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Photoinduced electron transfer in [10]CPP \supset C₆₀ oligomers with stable and well-defined supramolecular structures[†]

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Recent synthesis of a new type of polymer containing conjugated cycloparaphenylenes (CPP) macrocycles interconnected by a linear conjugated backbone opens up great potential of cyclic π -conjugated materials in organic photovoltaics. In this work, I report a theoretical study of the ground and excited state properties of such polymers and investigate an effect of inclusion of fullerene molecules into polymer chains. MD simulations reveal that oligomers (**[10]CPP_Fused \supset C₆₀**)₂₄ and (**[10]CPP_Fused \supset C₆₀**)₃₂ with π -extended CPPs tend to form stable, helix-like structures. I show that photoinduced electron transfer from the CPP-based polymer to C₆₀ fullerene is favorable and occurs on a nanosecond time scale. The hole- and excess-electron transfer rates are found to be significantly higher than the corresponding charge recombination rates.

Introduction

Attention to molecules with radial π -conjugation has grown very rapidly in recent decades. Rare examples of cyclic structures with radial π -conjugation, such as paraphenylacetylenes,^{1,2} picotubes,^{3,4} or cyclocarbons,^{5–7} have been appearing in the literature since the early 1990s, but have been associated with extremely difficult and low-yield advanced organic synthesis. This fact significantly hindered the wide use of such cyclic structures. However, in 2008, Jasti and Bertozzi and co-workers invented a new strategy for the synthesis of cycloparaphenylenes (CPPs) in acceptable yields.⁸ The unusual hoop-shaped structure with radially conjugated π -electron systems of these cycloparaphenylenes leads to significant progress in the development of protocols for the synthesis of various CPPs.^{9,10} Modern approaches have made it possible to obtain CPPs with a diameter from 7 to 28 Å, precisely controlled by the numbers of units in the macrocycle.^{11,12} Nanosized concave cavities of CPPs have been prone to the formation of multiple non-covalent interactions and widely used in supramolecular chemistry.¹³ The first host–guest complex of CPP was reported by Iwamoto *et al.* in 2011.¹⁴ The authors found that [10]CPP with 10 phenylene units has a nearly ideal diameter (13.8 Å) to accommodate C₆₀ fullerene. Considering that tuning the

fullerene properties by functionalization is a non-trivial task, a supramolecular approach to changing the fullerene behavior seems to be a very attractive method for obtaining new fullerene-based systems. Thus, special attention has been paid to CPP's donor–acceptor complexes due to their potential in organic photovoltaic applications.^{13,15,16}

However, the practical usage of such systems in organic electronics is somewhat difficult due to issues typical for small molecules. They suffer from inhomogeneous film formation and create sharp grain boundaries as effective charge carrier traps due to their high crystallinity.^{17–19} In contrast, conjugated polymers form layered structures with a large domain size, as well as the orientation of crystallites with respect to neighboring domains and transport directions.^{19,20} This, in turn, makes the transport properties much less sensitive to the relative lattice orientation and promotes conduction in the device. Despite all the intriguing features of CPPs, they have not previously been utilized in polymeric materials. In 2019, Du and co-workers²¹ reported the first example of a π -conjugated polymer composed of functionalized [8]CPP. The synthetic strategy used is based on the generation of a cyclic molecule from an appropriate curved and bifunctionalized fragment using Pd-catalyzed Suzuki coupling. The subsequent polymerization was carried out using a Ni-mediated Yamamoto reaction. The resulting polymer had a low polydispersity index of 1.44 and consisted of approximately 22 repeating units. Measurements have shown good carrier mobility when using this polymer as a hole- and electron-transport layer.²¹

Peters *et al.*²² also described the synthesis and electronic properties of π -conjugated polymers that combine the linear

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intramolecular conjugation pathways (phenyl/thienylacetylene) with the emerging properties of radial [6]/[8] CPP segments. Gel permeation chromatography data for most of the synthesized polymers indicated the presence of 7 to 27 repeating units. Photophysical measurements demonstrated that the combination of linear and radial π -conjugations provides more additive properties, but creates new electronic processes. The effect of the communication of CPP units in polymers was also considered. Authors demonstrate that the electronic properties of polymeric [8]CPPs with disjointed pi-conjugated substituents arise from multiple operative radial/linear conjugation pathways, as the disjoint pattern results in both *ortho* and *meta* connections to the CPP ring.²³

More recently, Wang *et al.*²⁴ have developed a novel all-carbon polymeric structure containing conjugated macrocycles linked together by a linear poly(*para*-phenylene) backbone acting as a host for the C₆₀ fullerene. The procedure for synthesizing a new [10]CPP-based polymer was the same as that for the original [8]CPP-based polymer. Fullerene was incorporated into polymeric [10]CPP at the last stage. The polydispersity index of the [10]CPP polymer was found to be 2.66, indicating a broad molecular weight distribution. MALDI-TOF mass spectrometry detected fractions containing up to 17 repeating units. Transient absorption measurements showed that, upon photoirradiation, an electron is transferred from the photoexcited polymeric unit to the C₆₀ fullerene. In addition, the measurement of electron- and hole-mobility clearly showed that the incorporation of C₆₀ into a [10]CPP-based polymer leads to an improvement in the conductive properties.

Herein, I discuss the effect of a linker between CPPs and the inclusion of C₆₀ fullerene into the polymer chain of CPPs on the spatial structure of $([10]\text{CPP})_n$ and quantitatively characterize photoinduced electron transfer (PET), hole transfer (HT), and excessive electron transfer (EET) processes occurring in the all-carbon conjugated $([10]\text{CPP_Fused}\supset\text{C}_{60})_n$ polymer.

