

# Nonadiabatic dynamics of cycloparaphenylenes with TD-DFTB surface hopping

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## Abstract

We implemented a version of the decoherence-corrected fewest switches surface hopping based on linear-response time-dependent density functional tight binding (TD-DFTB), enhanced by transition density analysis. The method has been tested for the gas-phase relaxation dynamics of two cycloparaphenylene molecules, [8]CPP and [10]CPP, explaining some important features of their nonadiabatic dynamics, such as the origin of their long fluorescence lifetimes (related to the slow radiative emission from the  $S_1$  state) and the trend of increasing the fluorescence rate with the molecular size (related to an increase in the  $S_1$ - $S_0$  energy gaps and oscillator strengths in the larger molecule). The quality of the TD-DFTB electronic structure information was assessed through four quantities: excitation energies; charge-transfer (CT) numbers, which estimates the charge transfer character of states; participation ratio (PR), which describes delocalization of electronic density; and participation ratio of natural transition orbitals (PRNTO), which describes the multiconfigurational character of states. These quantities were computed during dynamics and recomputed for the same geometries with the higher level long-range corrected TD-LC-DFTB and a lower level single-determinant approximation for the excited states, SD-(LC)-DFTB. Taking TD-LC-DFTB as the standard, TD-DFTB underestimates the excitation energies by about 0.5 eV and overestimates CT and PR. SD-DFTB underestimates excitation energies and overestimates CT to the same extent that TD-DFTB does,

but it predicts reasonable PR distributions. SD-LC-DFTB leads to an extreme overestimation of the excitation energies by approximately 3 eV, overestimates the charge transfer character of the state, but predicts the PR values very close to the ones obtained with TD-LC-DFTB.

## 1. Introduction

Nonadiabatic effects influence the underlying relaxation mechanisms of a range of processes occurring after photoexcitation in systems of all sizes, spanning from single molecules to solid-state materials. Some of these nonadiabatic processes such as charge and energy transfer, electron-hole separation, singlet fission, etc.<sup>1-5</sup> have attracted a lot of attention due to their potential to aid the development of photonic and photovoltaic materials. Therefore, a number of research groups have been aiming to develop and extend suitable theoretical algorithms for simulation of nonadiabatic processes in extended systems.<sup>6-8</sup> Since the dimensionality of extended systems is a major limiting factor in their excited-state dynamics simulations, the methods that have been used for this purpose usually rely on mixed quantum-classical nonadiabatic dynamics approaches, such as the trajectory surface hopping in its different variants.<sup>6-7,9</sup> These methods are often associated with fast semiempirical algorithms for electronic structure computations.<sup>7,10-15</sup>

The density functional tight binding (DFTB)<sup>16</sup> method bridges the gap between first-principles electronic structure methods and empirical tight-binding approaches. DFTB can be seen as an approximation of density functional theory (DFT) at the generalized-gradient approximation (GGA) level. The basic algorithms implemented in the DFTB+ code,<sup>17</sup> which we use here, are the non-self-consistent charge (non-SCC)<sup>18-19</sup> and SCC-DFTB (DFTB2 and DFTB3)<sup>16</sup> for the electronic ground state. The treatment of dispersion interactions (+D) is possible in the DFTB+ code via Lennard-Jones potentials,<sup>20</sup> with the Slater-Kirkwood polarizable atomic model,<sup>21</sup> and on the level of the DFT-D3 method.<sup>22</sup> Electronic excitations can be calculated with linear-response time-dependent approach (LR TD-DFTB).<sup>23</sup> A recent extension of the method now also allows for computations using long-range corrected (LC) functionals,<sup>24-25</sup> which provide a better description of state localization and charge transfer excitations. The low computational cost of these algorithms renders levels up to LR TD-(LC)-DFTB3 (+D3) suitable for simulations of excited states in extended systems.

