptor in the molecular backbone, while the rest of the CPP subunits are considered donors, because their bent π -scaffold results in higher

energy for the highest occupied molecular orbital (HOMO) compared to the linear counterparts. With this design concept, Itami reported two D–A nanohoops with anthraquinone and tetracyanoanthraquinodimethane installed as acceptors.¹¹ Jasti reported pyridinium embedded nanohoops with various hoop sizes¹² and then a D–A nanohoop containing a benzothiadiazole (BT) moiety.¹³ The second type involves aligning D and A units alternately, often with phenylene acting as a spacer. Examples of this genre include molecular systems prepared by Nuckolls,¹⁴ Wang,¹⁵ Tanaka,¹⁶ Li,¹⁷ and Tan,¹⁸ in which imide, fluorenone, ester, diketopyrrolopyrrole, and BT were employed as acceptors and thiophenylene and phenylene were introduced as donors (Fig. 1b). Applications in the fields of organic electronics¹⁹ and bioimaging²⁰ have started to emerge, even though there are now just a few examples available.

Nanohoops with D–A motifs are particularly intriguing for bioimaging, considering their low cytotoxicity.²⁰ However, applications in biological systems usually require a red emission with a high fluorescence quantum yield (FQY). Unfortunately, most of the abovementioned D–A bearing nanohoops experienced a significant drop in FQY with red-shifted emission, with only a few BT-containing candidates retaining a moderate-to-high FQY with an emission wavelength above 500 nm.¹³ To further meet the application demand, an expanded library of molecules with a well-established structure–property relationship is highly desired. To this end, Jasti *et al.* conducted a systematic theoretical analysis with 18 D–A motifs theoretically examined, thus providing

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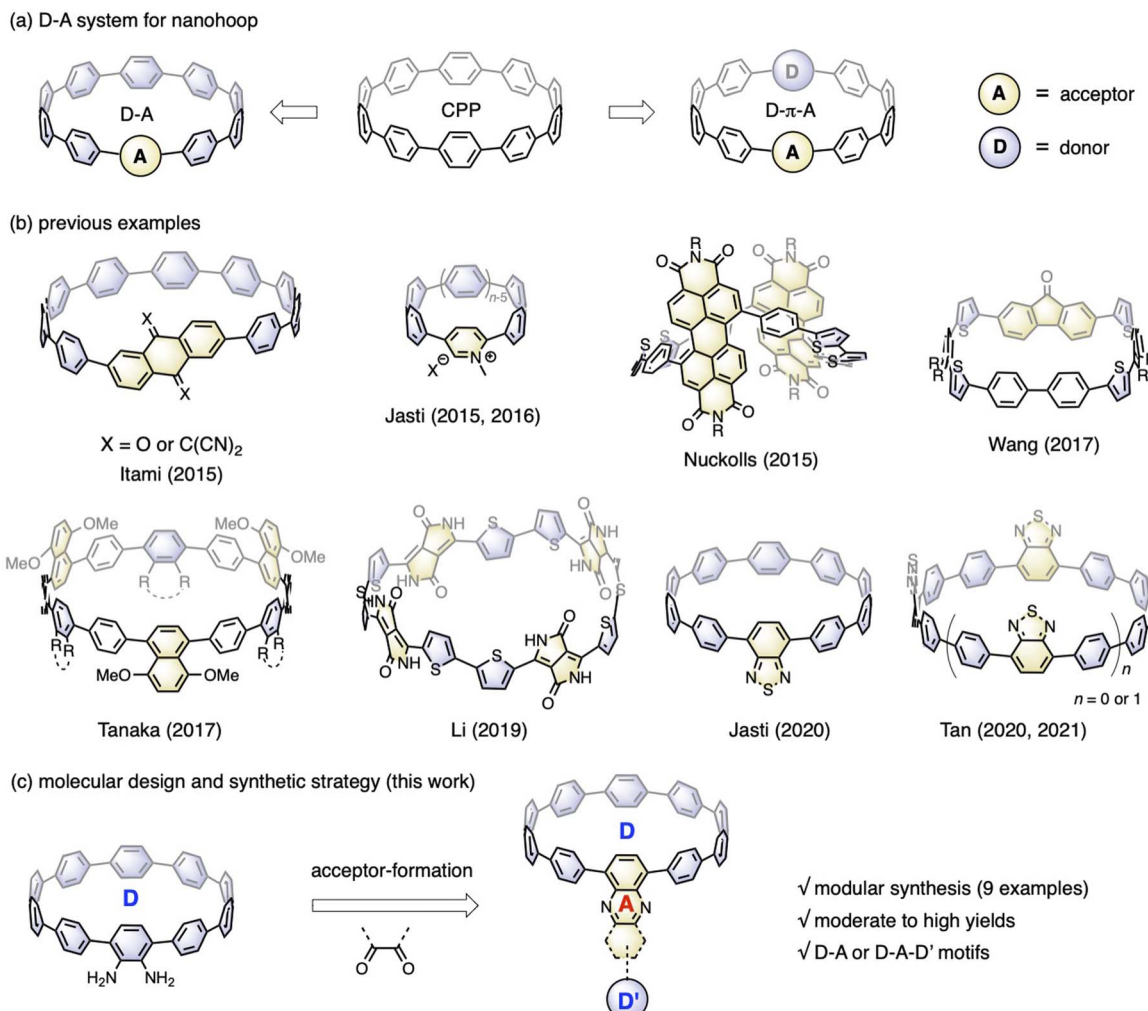


Fig. 1 (a) Two types of nanohoops bearing D-A moieties; (b) previously reported examples of D-A nanohoops; (c) molecular design and synthetic strategy for D-A nanohoops.

a theoretical blueprint for the molecular design.²¹ To make these motifs synthetically accessible, a modular synthetic approach is desirable.