Results and discussion

Polymeric complex $([10]\text{CPP}\supset\text{C}_{60})_n$

In photovoltaic devices, the absorption of photons creates excitons (coupled electron-hole pairs). Their subsequent charge separation and carrier mobility determine the overall performance of the device. Thus, not only charge separation/recombination rates, but also hole and electron transfer rates are extremely important characteristics for solar cells and artificial photosynthetic systems. The possession of the unique structural and physical properties of long π -extended [10]CPP-based polymers encourages the investigation of their response to photoexcitation. Being a rather flexible polymeric chain where monomers are interconnected by a single bond, $([10]\text{CPP})_n$ and $([10]\text{CPP}\supset\text{C}_{60})_n$ can have a significant conformational diversity. To gain insight into the dynamic shape of these polymers, I ran a series of molecular dynamics (MD) simulations for 8-unit oligomers. The simulations were performed using a classical Merck molecular force field (MMFF)

with a step size of 0.5 fs up to a total of 5 ns at 300 K within the *NVT* ensemble as implemented in the Tinker-HP program^{25,26} (for details see the ESI†). The performed MD simulations revealed interesting features of the $([10]\text{CPP})_8$ and $([10]\text{CPP}\supset\text{C}_{60})_8$ chains. In particular, the studied oligomers formed separate domains in which the units are in the *syn* or *anti* position to each other (Fig. 1a and b). As seen in the $([10]\text{CPP})_8$ oligomer, *syn* and *anti* units alternate. There are two conformations observed for this oligomer: conformation 1 was from 0.7 to 3.9 ns and conformation 2 was from 3.9 to 5 ns. They differ in the orientation of the terminal [10]CPP unit. In conformation 1, this unit is in *syn* orientation with the two adjacent units, while in conformation 2, this terminal unit adopts *anti* orientation. The representative structures of both conformers are shown in Fig. S1 (ESI†). The incorporation of the C₆₀ fullerene leads to an increase in the size of *syn*-oriented units. In the $([10]\text{CPP}\supset\text{C}_{60})_8$ oligomer, this domain increases to 3 monomeric units.

The mobility of electrons and holes are determined by the value of electronic coupling. It can be assumed that a large distance between the units in the *anti* configuration compared to those in the *syn* configuration greatly reduces the electronic coupling and thus affects the charge transport rate. Thus, the size of the *syn* domain and the number of units in the *anti* conformation are very important for efficient charge transport in such a polymer. To validate the aforementioned statement, electronic couplings were calculated for 20 randomly selected structures (10 *syn* and 10 *anti* conformations) extracted from the MD trajectory of the $([10]\text{CPP})_8$ oligomer. The results revealed that the average coupling value for the *syn* conformers is twice higher than the average coupling value for the *anti* conformers.

Based on the above, it can be assumed that an increase in the electron and hole mobility of the $([10]\text{CPP}\supset\text{C}_{60})_n$ polymer compared to the $([10]\text{CPP})_n$ polymer^{21,24} is due to an increase in the number of *syn*-oriented units. I hypothesized that the cyclodehydrogenation of the arylated CPP (monomeric unit), leading to the formation of the π -extended fragment (Fig. 1c), would increase the rigidity of the entire system and facilitate the *syn* conformation. To test this assumption, I performed MD simulations for the $([10]\text{CPP_Fused}\supset\text{C}_{60})_8$ oligomer. Already after 1.5 ns, the system formed a stable semicircular structure, in which all 8 monomeric units are in the *syn* conformation. When comparing the three considered 8-unit oligomers, a significant fluctuation of the RMSD values for $([10]\text{CPP_Fused}\supset\text{C}_{60})_8$ attracts attention. It was found that the frames with higher RMSD values correspond to a more compressed semicircular structure with high curvature. In turn, the frames with smaller RMSD values correspond to a more open structure with low curvature. The corresponding structures are shown in Fig. S2 (ESI†).

Inspired by the results, I ran simulations for polymers whose size is similar to that measured experimentally.^{21,24} Two polymers, $([10]\text{CPP_Fused}\supset\text{C}_{60})_{24}$ and $([10]\text{CPP_Fused}\supset\text{C}_{60})_{32}$, containing 24 and 32 monomeric units were analyzed. The simulations were performed under the same conditions as for