In this work, we implemented an interface between DFTB+<sup>17</sup> and Newton-X,<sup>26-27</sup> allowing to run decoherence-corrected fewest switches trajectory surface hopping (FSSH) with the TD-DFTB method<sup>23</sup> available in the recently released 1.3 version of the DFTB+ code. There are reports of few other implementations of nonadiabatic dynamics with mixed quantum-classical methods based on DFTB. Jakowski and Morokuma have implemented Ehrenfest dynamics based on Liouville-von Neumann equation

for density matrices propagation and made several applications with the SCC-DFTB method.<sup>28</sup> Ehrenfest dynamics based on real-time Kohn-Sham DFTB has been described in Refs. <sup>14, 29</sup>. Surface hopping dynamics, also based on real-time Kohn-Sham DFTB, is discussed in Refs.<sup>10, 15</sup>. In these surface hopping implementations, the excited states are represented by single Kohn-Sham determinants in a basis of stationary orbitals (SD-DFTB), which may lead to an inaccurate description of adiabatic crossings.<sup>30</sup> Surface hopping with a more robust description of the excited states should be obtained via linear-response TD-DFTB. Nonadiabatic dynamics with linear-response TD-DFTB, which has also been implemented by Mitrić *et al.*<sup>11</sup> based on an earlier version of DFTB, is the basis of our developments here.

Our surface hopping based on linear-response TD-DFTB implementation relies on a general interface between the newest versions of the DFTB+ and Newton-X codes, which are both publicly available free of charge. It enables diverse new features to improve the description of ground and excited state energies, such as the DFTB third-order expansion (DFTB3 model)<sup>31</sup> and onsite corrections for TD-DFTB,<sup>32</sup> to accelerate the dynamics, such as fast nonadiabatic coupling calculations based on the recently proposed orbital derivative method;<sup>33</sup> and to enhance the data analysis, via a direct module for advanced transition density analysis based on the program TheoDORE.<sup>34</sup> Moreover, this interface is ready to profit from analytical energy gradients for TD-DFTB based on long-range corrected functionals,<sup>24-25</sup> which are currently under development.

The FSSH/TD-DFTB dynamics (on the GGA level of functional) has a potential for the simulation of processes where the charge-transfer states are not involved in dynamics (e.g. relaxation processes of small molecules in solutions, where both solute and solvent molecules are treated with TD-DFTB, see e.g. Ref.<sup>11</sup>), or in the cases where the occurring charge-transfer states are of short-distance (like for instance in the process of hot electron injection in photovoltaic materials, see e.g. Ref.<sup>35</sup>). Even in cases where TD-DFTB is not expected to work well, mainly due to artifacts in the description of charge-transfer states, a multilayer approach may be devised,<sup>36</sup> where dynamics is propagated with TD-DFTB and single points are recalculated at higher level. Naturally, in such a case, time-dependent properties (population lifetimes, for instance) may not be adequately described, while statistical distributions of the excited-state in equilibrium may still deliver relevant information.

As a test case for this new implementation, we choose to study the nonadiabatic dynamics of two cycloparaphenylenes, [8]CPP and [10]CPP. Cycloparaphenylenes are hoop-shaped conjugated molecules composed of repeating phenylene units bonded in *para* positions to form a ring. They have been synthesized by several experimental groups<sup>37</sup> and since then studied experimentally<sup>38-41</sup> and theoretically<sup>42-44</sup> in detail. They received a lot of attention because of their potential use as precursors in the synthesis of armchair

carbon nanotubes<sup>45</sup> and as efficient emitters of tunable wavelengths.<sup>46</sup> CPPs show unusual optical properties, markedly distinct from the linear paraphenylenes,<sup>47</sup> such as size-independent absorption wavelengths, blue shifting of emission peaks with an increase in the ring size<sup>39</sup>, double-peaked emission spectra,<sup>39</sup> among others. CPP excitations are characterized by long relaxation lifetimes in the nanosecond domain.<sup>41, 48</sup> Small CPPs are also interesting due of their large quantum yields for singlet oxygen production in the reaction with triplet oxygen.<sup>41, 48</sup>

As we shall discuss, nonadiabatic dynamics with LR TD-DFTB explore states with somewhat large CT character. This is not surprising given that conventional DFTB is parameterized for PBE, where exactly this same type of problem is expected to occur.<sup>49</sup> Recent results for (LR) TD-LC-DFTB make clear that this level provides correct energy character, fixing the main deficiencies of conventional DFTB.<sup>25</sup> Since long-range corrected functionals are indispensable for the correct description of charge-transfer states, but the analytical gradients for long-range corrected TD-DFTB are under development at the moment, we resort to a multilayer dynamics scheme, where the conventional simulations are performed with TD-DFTB and the TD-LC-DFTB computations of the excited states populated during dynamics are performed *a posteriori*. This approximation precludes us of drawing out conclusions about states' lifetimes, because of the mismatch between TD-DFTB and TD-LC-DFTB gradients. Still, plenty of important insights about relaxation mechanisms could be deduced, primarily because the long-enough dynamics provides mapping of the excited states hypersurfaces, particularly the  $S_1$  hypersurface, and thus fluorescence and internal conversion processes can be studied, as well as the properties of occurring excitons.