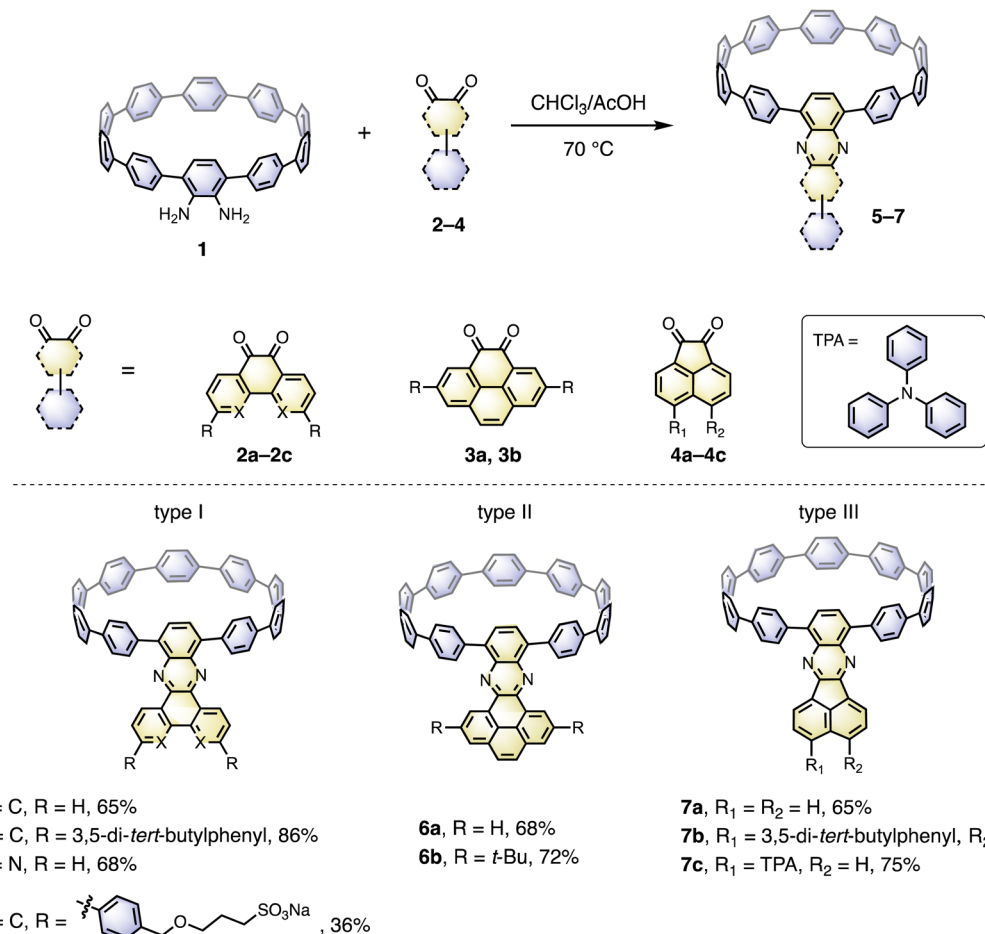
According to the nanohoop-forming strategies discovered by Jasti/Bertozzi,^{8a} Yamago,²² Itami,²³ and Tsuchido/Osakada,²⁴ the donor and acceptor units can be introduced before or during the macrocyclization stage. When the donor and acceptor moieties are changed using this synthetic strategy, synthons must be changed from the beginning of the synthetic route, which slows down the rapid expansion of the material scope. In a previous study, we prepared a diamino-[10]CPP derivative *via* a multi-step synthetic sequence and discovered that it could undergo a cyclocondensation reaction efficiently with tetraketone compounds.²⁵ The cyclocondensation reaction could transform the formerly electron-donating amino-containing phenylene into an electron-withdrawing moiety. Additionally, stronger electron donors (D') can be readily introduced by a transition-metal-catalyzed cross-coupling reaction. Therefore, starting from diamino-[10]CPP as a key precursor in this study, a series of D-A or D-A-D' systems are readily accessible by simply changing the diketone moieties

(Fig. 1c). The obtained molecular structures are similar to those of CPPs embedded with polycyclic aromatic hydrocarbons (PAHs) reported by Du,²⁶ but with an additional D-A feature. This modular synthetic strategy enabled the accumulation of nine congeners with deliberate structural design, and their electronic structures, host-guest chemistry and ultrafast photophysics were systematically investigated.

Results and discussion

Synthesis and electronic structures

The cyclocondensation reaction between diamine and diketone is a robust, metal-free reaction commonly adopted to construct complicated polycyclic aromatic systems.²⁷ We discovered that this reaction could be readily applicable to a [10]CPP diamine derivative (**1**) and various diketones (**2-4**) under standard reaction conditions containing only chloroform and acetic acid as solvents (Scheme 1). The diketones were either commercially available or easily prepared according to the literature procedures.²⁸ The reactions proceeded with moderate to good yields, and the workup and



Scheme 1 Synthesis of nanohoops with a D-A structure.

purification procedures were simple and handy (see ESI† for details). After the condensation, the formerly electron donating phenylene with a diamino-group was converted into an electron-withdrawing unit, thus forming a D-A nanohoop. With different diketones, phenanthrene- (type I, 5a–5d), pyrene- (type II, 6a, 6b) and acenaphthylene-type (type III, 7a–7c) congeners were prepared. Notably, compound 5c with further nitrogen-doping could also be obtained, representing an attractive motif for metal complexation.²⁹ In addition, aliphatic or aromatic substituents can be readily introduced onto the acceptor units, to improve the solubility and to further tune the electronic nature of the nanohoops. Consequently, a water-solubilizing congener 5d was prepared by introducing two water solubilizing sulfonate substituents (Scheme S2†).³⁰ An asymmetric D-A-D' motif 7c, with vertically aligned CPP as a weaker donor and triphenylamine (TPA) as a stronger donor, was also successfully prepared.