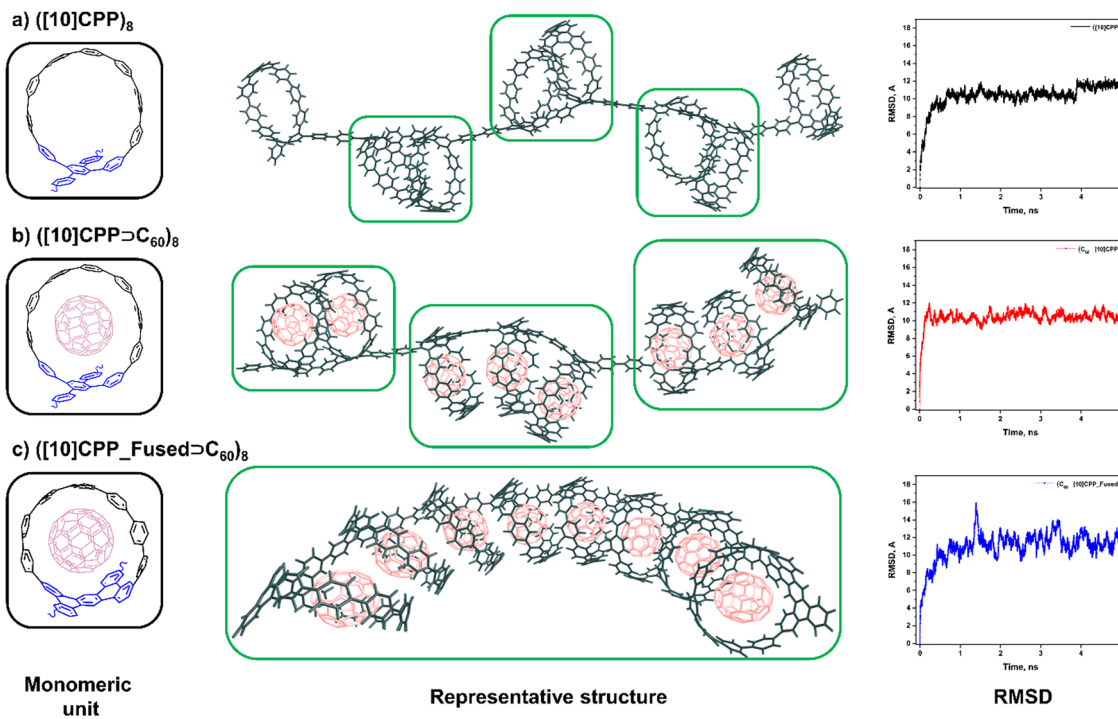


Fig. 1 Structure of monomeric units, 8-unit oligomers and RMSD diagrams for 5 ns of MD simulation runs for the (a) fullerene free $([10]\text{CPP})_8$ oligomer, (b) fullerene encapsulated $([10]\text{CPP} \supseteq \text{C}_{60})_8$ oligomer and (c) $([10]\text{CPP_Fused} \supseteq \text{C}_{60})_8$ oligomer containing an π -extended fragment.

oligomers containing 8 units, up to 10 ns in total. It is interesting to note that both systems formed stable helical structures (Fig. 2). The formation of the helical structures can be seen in Movies 1 and 2 (ESI[†]).

As seen in Fig. 2, the $([10]\text{CPP_Fused} \supseteq \text{C}_{60})_{24}$ polymer acquires a stable helical configuration after about 4 ns, while the $([10]\text{CPP_Fused} \supseteq \text{C}_{60})_{32}$ system takes about 7 ns to complete the formation process. As soon as the helical structure is formed, both considered systems retain it throughout the simulation time. RMSD graphs with the zoomed region of interest are presented in Fig. S3 (ESI[†]). It is important to note that the helical structures are very stable. To examine the thermal stability of the formed structure, MD simulations were

performed for 5 ns at 500 K on each of two helical structures independently obtained by MD simulations at 300 K. High temperature did not lead to the disruption of the formed structures or their noticeable distortion (see Fig. S4, ESI[†]).

Ground state properties of the $[10]\text{CPP_Fused} \supseteq \text{C}_{60}$ monomer

The possibility of photoinduced electron transfer (PET) in non-covalently bound donor–acceptor dyads consisting of C₆₀ as an electron acceptor and cycloparaphenylenes (CPP) as an electron donor has been previously reported for several systems.^{13,27–29} To study the possible PET processes in the polymer systems, a monomeric inclusion complex $[10]\text{CPP_Fused} \supseteq \text{C}_{60}$ (U1)

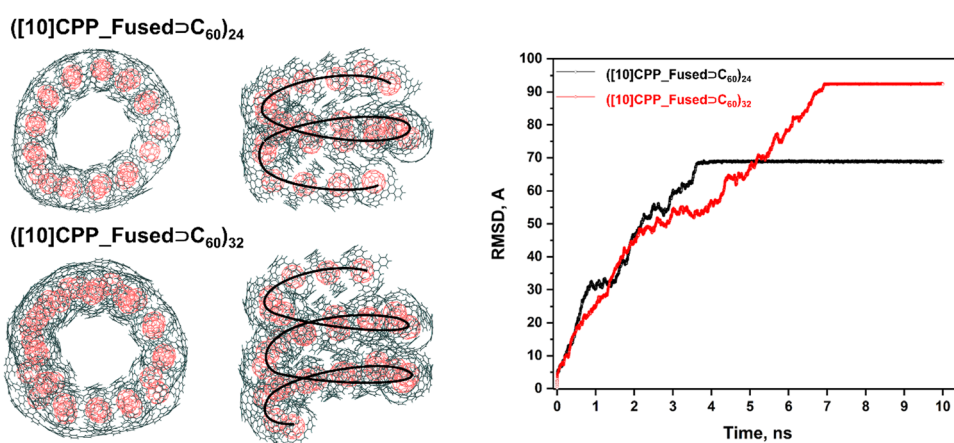


Fig. 2 Representative structure of the $([10]\text{CPP_Fused} \supseteq \text{C}_{60})_{24}$ and $([10]\text{CPP_Fused} \supseteq \text{C}_{60})_{32}$ oligomers and RMSD diagrams for 10 ns of MD simulation runs.