The Newton-X/DFTB+/TheoDORE interface that we present in this paper allows also the analysis of transition density matrices of the states populated during dynamics. We utilized this feature to perform the statistical analysis of excitation energies, charge transfer, density delocalization, and multiconfigurational character for a large ensemble of points generated during the dynamics, based on different approximations of the  $S_1$  state wave functions, (LR) TD-(LC)-DFTB and SD-(LC)-DFTB. This comparison allows us to derive a comprehensive picture about what kinds of deviations in the electronic densities are caused by an exploration of the configurational space by GGA functionals and single determinant approximations. This analysis confirmed that the full LR TD-DFTB representation of the wave function is a better approximation for the excited states than the single-determinant one, often applied in the simulations of nano-sized systems.<sup>10</sup>

This paper is organized as follows. In the section Method Description, we briefly review the fewest switches surface hopping and TD-DFTB methods, providing details on the interfaces between DFTB+, Newton-X, and TheoDORE codes. Computational Details contains the technical information about the methods used

for the actual calculations. In the section Results and discussion, we first discuss the reliability of the description of the excited states of the [8]CPP and [10]CPP on the TD-DFTB level, comparing the vertical excitations with TD-DFT results calculated with three different functionals. Next, we discuss the TD-DFTB excited state dynamics of cycloparaphenylenes. Finally, we statistically analyze the quality of the TD-DFTB electronic structure information produced during the dynamics and evaluate the SD-(LC)-DFTB levels.

## 2. Method Description

### 2.1. Fewest switches trajectory surface hopping (FSSH)

FSSH is a mixed quantum-classical approach to quantum dynamics, which approximates the wave packet motion by a swarm of independent nuclear trajectories, each one evolving classically on a single electronic state.<sup>50</sup> Nonadiabatic transitions between states are introduced by a stochastic algorithm which may instantaneously switch the surface on which the trajectory is propagated, based on evaluating of hopping probabilities.<sup>50</sup> The hopping probabilities in FSSH are obtained by expanding the time-dependent electronic wave function as

$$\Psi(\mathbf{r}; \mathbf{R}(t)) = \sum_K c_K(t) \psi_K(\mathbf{r}; \mathbf{R}(t)). \quad (1)$$

$\psi_K(\mathbf{r}; \mathbf{R}(t))$  is the adiabatic electronic wave function for state  $K$ , which is a function of the electron coordinates  $\mathbf{r}$  and parametrically dependent on the classical nuclear coordinate  $\mathbf{R}$ .  $c_K(t)$  are complex time-dependent expansion coefficients. The nuclear trajectories  $\mathbf{R}(t)$  are propagated by integrating Newton's equations on the active state  $M$ .

Simultaneously to the propagation of Newton's equations, a local-approximation of the time-dependent electronic Schrödinger equation (TDSE) is integrated to obtain the expansion coefficients,  $c_K(t)$ . In the adiabatic representation, it is given by

$$i\hbar \frac{dc_J}{dt} = E_J c_J - i\hbar \sum_K \sigma_{JK} c_K, \quad (2)$$

where  $E_J$  is the energy of the electronic state  $J$  and the bra-ket in the second term of the right side represents the time-derivative nonadiabatic coupling (NACs) between states  $J$  and  $K$

$$\sigma_{JK} = \left\langle \psi_J \left| \frac{d\psi_K}{dt} \right. \right\rangle. \quad (3)$$

The coefficients  $c_I$  are further corrected for decoherence effects with the simplified decay of mixing method (SDM)<sup>51</sup> according to

$$\begin{aligned} \bar{c}_K &= c_K e^{-\Delta t/\tau_{KM}}, \quad \forall K \neq M, \\ \bar{c}_M &= \frac{c_M}{|c_M|} \left[ 1 - \sum_{K \neq M} |\bar{c}_K|^2 \right]^{1/2}. \end{aligned} \quad (4)$$