The molecular structures of nanohoops were elucidated using nuclear magnetic resonance (NMR) spectra and high-resolution mass spectra. After extensive trials, we were unable to obtain single crystal structures, but were able to obtain the crystal structure of 6a after its encapsulation with C₆₀ (*vide infra*), which supported the formation of the nanohoop structure. To investigate the electronic features resulting from the D-A motifs, density

functional theory (DFT) calculations at the M06-2X/6-311G** level were conducted to provide the frontier molecular orbitals of model compounds of each type, 5a, 6a and 7a, together with the D-A-D' motif 7c (Fig. 2). In comparison to [10]CPP with a fully delocalized HOMO and the lowest unoccupied molecular orbital (LUMO), spatial separation of frontier orbitals was observed for all four molecules, with the HOMO localized on the donor and LUMO distributed on the acceptor. A concomitant reduction of the LUMO and elevation of the HOMO were also found, leading to a narrowed HOMO–LUMO gap. The connection of an additional strong TPA donor in 7c led to elevation of the HOMO and reduction in LUMO levels. The cyclic voltammograms of 5a, 6a, 7a and 7c showed only oxidation waves, and half-wave potentials were in line with the calculated HOMO levels (Fig. S1 and Table S1†). According to the time-dependent DFT (TD-DFT) calculation, the HOMO → LUMO transition, which was Laporte forbidden for [10]CPP,³¹ was partially allowed for the D-A nanohoops in this study because of the broken symmetry in orbitals.

Host–guest complexation with C₆₀

Recently, several C₆₀ hosts have been prepared to realize different functions.³² As a well-studied C₆₀ host, [10]CPP possessed ideal size complementarity for encapsulating C₆₀ with noncovalent van der

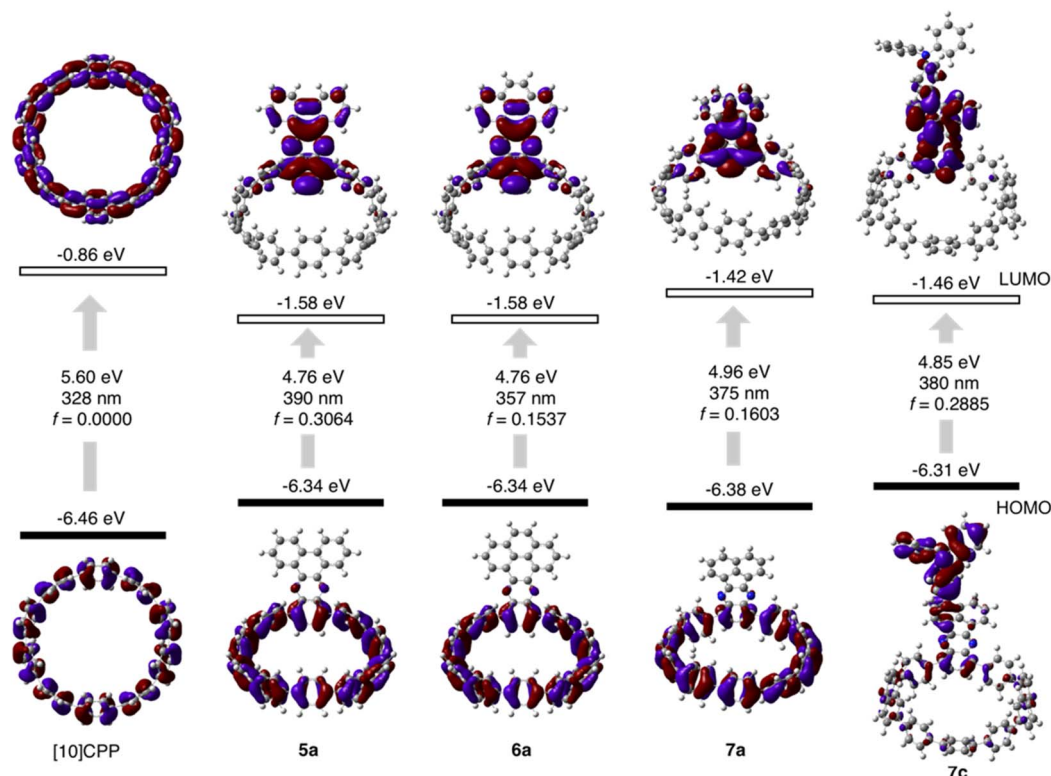


Fig. 2 Calculated frontier orbital distributions and energy levels of [10]CPP and D-A nanohoops in this study.