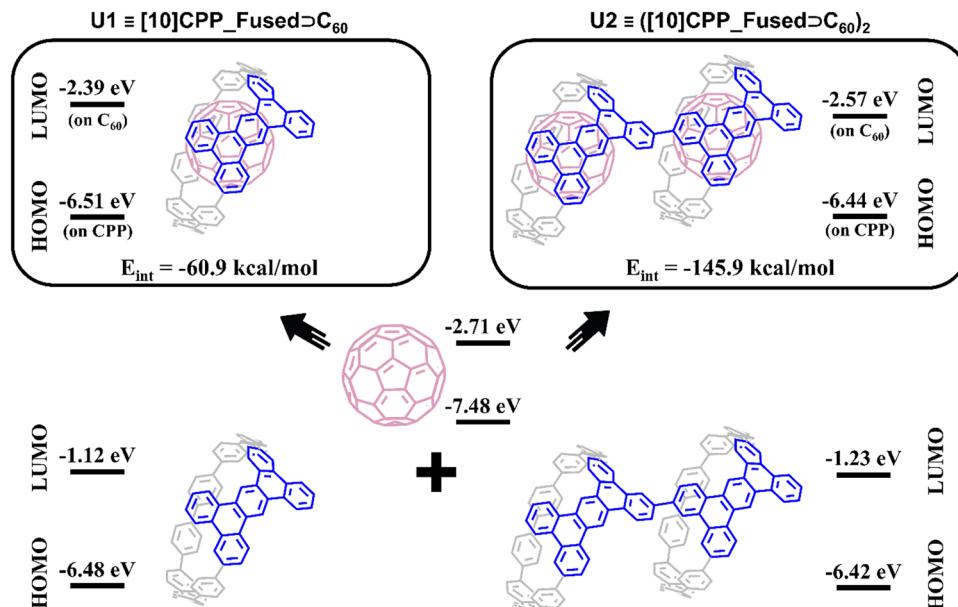


Fig. 3 Structure and HOMO/LUMO energies of **U1** and **U2** complexes and their subunits.

and its dimer $[(10)\text{CPP_Fused} \supset \text{C}_{60}]_2$ (**U2**) were considered. Geometries of **U1** and **U2** complexes (Fig. 3) were optimized using the BLYP-D3(BJ)/def2-SVP level of theory.^{30–35} (The full computational details are provided in the ESI†). For the **U1** system, the interaction energy is $-61.0 \text{ kcal mol}^{-1}$ (Table 1). This value is similar to the energy estimated for the complexes of fullerene with unmodified **[10]CPP** at the same level of theory.^{36,37} It is worth mentioning that the interaction energy in the **U2** complex was found to be superadditive. The energy is $-145.9 \text{ kcal mol}^{-1}$, which is greater than the sum of energies of pair interactions (between the host dimer and each of fullerenes). The observed superadditivity can be explained by the favorable interaction of fullerenes with each other. For details see Table S2 and Fig. S5 (ESI†). A similar effect was observed earlier for carbon nano-onions and other carbon-rich complexes.^{38,39}

As seen in Fig. 3, the HOMO and LUMO in both **U1** and **U2** complexes are localized on **[10]CPP_Fused** and C_{60} units, respectively. The formation of complexes has a rather weak effect on the HOMO and LUMO energies of the fragments. The HOMO energy changes within 0.05 eV, whereas the increase in

the LUMO energies was found to be 0.32 and 0.14 eV for **U1** and **U2**, respectively. The small differences in the HOMO and LUMO energies of complexes and individual fragments indicate the absence of charge separation in the ground state of the complexes. The performed population analysis does not reveal any significant charge transfer between the host **[10]CPP_Fused** unit and C_{60} guest molecules (Table S3, ESI†).

To analyze the nature of the host–guest interactions, I performed the Morokuma-like energy decomposition analysis (EDA)^{40–42} implemented in the ADF program at the BLYP-D3(BJ)/TZP level of theory. The EDA decomposes the interaction energy into four components: electrostatic (ΔE_{elstat}), Pauli repulsion (ΔE_{Pauli}), orbital interactions (ΔE_{oi}), and dispersion correction (ΔE_{disp}). This allows one to estimate the role of specific interactions. The EDA results listed in Table 1 indicate that the host–guest interactions in the complexes are quite similar. Among the binding forces ($\Delta E_{\text{elstat}} + \Delta E_{\text{oi}} + \Delta E_{\text{disp}}$), the dispersion term prevails with a contribution of 63–64%. The second largest term is electrostatic attraction with a contribution of 23–24%. Finally, the orbital interactions provide only 12–13% of the total stabilization interactions.

Table 1 EDA results for **U1** and **U2** complexes

Complex	Interaction scheme	Energy terms ^a , keal mol^{-1}				
		ΔE_{Pauli}	ΔE_{elstat}	ΔE_{oi}	ΔE_{disp}	ΔE_{int}
U1	$E_{\text{int}}^{\text{TOTAL}}$	83.00	-34.28 (23.8%)	-17.44 (12.1%)	-92.23 (64.1%)	-60.95
U2	$E_{\text{int}}^{\text{[(10)CPP_Fused}_2 \cdots \text{C}_{60}^1]}$	102.42	-41.60 (24.0%)	-22.60 (13.0%)	-109.39 (63.0%)	-71.16
	$E_{\text{int}}^{\text{[(10)CPP_Fused}_2 \cdots \text{C}_{60}^2]}$	99.35	-39.32 (23.6%)	-22.16 (13.3%)	-105.39 (63.2%)	-67.52
	$E_{\text{int}}^{\text{C}_{60}^1 \cdots \text{C}_{60}^2}$	20.38	-7.81 (28.2%)	-4.89 (17.6%)	-15.04 (54.2%)	-7.36
	$E_{\text{int}}^{\text{TOTAL}}$					-145.85

^a The percentage contributions to the sum of attracting energies ($\Delta E_{\text{elstat}} + \Delta E_{\text{oi}} + \Delta E_{\text{disp}}$) are given in parentheses.