In these equations,  $M$  is the active state and the decoherence time  $\tau_{KM}$  is given by the phenomenological equation

$$\frac{1}{\tau_{KM}} = \frac{|E_K - E_M|}{\hbar} \left( 1 + \frac{\alpha}{K_n} \right)^{-1}, \quad (5)$$

where  $E_I$  is the potential energy of state  $I$ ,  $K_n$  is the classical kinetic energy of the nuclei, and  $\alpha$  is a parameter whose recommended value is 0.1 Hartree.

The FSSH probability<sup>50</sup> between the active state  $M$  and another state  $K$  at each time step is computed as

$$P_{M \rightarrow J} = \max \left[ 0, \frac{-2\Delta\tau}{|c_M|^2} \sigma_{MJ} \operatorname{Re}(\bar{c}_M^* \bar{c}_J) \right], \quad (6)$$

where  $\Delta\tau$  is the time step for the integration of the TDSE. Thus, once the initial conditions (positions and velocities) are created, to perform the FSSH simulations, one must solve the electronic TDSE at each nuclear position given by Newton's equations to obtain the excitation energies, gradients, and to evaluate time-dependent NACs. These quantities are interpolated between two classical time steps to be used in the integration of the TDSE.

Our FSSH/TD-DFTB implementation profits of the algorithmic infrastructure already implemented in Newton-X. In the standard way, time-dependent NACs are computed by finite differences method as proposed by Hammes-Schiffer and Tully.<sup>52</sup> In this approach, TD-NACs are given in terms of excited-state wave function overlaps between two subsequent nuclear time steps,

$$\sigma_{JK}(t) \approx \frac{1}{4\Delta t} \left[ 3S_{JK}(t) - 3S_{KJ}(t) - S_{JK}(t - \Delta t) + S_{KJ}(t - \Delta t) \right], \quad (7)$$

where

$$S_{JK}(t) = \langle \psi_J(t - \Delta t) | \psi_K(t) \rangle. \quad (8)$$

The wavefunction overlaps  $S_{JK}$  can be evaluated on a basis of Slater determinants (determinant derivative method).<sup>53</sup> The TD-NACs can be alternatively evaluated with the recently implemented orbital derivative method<sup>54</sup> developed by Izmaylov’s group,<sup>33</sup> which relies on time derivatives in a basis of molecular orbitals. In this case, the calculation of the  $S_{JK}$  overlaps is skipped, and the NACs are directly computed in terms of molecular orbital (or KS orbital) overlaps

$$s_{ij}(t) = \langle \chi_i(t - \Delta t) | \chi_j(t) \rangle. \quad (9)$$

Instead of the integration of the TDSE equation, having  $S_{JK}$  computed with either method described above, the time-dependent coefficients  $c_K$  can also be obtained via the local diabaticization method<sup>55</sup> proposed by Granucci and Persico.<sup>13</sup> In this case, the coefficients are obtained by a unitary transformation between two time steps, with the help of an adiabatic-to-diabatic transformation matrix obtained under the condition  $\sigma_{JK} = 0$ .

## 2.2. The basics of the TD-DFTB method

The basic idea of the density functional based tight binding (DFTB) method is to perform a Volterra expansion of the Kohn-Sham (KS) DFT total energy functional around a reference electronic density ( $\rho_0$ ),<sup>16</sup>

$$E^{DFTB}[\rho] = \underbrace{E^0[\rho_0]}_{DFTB1} + \underbrace{E^1[\rho_0, \delta\rho]}_{DFTB2} + \underbrace{E^2[\rho_0, (\delta\rho)^2]}_{DFTB3} + E^3[\rho_0, (\delta\rho)^3] + \dots, \quad (10)$$

where  $\rho_0$  is the sum of neutral atomic densities. Within the DFTB+ code, the second-order (SCC-DFTB or DFTB2)<sup>16</sup> and the third-order (DFTB3)<sup>56</sup> expansions of the density functional have been implemented.