Waals (vdW) interaction.³³ Considering that C_{60} is a well-known acceptor itself, it is interesting how nano hoop hosts with D-A features could influence the complexation. We first disclose the complexation in the solid state. The single crystal structure of $6a \supset C_{60}$ was obtained by slow diffusion of *n*-hexane into a $CHCl_3$ solution of $6a$ and C_{60} at $-20\text{ }^\circ\text{C}$. Note that the single crystal of the complex was easier to achieve than the free-standing $6a$, suggesting that C_{60} serves as a template for the crystal growth.³⁴ As shown in Fig. 3a, a 1:1 host-guest complex formed with the C_{60} sited in the center of the nano hoop. The intermolecular distances of the host and the guest were in the range of 3.4–3.7 Å, indicating a dominant vdW interaction. The C_{60} molecules were separated by a distance of 3.2 Å, and the continuous vdW contact between the C_{60} molecules suggested the possibility of a conductive pathway for electrons, which is potentially useful for electronic applications. Although the incorporation of the π -extended acceptor had an insignificant influence on the circular shape of the nano hoop (Fig. S2†), the torsional angles of the acceptor and the adjacent phenyl (52° and 54°) were larger than the rest of the CPP units (45° on average, Table S3†). This resulted in a tilting dihedral angle of 120° between the acceptor unit and the CPP mean plane. In the packing structure, each molecule was zipped together by the π - π interaction between neighbouring C_{60} and between the acceptor unit and phenyl (Fig. 3b), to form a one-dimensional stacking. Such interactions sewing the complexes together can be further visualized by d_e mapping from the Hirshfeld analysis,³⁵ which demonstrated a similar pattern of close contact in the interior of the acceptor and exterior of the phenyl ring (Fig. 3c). The solvent molecules of *n*-

hexane and $CHCl_3$ were filled in the curved space between the two complexes (Fig. S4†). The frontier orbitals of the complex were calculated using DFT at the LC-BLYP/6-311G* level³⁶ with basis set superposition error (BSSE) correction.³⁷ The HOMO was mainly localized on the host and the LUMO on the guest, similar to the

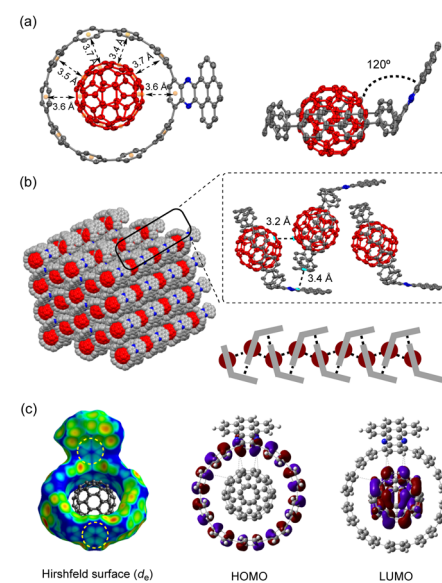


Fig. 3 (a) Top view and side view of the X-ray crystallographic structure of $6a \supset C_{60}$ at 160 K ; (b) the packing structure; (c) Hirshfeld surface analysis and frontier molecular orbitals of $6a \supset C_{60}$.



Table 1 Binding constants and Gibbs energies of complexes measured in *o*-DCB

	K_a (M^{-1})	ΔG at 298 K (kcal mol $^{-1}$)
5a \supset C ₆₀	$(9.10 \pm 0.12) \times 10^5$	-8.10 ± 0.01
6a \supset C ₆₀	$(1.05 \pm 0.21) \times 10^6$	-8.18 ± 0.02
7a \supset C ₆₀	$(7.69 \pm 0.12) \times 10^5$	-7.79 ± 0.01
7c \supset C ₆₀	$(7.61 \pm 0.60) \times 10^5$	-7.78 ± 0.04
[10]CPP \supset C ₆₀ (ref. 33a)	$(6.02 \pm 0.18) \times 10^3$	-5.15 ± 0.02

case of [10]CPP, whereas the LUMO+3 orbital was distributed on the acceptor unit (Fig. S19†). The calculation suggested that the charge transfer between the host and guest dominated in the complex.

The binding behavior was then investigated in the solution phase with four nanohoops, 5a, 6a, 7a and 7c, as model compounds. Upon the addition of C₆₀ into the *o*-DCB solution of hosts, instant fluorescence quenching was observed, in agreement with the charge transfer nature of the complex (Fig. S19†). The binding stoichiometry in solution was estimated to be 1:1, based on the titration experiments at different concentrations that are all well fitted to the 1:1 model (Fig. S9 and Table S4†).³⁸ The ¹H NMR spectrum of 6b \supset C₆₀ measured in CDCl₃ at -30 °C shows a significant change in chemical shifts of the phenylene protons compared to the free-standing 6b, whereas the protons on nitrogen-containing PAH were less affected (Fig. S10†). As presented in Table 1, triplicate fluorescence quenching titration in *o*-dichlorobenzene (*o*-DCB) gave similar binding constants for all four complexes, in the order of 10^5 to 10^6 M $^{-1}$, which was two to three orders of magnitude higher than that of [10]CPP \supset C₆₀ measured in the same solvent (Fig. S5–S8†).^{33a} The enhancement in binding could be due to (1) an enlargement of the π – π interaction area^{26a} and (2) an elevation in the HOMO level of the D–A nanohoop which facilitated the strong electrostatic interactions between the host and guest.

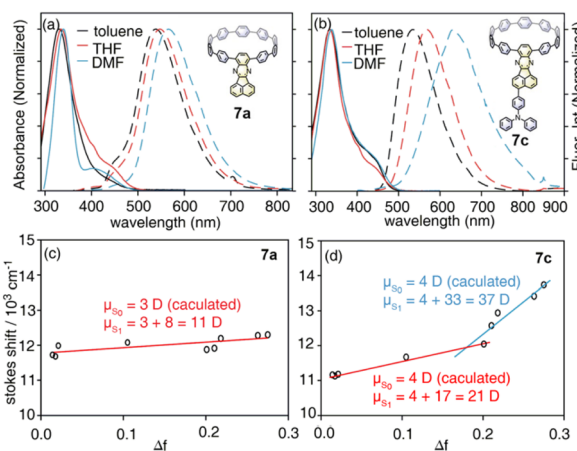


Fig. 4 Absorption (solid lines) and fluorescence spectra (dashed lines) of (a) 7a and (b) 7c in selected solvents, and the corresponding Lippert–Mataga fittings of (c) 7a and (d) 7c.