To gain access to the host–guest interaction topology, I performed a series of QTAIM (Bader's Atoms in Molecules theory) and NCI (non-covalent interaction index) calculations for the **U1** and **U2** systems. Various QTAIM characteristics at the bond critical points (BCPs) can provide information on the nature of particular interaction in supramolecular complexes.⁴³ In the case of **U1**, the analysis revealed only C···C interactions (see Table S4, ESI†). The observed intermolecular interactions have $\rho(r)$ values in the range of 0.005–0.008 a.u., while the Laplacian $\nabla^2\rho(r)$ values are positive, indicating that these interactions are of closed-shell type. Moreover, $-G(r)/V(r)$ values at all BCPs are greater than 1, demonstrating the non-covalent nature of the interaction. Taking into account that the host and guest units are the π -conjugated systems, the C···C interactions can be assigned to $\pi\cdots\pi$ interactions. In the case of the **U2** complex, two types of interactions were observed: interactions between the CPP and C_{60} within one unit, intraunit interactions, and interactions between CPPs and fullerenes of different units – interunit interactions. Intraunit interactions in **U2** are very similar to those in the **U1** complex and can be assigned to pure $\pi\cdots\pi$ interactions. For interunit interactions, the C···C interactions between fullerenes and the CH···C interactions between CPPs and fullerenes of different units were observed, which can be assigned to $\pi\cdots\pi$ and C–H··· π interactions. QTAIM molecular graphs for the **U1** and **U2** complexes are given in Fig. S6 (ESI†). NCI confirms the QTAIM data. The green NCI isosurfaces between CPP and C_{60} of the same unit are evenly distributed throughout the cavity and have the similar shape for both complexes, indicating weak van der Waals interactions. In addition, the green isosurface between fullerenes and CPPs of different units was found. Thus, the results of QTAIM and NCI analyses are fully coincident and confirm a set of weak van der Waals interactions between neighboring units. The reduced density gradient (RDG) plots and NCI isosurfaces are presented in Fig. S7 and S8 (ESI†).

Singlet excited states

Simulations of excited states were performed using the TD-DFT method with the CAM-B3LYP-D3(BJ)/def2-SVP scheme.^{32–35,44} The CAM-B3LYP functional is very well suited for modeling charge transfer processes in fullerene-based complexes.⁴⁵ To characterize the properties of excited states, the **U1** and **U2** systems were divided into 2 and 4 fragments, respectively: host **[10]CPP** and guest C_{60} . The electron density distribution was analyzed for the 80 lowest excited states. Three types of excited states were identified: (1) locally excited (LE) states, where excitation is mostly localized either on the guest or the host molecule, with charge separation smaller than 0.1e (CS < 0.1e); (2) charge transfer (CT) states showing significant charge separation (CS > 0.8e); and (3) mixed states, where both LE and CT states contribute substantially (0.1e < CS < 0.8e).^{46,47}

In the gas phase, the 80 lowest vertical singlet excitation energies of the **U1** complex are found in the range from 2.51 to 4.41 eV. Table 2 contains the characteristics of the lowest excited states of each type. The LE1 state localized on C_{60} has the lowest energy, and is associated with a HOMO–4 to

LUMO+1 transition. The LE2 state with the exciton localized on **[10]CPP_Fused** lies by 0.82 eV higher in energy at 3.33 eV and corresponds to the HOMO to LUMO+3 transition. The lowest CT1 state with 0.98e transferred from **[10]CPP_Fused** to C_{60} lies at 2.74 eV, and corresponds to the HOMO → LUMO transition. Such CT state can be described as $[\text{10}]\text{CPP}_\text{Fused}^+ \supset C_{60}^-$. Among the computed excited states, only this type of charge separated state was observed. The molecular orbitals representing the LE1, LE2, and CT1 states for the **U1** complex are depicted in Fig. 4.

The energies of the 80 lowest-lying singlet excited states of **U2** due to the higher density of state found in a narrower range of energies compared to the **U1** complex and vary from 2.47 to 3.59 eV. The energies of LE1 and LE2 states in both complexes are almost identical. In **U2**, two types of CT states were identified, unlike **U1**. The first CT type, similar to **U1**, corresponds to the electron transfer from **[10]CPP_Fused** to C_{60} within the same monomeric unit. The energy of the CT1 state in **U2** is very similar to that in **U1** and equals to 2.64 eV. The second type of the CT state, corresponding to the electron transfer from **[10]CPP_Fused** of one monomer to C_{60} of another monomer, was found at 3.11 eV (0.45 eV higher than CT1). This state can be described as $([\text{10}]\text{CPP}_\text{Fused}^+ \supset C_{60}) - ([\text{10}]\text{CPP}_\text{Fused} \supset C_{60}^-)$. For the **U2** complex, within the computed 80 excited states, there are no states with a high oscillator strength. To get a clue about highly absorptive states, an additional 20 states were calculated. The state of interest has been found at 3.83 eV.