The total energy within DFTB is derived employing several approximations.<sup>16</sup> A minimal basis set of localized atomic orbitals, obtained by solving the DFT equations for atoms, is used in the expansion of KS orbitals. The matrix elements of the part of the Hamiltonian that depends only on the reference density and orbitals are evaluated based on DFT computations. Its diagonal elements are simply equal to the computed atomic orbital energies, and the nondiagonal elements are precomputed for pairs of elements for a range of

internuclear distances and tabulated in so-called Slater-Koster (SK) files.<sup>16</sup> Overlaps of atomic orbitals are likewise precomputed and stored.<sup>16</sup>

The terms of the energy expression depending on the electronic density fluctuation  $\delta\rho$  are simplified using the Mulliken approximation by means of atomic point charges.<sup>16</sup> Remaining integrals are parameterized using Hubbard-like parameters for each atom computed with DFT. Furthermore, the nuclear repulsion term and the terms which depend only on the reference density are approximated as a sum of short-range repulsive potentials which depend only on diatomic distance.<sup>16, 56-57</sup>

The LR TD-DFTB method applies a similar procedure as LR TD-DFT in the computation of excitation energies,<sup>23</sup> which are given as the eigenvalues  $\omega$  of the equation

$$\begin{bmatrix} \mathbf{A} & \mathbf{B} \\ \mathbf{B}^* & \mathbf{A}^* \end{bmatrix} \begin{bmatrix} \mathbf{X} \\ \mathbf{Y} \end{bmatrix} = \omega \begin{bmatrix} 1 & 0 \\ 0 & -1 \end{bmatrix} \begin{bmatrix} \mathbf{X} \\ \mathbf{Y} \end{bmatrix}, \quad (11)$$

where the matrix elements are written in terms of orbital transitions from occupied orbitals ( $i, j$ ) into virtual orbitals ( $a, b$ ) as

$$\begin{aligned} A_{ia,jb} &= \delta_{ij} \delta_{ab} (\varepsilon_a - \varepsilon_i) + (ia||jb), \\ B_{ia,jb} &= (ia||bj). \end{aligned} \quad (12)$$

In these equations,  $(ia||jb)$  are two-electron integrals that involve both Coulomb and exchange-correlation kernels and  $\varepsilon$  represents the orbital energies from a previous DFTB ground state calculation. In the framework of TD-DFTB, the two-electron integrals are calculated applying the generalized Mulliken approximation, according to which the transition densities between different Kohn-Sham orbitals are represented by point charges.<sup>23</sup> The LR TD-DFTB method has recently been expanded beyond the generalized Mulliken approximation by introducing onsite exchange-like integrals.<sup>32</sup> This correction improves the description of  $n\pi^*$  and  $\sigma\pi^*$  states, which were not well described in the original version of LR TD-DFTB.<sup>32</sup>

Moreover, long-range corrected (LC) exchange-correlation functionals have also been implemented and benchmarked in the DFTB method (LC-DFTB)<sup>24</sup> and in the LR TD-DFTB method for excited state simulations (TD-LC-DFTB).<sup>24</sup> Benchmark calculations on a set of small chromophores showed that the TD-LC-DFTB method can treat charge transfer excitations accurately. For this class of excitations, DFTB (like any DFT method) using the generalized-gradient approximation underestimates excitation energies strongly.<sup>25</sup> Although the lack of analytical excited-state gradients within TD-LC-DFTB precludes the use of

this approach in surface hopping simulations at the moment; we will still use it here as benchmark results for single point calculations.

### 2.3. Time-dependent wave function overlaps with TD-DFTB

As discussed above, the core quantity to compute the hopping probabilities are NACs, which can be calculated in terms of the wave function overlaps  $S_{JK}$  (Eq. (8)) or molecular (or KS) orbital overlaps  $s_{ij}$  (Eq. (9)), depending on the approach chosen. The algorithm for computation of these quantities has been already implemented in the Newton-X code for the calculation of the TD-DFT NACs.<sup>53</sup> Here we only describe them briefly to point out the approximations used in the case of TD-DFTB.