Optical and excited state properties

The absorption and emission spectra of 9 synthesized D–A nanohoops were measured in CH₂Cl₂ and water solutions (Fig. 4, S14 and S16†) and the data are summarized in Table 2. All compounds showed maximum absorptions around 330 nm, which is typical of the CPP structure. The shoulder peaks in the region of 400–500 nm are from the partly allowed HOMO \rightarrow LUMO transition (Fig. 2). Compared to the emission of the parent [10]CPP (466 nm), the emissions were substantially red-shifted to a region of 540–610 nm, falling in the optical window desirable for biological applications. Gratifyingly, a moderate FQY was retained at a level of 37–67%, in contrast to other nitrogen containing D–A CPPs.¹² This could be due to the absence of a nonradiative relaxation channel in our system. Note that 5d exhibits a FQY of 16% in an aqueous solution, which is comparable to Jasti's water soluble nanohoop but with a 69 nm redshift in emission (510 nm vs. 579 nm).^{30a} The plot of wavenumber against the solvent parameter $E_T(30)$ ³⁹ revealed positive solvatofluorochromism due to intramolecular charge transfer (Fig. S15†).

With model systems 7a and 7c, the photophysics of D–A and D–A–D' nanohoops were further investigated using steady and ultrafast spectroscopic measurements. As shown in Fig. 4, the absorption spectra of 7a and 7c were dominated by a pronounced peak at approximately 330 nm, corresponding to a local excited state of CPP rings (LE_{CPP}) as revealed by natural transition orbital analysis (NTO, see Fig. S20†),⁴⁰ *i.e.* S₀ \rightarrow S₃ (7a, 3.590 eV) and S₀ \rightarrow S₄ states (7c, 3.585 eV) with pronounced oscillator strength. As forbidden transitions, the local excited states of the acceptor (LE_A) were also revealed through calculations for 7a (S₂) and 7c (S₃) with low oscillator strength. Meanwhile, weak absorption in the 400–500 nm regime was observed for 7a and 7c as a shoulder of the LE_{CPP} peak. For 7a (D–A), the NTO analysis (Fig. S20a†) indicated pronounced charge transfer (CT_{CPP \rightarrow A}) characteristics of the lowest-lying S₁ state (\sim 3.3 eV) below the LE_A (S₂) state. However, the case of 7c (D–A–D') was more complicated, *i.e.* both S₁ and S₂ states exhibited a CT character but corresponded to different donors. Under weak solvation (calculated using PCM = toluene), the S₀ \rightarrow S₁ transition of 7c corresponds to CT_{CPP \rightarrow A} whereas TPA acts

Table 2 Optical properties of [10]CPP and D–A nanohoops in this study

	λ_{abs} /nm	λ_{em} /nm	FQY (%)	Solvent
5a	334, 448, 453	590	67	CH ₂ Cl ₂
5b	333, 418	590	53	CH ₂ Cl ₂
5c	331	602	43	CH ₂ Cl ₂
5d	337, 394, 417	593	56	CH ₂ Cl ₂
5d	338, 392, 429	579	16	H ₂ O
6a	333, 423, 447	593	60	CH ₂ Cl ₂
6b	322, 428, 449	593	52	CH ₂ Cl ₂
7a	329	555	40	CH ₂ Cl ₂
7b	333	542	37	CH ₂ Cl ₂
7c	332	579	65	CH ₂ Cl ₂
[10]CPP ^{8c}	338	466	65	CH ₂ Cl ₂



Table 3 The TD-DFT (M06-2X/6-311G**, PCM = toluene) calculated excitation energy, oscillator strength and tentative assignment for low-lying excited states of **7a** and **7c**

7a (CPP-A)				7c (CPP-A-TPA)		
States	Excitation energy (eV)	Oscillator strength	Assignment	Excitation energy (eV)	Oscillator strength	Assignment
S ₁	3.3005	0.1603	CT _{CPP→A}	3.2590	0.2885	CT _{CPP→A}
S ₂	3.4710	0.0080	LE _A	3.3421	1.0278	CT _{A←TPA}
S ₃	3.5901	1.0510	LE _{CPP}	3.4929	0.0021	LE _A
S ₄	3.6597	0.2025	CT _{CPP→A}	3.5852	0.7621	LE _{CPP}

as a donor (CT_{A←TPA}) for S₀ → S₂ with an even higher oscillator strength (Fig. S20b†). Two plausible CT states (CT_{CPP→A} and CT_{A←TPA}) might be involved in the relaxation of **7c** due to its unique asymmetric D–A–D' structure with weak and intense donors, *i.e.* CPP (D) and TPA (D'), respectively.

To verify the assignment, the quantitative CT/LE contribution of each excited state was estimated using hole–electron analysis of **7a** and **7c** (see Table S11†)^{41,42} Regarding D–A nanohoop **7a**, the LE_{CPP} state (S₃) was confirmed with approximately 85% hole–electron overlapping and LE% up to approximately 94%. However, a weak (~24%) CT character was observed for the S₁ state (CT_{CPP→A}). Note that the calculated LE/CT contribution is highly dependent on the pre-setting of D/A fragments.^{41,43} The CPP was set as a donor for **7a**, but more options must be considered for asymmetric D–A–D' (**7c**). Without loss of generality, we calculated CT% of each excited state of **7c** with three pre-settings (Table S11†): (1) TPA as a donor, the S₂ state exhibits an ~31% CT character; (2) CPP as a donor, the S₁ state exhibits approximately a 19% CT character; (3) TPA and CPP as donors, both S₂ and S₁ exhibit a CT character of approximately 37% and 24%, respectively. The results confirmed that both S₁ and S₂ of **7c** were CT states but with different donors (*i.e.* CT_{CPP→A} and CT_{A←TPA}, respectively). The higher CT% and oscillator strength of CT_{A←TPA} indicate that TPA is stronger than CPP for donating electrons. According to the results, the existence of TPA might lead to dramatically changed photophysics of **7c**. The LE_{CPP} (S₄) and LE_A (S₃) states were also confirmed, and the assignment of S₁ to S₄ states of **7a** and **7c** is summarized in Table 3.