Effects of environment on excited states and electron transfer rates

To investigate the influence of the polar environment on electronic excitations, a well-proven COSMO-like model^{28,36,37,39,48,49} with dichloromethane (DCM) as a solvent was applied. DCM is a commonly used solvent in the measurement of the photophysical properties of carbon-based compounds. The dipole moments of the **U1** and **U2** complexes were calculated to be 0.09 and 0.76 D, respectively. The small dipole moment of the studied complexes can be explained by their high symmetry. The ground state (GS) solvation energies are –0.49 and –0.89 eV for **U1** and **U2**, respectively (Fig. 5). Expectedly, the solvation energies for the LE states are very similar to the GS. In turn, the solvation energies for intraunit CT1 states were found to be –0.78 and –1.13 eV. The relatively small increase in the solvation energies can be explained by a high degree of charge delocalization in both fragments and their close proximity. The interunit CT2 state observed only in the **U2** complex has a solvation energy of –1.43 eV. The computed CT2 solvation energy is almost twice as strong as for the CT1 state, which is rationalized by the noticeable spatial separation of donor and acceptor units in the CT2 state. The dipole moment differences between the GS and CT1/CT2 states in the **U2** complex were found to be 5.9 and 32.6 D, respectively. Detailed solvation data for both complexes are given in Table S5 (ESI†).

It is important to note that stabilization of the CT1 state by the solvent in both **U1** and **U2** complexes is sufficient to reorder LE1 and CT1 states, making the latter the lowest excited state



Table 2 Excitation energies (E_x , eV), main single excited configuration (HOMO(H)–LUMO(L)) and its weight (W), oscillator strength (f), extent of charge transfer (CT, e) or localization of exciton (X) computed for **U1** and **U2** complexes in the gas-phase (VAC) and dichloromethane (DCM)

	Supramolecular host-guest system			
	U1		U2	
	VAC	DCM	VAC	DCM
LE1 (fullerene C_{60})				
E_x	2.511	2.503	2.473	2.474
Transition (W)	H–4–L (0.53)	H–4–L+1 (0.35)	H–8–L (0.28)	H–8–L (0.28)
f	<0.001	<0.001	<0.001	<0.001
X	0.947	0.946	0.625	0.616
LE2 (host $[10]\text{CPP_Fused}$) ^a				
E_x	3.330	3.335	3.222	3.232
Transition (W)	H–L+3 (0.47)	H–L+3 (0.43)	H–L+6 (0.24)	H–L+6 (0.25)
f	0.066	0.102	0.008	0.012
X	0.890	0.867	0.880	0.682
Most absorptive transition (localized on $[10]\text{CPP_Fused}$)				
E_x	3.802	3.805	3.832	3.834
Transition (W)	H–2–L+3 (0.31)	H–2–L+3 (0.31)	H–1–L+6 (0.17)	H–L+6 (0.11)
f	1.587	2.224	2.248	2.518
X	0.811	0.881	0.989	0.990
CT1 intraunit ($[10]\text{CPP_Fused} \rightarrow \text{fullerene } C_{60}$)				
E_x	2.741	2.449	2.667	2.429
Transition (W)	H–L (0.93)	H–L (0.89)	H–1–L (0.66)	H–1–L (0.55)
f	0.002	0.002	<0.001	<0.001
CT	0.976	0.978	0.979	0.978
CT2 interunit ($[10]\text{CPP_Fused} \rightarrow \text{fullerene } C_{60}$)				
E_x	n/a ^b		3.112	2.578
Transition (W)			H–L+2 (0.57)	H–L+2 (0.51)
f			<0.001	<0.001
CT			0.992	0.987

^a Mixed states with significant contributions of both LE and CT. ^b n/a – states of interest are not possible for the **U1** complex.

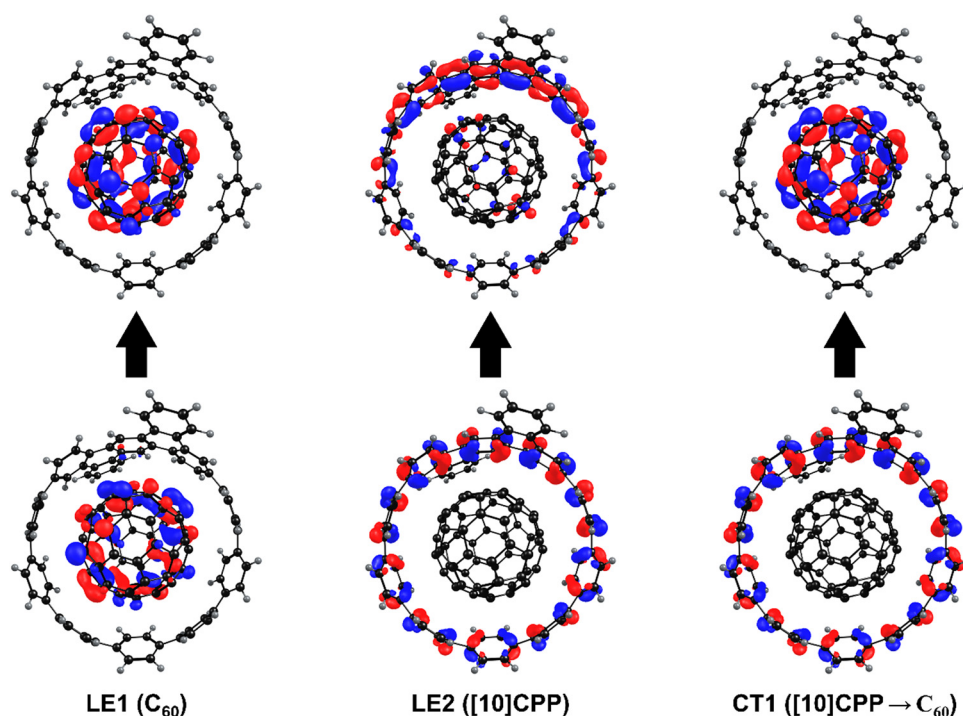


Fig. 4 Molecular orbitals representing the LE1, LE2, and CT1 states for the **U1** complex.