The Casida approximation—stating that the excited state wave function can be represented using a CIS-like expansion on the basis of singly-excited Slater determinants—is applied.<sup>58</sup> This approximation was previously utilized in the implementation of the surface hopping dynamics with the TD-DFT method.<sup>59-61</sup> The number of the singly-excited Slater determinants that is obtained by applying a single excitation operator to the ground state Slater determinant is  $N_{det} = N_{occ}N_{virt}$ , and the form of the excited state wave function is

$$|\psi_J\rangle = \sum_{i,a} (X + Y)_{ia}^J |\Phi_{ia}\rangle, \quad (13)$$

where the CIS coefficients  $\mathbf{X} + \mathbf{Y}$  are the eigenvectors of Eq. (11) and  $|\Phi_{ia}\rangle$  are the singly-excited Slater determinants.

The overlap of the excited state wave functions represented in the basis of Slater determinants at time steps  $t$  and  $t + \Delta t$  is

$$\langle \psi^I(t) | \psi^J(t + \Delta t) \rangle = \sum_{ia,jb} (X + Y)_{ia}^{I*} (X + Y)_{jb}^J \langle \Phi_{ia}(t) | \Phi_{jb}(t + \Delta t) \rangle. \quad (14)$$

The overlap of two Slater determinants is computed as the determinant of the overlap matrix of KS orbitals  $\chi_i$  obtained at two subsequent time steps.<sup>62</sup> Furthermore, the overlaps of KS orbitals (Eq. (9)) are represented in terms of overlaps of atomic orbitals  $\phi_\mu$  at two subsequent geometries from the dynamics:

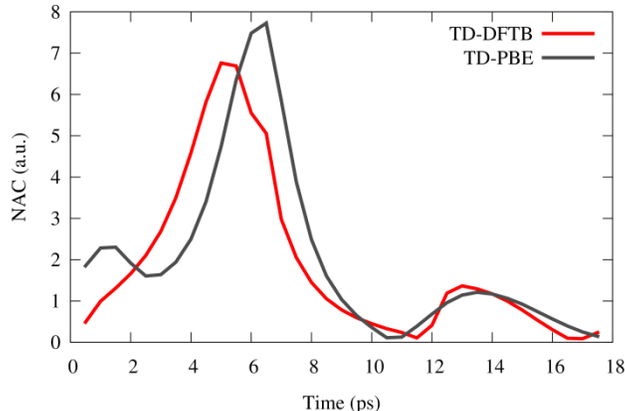
$$\langle \chi_i(t) | \chi'_j(t + \Delta t) \rangle = \sum_{\mu\nu} c_{\mu i}^*(t) c_{\nu j}(t + \Delta t) \langle \phi_\mu(t) | \phi'_\nu(t + \Delta t) \rangle. \quad (15)$$

These atomic orbitals do not form an orthonormal set because they are computed at subsequent geometries in the dynamics, which slightly differ from each other. As already mentioned, the atomic orbital overlaps are precomputed in DFTB at different internuclear distances and stored in Slater-Koster (SK) files. Thus, during an SCC-DFTB computation, these integrals are not calculated, but they are read from the SK files.

During the dynamics, it might happen that some of the atom pairs from two subsequent geometries are displaced with respect to each other by very short distances, which are not covered within the precomputed atomic orbital overlaps because they are not relevant for simulations of single molecules. In this case, we evaluate NACs by applying a linear extrapolation using the computed values from the last two time steps. This problem does not happen with the 3<sup>rd</sup> order DFTB and the 3ob Slater-Koster set optimized for it, because atomic orbital overlaps in the standard 3ob set are computed for interatomic distances larger than 0.02 Bohr. This problem might occur in simulations with the 2<sup>nd</sup> order DFTB, where the mio Slater-Koster set should be used, in which the atomic orbital overlaps are defined for minimum interatomic distance of 0.4 Bohr. Therefore, we suggest using the 3<sup>rd</sup> order DFTB and the 3ob basis set to circumvent the extrapolation of the nonadiabatic couplings.

Since DFTB is a valence-electron-only method, core occupied orbitals do not appear in the construction of Slater determinants. This fact speeds up the computation of wave function overlaps for two reasons: (1) the dimension of the Slater determinants is reduced; (2) the number of Slater determinants is decreased from  $N_{occ}N_{virt}$  to  $(N_{occ} - N_{core})N_{virt}$ . The formal scaling of the determinant derivative method is hence reduced from  $N_{occ}^5 N_{virt}^2$  to  $(N_{occ} - N_{core})^5 N_{virt}^2$  and the scaling of the orbital overlap method is reduced from  $N_{occ} N_{virt}^2$  to  $(N_{occ} - N_{core})N_{virt}^2$ .