Due to the dipolar character, the energy level of CT states can be effectively tuned by the solvation effect,⁴⁴ *i.e.* solvatochromism. As CT_{A←TPA} is considered to be more dipolar than CT_{CPP→A}, more pronounced solvatochromism can be expected for CT_{A←TPA}, which was further verified (Fig. 4, bottom panel). As shown in Fig. 4, **7a** exhibited a linear dependence of the Stokes shift on solvent polarity (Δf), whereas the Lippert–Mataga fitting⁴⁵ resulted in approximately 10.95 debye dipole moment of the S₁ state (μ_{S1} , for details see ESI, Section 5.2†). In contrast, segmented dependence of the Stokes shift was observed with Δf for D–A–D' **7c**. In the weak solvation regime (low polarity solvents, $\Delta f < 0.2$), μ_{S1} was estimated to be approximately 20.98 debye. However, a different S₁ state of **7c** with almost doubled μ_{S1} (~37.10 debye) was observed in a strong solvation regime ($\Delta f > 0.2$), which was further confirmed using fluorescence lifetime measurements. As shown in Fig. S18,† the fluorescence decay (S₁ lifetime) of **7a** was similar in toluene, tetrahydrofuran (THF) and dimethylformamide (DMF),

corresponding to the emission of an identical S₁ state, *i.e.* CT_{CPP→A}.⁴⁶ However, the fluorescence lifetime of **7c** in DMF was significantly shorter than that in toluene and THF, indicating a different S₁ state of **7c** under weak and strong solvation conditions. Compared with the CT_{CPP→A} (S₁) state in low polarity solvents, we believe that the CT_{A←TPA} state of **7c** was energetically lowered down due to strong solvation in high polarity solvents, *e.g.* DMF, while solvatochromism of the CT_{CPP→A} state was less pronounced due to a lower dipole moment. Consequently, CT_{A←TPA} becomes the lowest-lying S₁ state of **7c** in DMF, whereas CT_{CPP→A} serves as a S₁ state in toluene and THF, which may lead to dramatically changed ultrafast relaxation dynamics of **7c** in solvents of different polarity.

Ultrafast charge transfer dynamics

The S₁ state plays an important role in the photophysics of organic chromophores.⁴⁷ Compared with the D–A nanohoop (**7a**), the coexistence of CT_{CPP→A} and CT_{A←TPA} states may provide the D–A–D' nanohoop (**7c**) with unique electronic relaxation where the energy levels of two CT states might be effectively tuned by solvation. This was further characterized by using femtosecond transient absorption (fs-TA) spectroscopy. Upon excitation at 330 nm, fs-TA spectra of **7a** and **7c** in toluene, THF and DMF solutions were measured in the 460–850 nm probe range with approximately 200 fs temporal resolution (Fig. 5).

As shown in Fig. 5(a–c), D–A nanohoops (**7a**) exhibited a similar fs-TA signal in toluene, THF and DMF. According to the TD-DFT calculation (Table 3), the LE_{CPP} state of **7a** is initially populated by 330 nm excitation, corresponding to a broad feature (550–850 nm, maximum at ~690 nm) which was consistent with the reported excited state absorption (ESA) band of [10]CPP.⁴⁸ The decay of the LE_{CPP} signal was accompanied by the rise of a new band at ~490 nm, corresponding to an ultrafast charge transfer process (LE_{CPP} → CT_{CPP→A}).⁴⁹

Meanwhile, spectral depletion at approximately 525 nm was observed in the initial TA spectra and red-shifts were observed in subsequent delay steps up to several picoseconds, which was attributed to dynamic Stokes shifting of the stimulated emission (SE) band, *i.e.* the solvation of the CT_{CPP→A} state. Note that the solvation of CT states is typically ultrafast in simple organic dipolar solvents,⁵⁰ *i.e.* down to the sub-picosecond time scale. Therefore, the SE dynamic shifting might not be fully disentangled from the LE_{CPP} → CT_{CPP→A} process due to the limitation of temporal