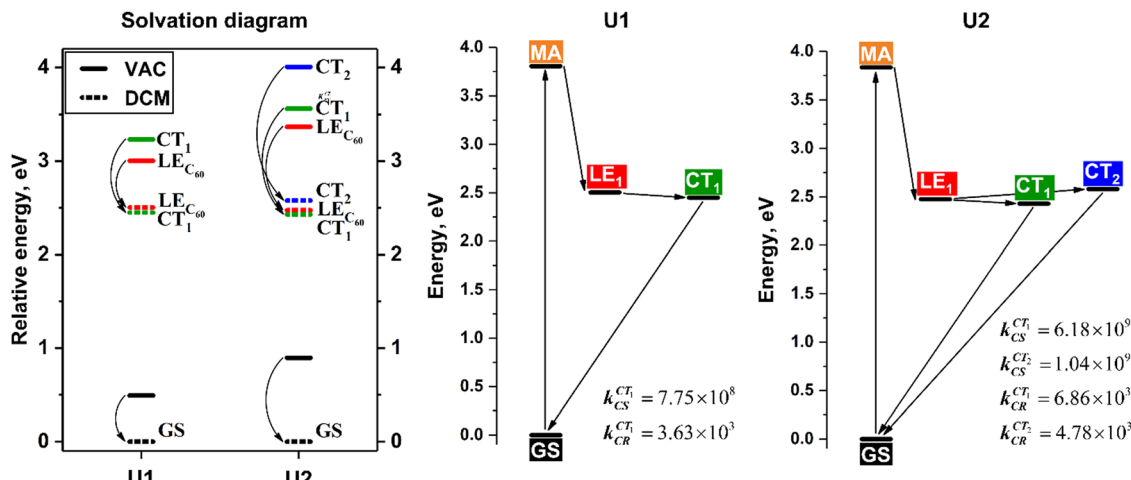


Fig. 5 Solvation diagram showing the energies of LE and CT states computed in vacuum (VAC) and dichloromethane (DCM) for **U1** and **U2** complexes and energy levels and ET/CR rates for the complexes in DCM.

Table 3 Gibbs energy ΔG^0 (in eV), electronic coupling V_{ij} (in eV), solvent (λ_s) and internal (λ_i) reorganization energy (in eV), Huang–Rhys factor (S_{eff}) and rates k_X (in s^{-1}) for charge separation and charge recombination processes computed for **U1** and **U2** complexes in DCM

Complex	Transition	ΔG^0 ^a	$ V_{ij} $	Reorg. energy		S_{eff}^b	k_X
				λ_i	λ_s		
U1	$\text{LE1} \rightarrow \text{CT1}$	-0.054	1.92×10^{-4}	0.200	0.161	1.008	7.75×10^8
	$\text{CT1} \rightarrow \text{GS}$	-2.449	9.73×10^{-3}	0.181	0.161	0.837	3.63×10^3
U2	$\text{LE1} \rightarrow \text{CT1}$	-0.045	5.39×10^{-4}	0.198	0.150	0.998	6.18×10^9
	$\text{LE1} \rightarrow \text{CT2}$	0.104	2.44×10^{-3}	0.210	0.288	1.059	1.04×10^9
	$\text{CT1} \rightarrow \text{GS}$	-2.429	1.14×10^{-2}	0.170	0.150	0.857	6.86×10^3
	$\text{CT2} \rightarrow \text{GS}$	-2.578	1.22×10^{-2}	0.155	0.288	0.781	4.78×10^3

^a Gibbs energy difference between the given states. ^b Effective value of the Huang–Rhys factor $S_{\text{eff}} = \lambda_i/\hbar\omega_{\text{eff}}$, where $\hbar\omega_{\text{eff}}$ is set to 1600 cm^{-1} .

(Fig. 5). In the case of the CT2 state in the **U2** complex, the stabilization is not enough to reorder this state with LE1, but the energy difference between LE1 and CT2 states in DCM has become less than 0.1 eV. Because of the structural similarity, it seems interesting to compare the behavior of **U1** and the well-studied [10]CPP \supset C₆₀. As was shown earlier for original [10]CPP, the charge transfer reaction is characterized by a small positive Gibbs energy. Thus, a more extended π -system of the [10]CPP_Fused nanoring leads to the stabilization of the energy of the CT state and makes charge separation thermodynamically favorable.

For both complexes under consideration, the CT states are characterized by a very weak oscillator strength and therefore cannot be directly populated by light absorption. However, they can be generated by the decay of the lowest LE states. To estimate the efficiency of a such population, I computed the electron transfer rates using the semi-classical method proposed by Ulstrup and Jortner.^{50,51} Within this approach, the intramolecular relaxation associated with ET is described by an effective vibrational mode, and the rate is controlled by four parameters: electronic coupling (V_{ij}) of the initial and final states, solvation reorganization energy λ_s , reaction Gibbs

energy ΔG^0 , and effective Huang–Rhys factor S_{eff} (for details see the ESI†). The computed parameters and rates in DCM are listed in Table 3. The rates were estimated using an effective frequency of 1600 cm^{-1} , which corresponds to the stretching of C=C bonds. Note that for similar systems the calculated rates of charge separation do not change significantly by varying the effective frequency from 1400 to 1800 cm^{-1} .^{52,53} For the **U1** and **U2** systems, the effect of frequency selection on the estimated ET rate was found to be negligible (Table S6, ESI†).