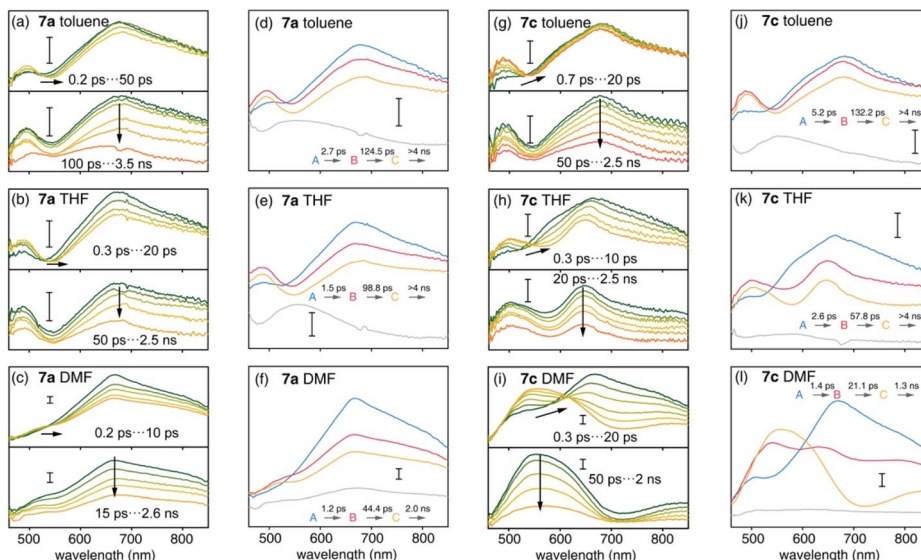


Fig. 5 Transient absorption spectra (a–c and g–i) and species-associated spectra (SAS) (d–f and j–l) of **7a** and **7c** in toluene (upper panel), THF (medium panel) and DMF (lower panel). Time constants were obtained from target analysis with a four step sequential model. The scale bar of signal intensity represents 1 mOD.

Table 4 Target analysis estimated time constants and their assignments

	$A \rightarrow B, \tau_1$ (ps)	Assignment	$B \rightarrow C, \tau_2$ (ps)	Assignment	$C \rightarrow, \tau_3$ (ns)	Assignment
7a	2.7	LE _{CPP} → CT _{CPP→A} and solvation of CT _{CPP→A}	124	Fast structural relaxation of CT _{CPP→A}	>4	Slow relaxation of CT _{CPP→A}
	1.5		98.8	Fast structural relaxation of CT _{CPP→A}	>4	Slow relaxation of CT _{CPP→A}
	1.2		44.4		2.0	
7c	5.2	LE _{CPP} → CT _{CPP→A} and solvation of CT _{CPP→A}	132	Fast structural relaxation of CT _{CPP→A}	>4	Slow relaxation of CT _{CPP→A}
	2.6		57.8	CT _{CPP→A} → CT _{A←TPA}	>4	Relaxation of CT _{A←TPA}
	1.4		21.1		1.3	

resolution. The subsequent relaxation of solvated CT_{CPP→A} was further observed in several nanoseconds.

The D–A–D' nanohoop (**7c**) in toluene exhibited nearly identical fs-TA spectra of **7a** (Fig. 5g) as expected, because CT_{CPP→A} still serves as the lowest-lying state for **7c** under weak solvation conditions. In THF, the ESA band (CT_{CPP→A}) was observed to be slightly extended to a longer wavelength (Fig. 5h). Furthermore, **7c** showed different TA responses under strong solvation conditions in DMF (Fig. 5i). The dynamic shifting of the SE band (CT_{CPP→A}) eventually leads to a pronounced ESA band in the 500–650 nm regime, which is similar to the reported ESA band of the solvated CT state with a TPA donor of a symmetric D–A–D chromophore (Fig. S21†).⁵¹ Furthermore, as discussed above, the energy level of CT_{CPP→A} and CT_{A←TPA} of **7c** can be effectively tuned by solvation. In DMF, the energy level of solvated CT_{A←TPA} becomes even lower than CT_{CPP→A} due to stronger solvation. As a result, relaxation of CT_{CPP→A} → CT_{A←TPA} was observed. Therefore, we assigned the formed broad ESA band of **7c** in DMF to the solvated CT_{A←TPA}, which is unobservable under weak solvation conditions.

We further conducted a quantitative target analysis on all TA data, which can be well reproduced using a sequential model containing four independent species (A → B → C → D), and

the estimated time constants are summarized in Table 4. The subtracted species-associated spectra (SAS) are illustrated in Fig. 5, while the concentration evolution of each species and time trace at the selected probe wavelength can be seen in Fig. S22 and S23† respectively.

The initial process (A → B) of **7a** exhibited ultrafast time constants ($\tau_1 < 3$ ps) and became shorter with increasing solvent polarity, which was consistent with the reported formation and solvation of the CT state.^{51,52} The increased CT_{CPP→A} band at 490 nm and red-shifted SE depletion on SAS of species B further confirmed our assignment of process A → B, *i.e.* LE_{CPP} → CT_{CPP→A} and solvation of CT_{CPP→A}. The subsequent process (B → C, τ_2) for tens of picoseconds with an unchanged TA shape was assigned to the fast relaxation of CT_{CPP→A}, in which structural relaxation might play a key role as widely reported for organic fluorescent chromophores.^{51,52} The slow relaxation of CT_{CPP→A} (C →, τ_3) leaves a structureless TA signal within nanoseconds, which might include several slow relaxation channels such as intersystem crossing, fluorescent decay or photochemical reactions. The relaxation mechanism of **7a** is summarized in Fig. 6a.

The target analysis on TA data of D–A–D' nanohoops (**7c**) in toluene led to comparable results of **7a**. As shown in Fig. 6c, the