In the studied complexes, the intraunit charge separation is characterized by a small negative Gibbs energy and reaction takes place in the normal Marcus region. The estimated charge separation rates were found to be 7.75×10^8 and 6.18×10^9 in **U1** and **U2**, respectively. The rate of the interunit charge transfer reaction for **U2** equals to 1.04×10^9 , which is slightly lower compared to the intraunit CT. The charge recombination reactions take place in a deep inverted Marcus region, and their rates are significantly lower compared to the charge separation rates.

In photovoltaic applications, the efficient rate of hole- and electron-extraction is equally important for the generation of long-lived CT states with a high quantum yield. The hole

Table 4 Gibbs energy ΔG^0 (in eV), electronic coupling V_{ij} (in eV), reorganization energy (in eV), rates k_X (in s^{-1}) and characteristic times τ (in ps) for hole transfer (HT) and excess electron transfer (EET) processes computed for the **U2** complex in DCM

Complex	Transition	ΔG^0	$ V_{ij} $	Reorg. energy		S_{eff}^a	k_X	τ
				λ_i	λ_s			
U2	HT	0.022	3.54×10^{-3}	0.232	0.140	1.117	9.08×10^{10}	11
	EET	0.085	1.36×10^{-2}	0.192	0.165	0.918	1.92×10^{11}	5

^a Effective value of the Huang–Rhys factor $S_{\text{eff}} = \lambda_i/\hbar\omega_{\text{eff}}$, where $\hbar\omega_{\text{eff}}$ is set to 1600 cm^{−1}.

transfer (HT) and excess electron transfer (EET) rates were computed for the **U2** complex (Table 4). The wave function of a hole in a good approximation can be written as a linear superposition of HOMOs on the individual **[10]CPP_Fused** fragments. In turn, for the description of the excess electron, a linear superposition of LUMOs on the C_{60} fullerene was used. The HT/EET rates were estimated by using Marcus theory.⁵⁴

It is interesting to compare the charge transport properties of the originally reported ($\text{C}_{60} \supset \text{[10]CPP}_n$) polymeric conjugate and the studied ($\text{[10]CPP_Fused} \supset \text{C}_{60}$) fused system in this work. Du and co-workers²⁴ reported that the incorporation of C_{60} into a polymer dramatically increases hole mobility, making it even higher than electron mobility ($\mu_H = 3.64 \times 10^{-4}$ cm² V^{−1} s^{−1} vs. $\mu_E = 8.06 \times 10^{-5}$ cm² V^{−1} s^{−1}). Our calculations predict that in the fused ($\text{C}_{60} \supset \text{[10]CPP_Fused}$)₂ complex the rate of EET is higher than the corresponding HT rate. Their predicted characteristic times are equal to 11 and 5 ps, respectively. The EET process acceleration for the fused system is apparently associated with its supramolecular structure. As I showed at the beginning, the structure of ($\text{[10]CPP_Fused} \supset \text{C}_{60}$)₈ consists of units only in the *syn* conformation, whereas ($\text{[10]CPP} \supset \text{C}_{60}$)₈ contains the units in both *syn*- and *anti*-conformations. As seen in Fig. 1, the distance between fullerenes in the *syn*-oriented units is much smaller than that in the *anti*, and therefore the excess electron transfer *via* fullerene units in ($\text{[10]CPP_Fused} \supset \text{C}_{60}$)_n is expected to be faster than in ($\text{[10]CPP} \supset \text{C}_{60}$)_n. It is important to note that the HT/EET rates are much higher than the corresponding charge recombination rates, allowing for efficient electron and hole extraction in a polymer solar cell.

Conclusions

In this work, the molecular dynamics simulations and DFT/TD-DFT approaches were combined to study the ground and excited state properties of the supramolecular system consisting of π -conjugated CPP-based polymers and C_{60} guests. I demonstrated that π -extension in **[10]CPP_Fused** \supset C_{60} increases the rigidity of the entire system, but most importantly facilitates the *syn* conformation of the monomeric units in the polymer that accelerates the charge transport through oligomer. MD simulations suggest that ($\text{[10]CPP_Fused} \supset \text{C}_{60}$)₂₄ and ($\text{[10]CPP_Fused} \supset \text{C}_{60}$)₃₂ oligomers tend to form stable, helix-like structures. The TD-DFT results obtained for the ($\text{[10]CPP_Fused} \supset \text{C}_{60}$)₁ and ($\text{[10]CPP_Fused} \supset \text{C}_{60}$)₂ host–guest complexes indicate the efficient PET from the **[10]CPP** unit to

the C_{60} fullerene. The ET occurs in the normal Marcus region on the nanosecond timescale. The hole transfer and excess electron transfer processes occur much faster than the corresponding charge recombination, which ensures their sufficient extraction. Overall, the results suggest that the proposed ($\text{[10]CPP_Fused} \supset \text{C}_{60}$)_n oligomer is a promising material for the ordered photoactive layer with the regular distribution of donor and acceptor phases for efficient charge separation.

Conflicts of interest

There are no conflicts to declare.

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