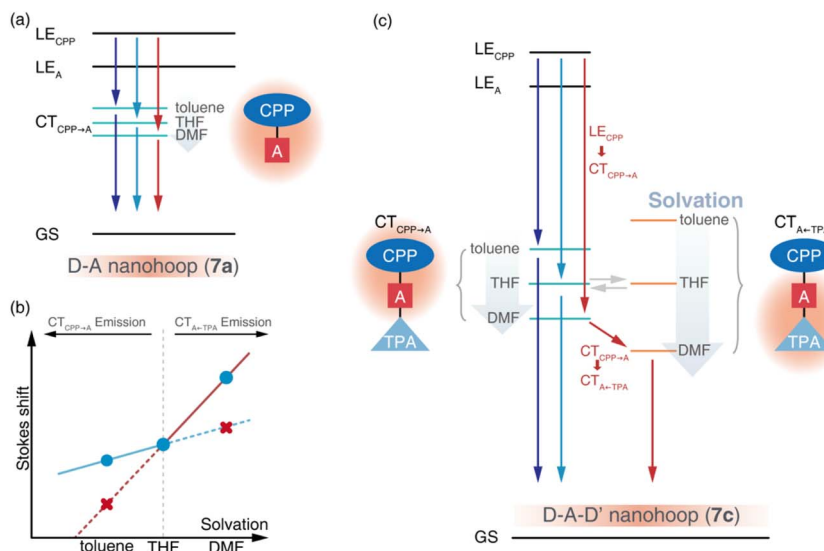


Fig. 6 Energy level diagrams illustrating the excited-state dynamics of (a) 7a and (b and c) 7c in solvents of different polarity. The arrows with a solid line represent transition between states, while the arrows as filled bands represent energetic level lowering of CT states due to solvation.

solvated $\text{CT}_{\text{CPP} \rightarrow \text{A}}$ is energetically lower than $\text{CT}_{\text{A} \leftarrow \text{TPA}}$ under weak solvation, *i.e.* in toluene. Consequently, solvated $\text{CT}_{\text{CPP} \rightarrow \text{A}}$ formed within $\tau_1 = 5.2$ ps and subsequently relaxed bi-exponentially ($\tau_2 = 132$ ps and $\tau_3 > 4$ ns) without involving $\text{CT}_{\text{A} \leftarrow \text{TPA}}$. In THF, the energy levels of solvated $\text{CT}_{\text{CPP} \rightarrow \text{A}}$ and $\text{CT}_{\text{A} \leftarrow \text{TPA}}$ might be nearly degenerate due to stronger solvation than in toluene. Therefore, a photostationary state including $\text{CT}_{\text{CPP} \rightarrow \text{A}}$ and $\text{CT}_{\text{A} \leftarrow \text{TPA}}$ might form within the observable $\tau_1 = 2.6$ ps, which was observed as the spectrally extended SAS of the TA response (species B and C, Fig. 5k) and further decayed bi-exponentially ($\tau_2 = 57.8$ ps and $\tau_3 > 4$ ns). Finally, the strong solvation in DMF changed the relative energy levels between $\text{CT}_{\text{CPP} \rightarrow \text{A}}$ and $\text{CT}_{\text{A} \leftarrow \text{TPA}}$. As shown in the Lippert–Mataga model of 7c (Fig. 6b), the energy level of solvated $\text{CT}_{\text{A} \leftarrow \text{TPA}}$ was below $\text{CT}_{\text{CPP} \rightarrow \text{A}}$ in DMF. As a result, the formed ($\tau_1 = 1.4$ ps) solvated $\text{CT}_{\text{CPP} \rightarrow \text{A}}$ subsequently relaxed to $\text{CT}_{\text{A} \leftarrow \text{TPA}}$ with a time constant of $\tau_2 = 21.1$ ps, corresponding to the observed broad ESA band which dominated the SAS of species C. The formed $\text{CT}_{\text{A} \leftarrow \text{TPA}}$ further decayed, with a faster decay rate ($\tau_3 = 1.3$ ns) than $\text{CT}_{\text{CPP} \rightarrow \text{A}}$ ($\tau_3 = 2.0$ ns), indicating their different states of origin.

The relaxation of the solvated CT state has been widely investigated in dipolar solvents.⁵³ Meanwhile, symmetric D–A–D and A–D–A chromophores have attracted attention due to the unique excited-state symmetry breaking relaxation,⁵⁴ *i.e.* the initially populated quadrupolar state decays to a dipolar state under strong solvation conditions. With model systems 7a and 7c, we demonstrated a unique photophysics mechanism of an asymmetric D–A–D' chromophore, in which the relative energy levels of two CT states (D–A and A–D') are highly dependent on solvation. As a result, the relaxation pathway of the D–A–D' chromophore can be effectively controlled using different solvation conditions.

Conclusions

In summary, this study described a versatile “late stage” synthetic approach that uses an effective metal-free cyclocondensation

reaction, which readily produces nanohoops with D–A or D–A–D' structures with tailored molecular design in moderate to high yields. Nine congeners, which belong to three structural genres, were synthesized and systematically investigated in terms of host–guest chemistry, optical properties and ultrafast charge transfer dynamics. Importantly, all compounds retain a moderate FQY (37–67%) with emission wavelengths ranging from 540–610 nm. Particularly, one of the nanohoops is water soluble with a FQY up to 16% at 579 nm in the aqueous solution, which is a feature desirable for biological applications. The solvation-controlled charge transfer relaxation of D–A–D' nanohoops was demonstrated using ultrafast spectroscopic measurements and TD-DFT calculations. The energy level of two CT states (D–A and A–D') can be effectively tuned by solvation of D–A–D' nanohoops, which leads to dramatically changed relaxation pathways in different solvents. We envision that the synthetic strategy presented in this study will rapidly expand the scope of nanohoop fluorophores with D–A structures, paving the way for practical applications particularly in the biological field.

Data availability

The ESI[†] contains a detailed description of the synthetic method, computational method and the supplementary spectroscopic and crystallographic data.

Author contributions

Z. S. and X. M. supervised the project. H. D., K. L., Q. Z., C. G. and Z. X. performed synthetic experiments and the study of host–guest chemistry. Z. G. and Y. W. performed spectral measurements and theoretical calculations. S. S. performed crystallographic analysis. All authors analyzed the data, discussed the results, and contributed to the manuscript writing.



Conflicts of interest

There are no conflicts to declare.

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