We tested the nonadiabatic couplings between  $S_3$  and  $S_2$  states in one sequence of the TD-DFTB surface hopping dynamics of thiophene by comparing the computed NACs between these states with the values computed from the TD-PBE wave functions imposing the geometries from the TD-DFTB dynamics. In the case of the TD-DFT wave functions, all occupied orbitals are used in the construction of ground state and excited Slater determinants. In both cases, the NACs were computed with the local diabaticization method.<sup>55</sup> The values of NACs obtained from TD-DFTB and TD-PBE wave functions are in excellent agreement (Figure 1), which indicates that the neglect of core molecular orbitals does not significantly affect the excited-state overlaps. Moreover, TD-DFTB excited-state wave functions seem to approximate their TD-DFT counterparts accurately also in regions of strong nonadiabatic coupling.



**Figure 1.** Comparison of nonadiabatic couplings (a.u.) between  $S_3$  and  $S_2$  states in the sequence of a trajectory of thiophene computed based on the TD-DFTB and TD-PBE wave functions.

#### 2.4. Excited-state transition density matrix analysis

To analyze the nature of the excited states occurring during TD-DFTB surface hopping dynamics, we interfaced the TheoDORE program with the DFTB+ program. TheoDORE is a program suite intended for the electronic wave function analysis based on state and transition density matrices between electronic states.<sup>63-65</sup> The implemented methods are particularly suitable for obtaining insight into excitonic structure in large conjugated systems<sup>66-67</sup> and are, therefore, a natural choice for analyzing CPPs. The most important part of this analysis refers to the determination of charge-transfer character of the excited states and their spatial localization properties by computing charge transfer and fragment participation ratio numbers. Additionally, the TheoDORE program also implements the computation of the natural transition orbitals of the excited states and their participation ratio of natural transition orbitals.

Segmenting a molecule into suitable fragments, one can evaluate the extent of charge transfer between fragments upon excitation by computing the value

$$\Omega_{AB}^J = \frac{1}{2} \sum_{\substack{a \in A \\ b \in B}} (D_{ab}^{0J})^2, \quad (16)$$

where  $a$  and  $b$  are molecular orbitals belonging to fragments  $A$  and  $B$ ,  $D_{ab}^{0J}$  is the element of the transition density matrix in the localized basis, for the transition from the ground state  $0$  into the excited state  $J$ .<sup>63</sup>

The charge transfer properties of the excited states are described with the charge transfer (CT) index, which is defined as a sum of the nondiagonal  $\Omega_{AB}^J$  elements divided by their total sum ( $\Omega$ ),

$$CT_J = \frac{1}{\Omega^A} \sum_{B \neq A} \Omega_{AB}^J. \quad (17)$$

They have values between 0, in the case of localized excitations (Frenkel excitons), and 1, in the case of charge transfer excitations.

The participation ratio (PR) defines the number of fragments over which an exciton is delocalized. It is defined as<sup>63</sup>

$$PR_J = \frac{1}{2} \left[ \sum_A \left( \sum_B \Omega_{AB}^J \right)^2 + \sum_B \left( \sum_A \Omega_{AB}^J \right)^2 \right], \quad (18)$$

where the first term quantifies the extent of the hole delocalization and the second term the extent of the electron delocalization over fragments.

A compact description of the excited states in terms of a relatively small number of transitions can be achieved in the basis of natural transition orbitals (NTOs).<sup>68</sup> The NTOs are obtained by a singular value decomposition of the transition density matrix represented in a molecular orbital basis,  $\mathbf{D}^{0,J}$ .

$$\mathbf{D}^{0,J} = \mathbf{U} \text{diag}(\sqrt{\lambda_1}, \sqrt{\lambda_2}, \dots) \mathbf{V}^T, \quad (19)$$

where the matrices  $\mathbf{U}$  and  $\mathbf{V}$  collect the sets of hole and electron orbitals, respectively, and  $\lambda_i$  are the weights of the corresponding transitions between them.<sup>63</sup>

The participation ratio of natural transition orbitals (PRNTO) defined as

$$PRNTO = \frac{(\sum_i \lambda_i)^2}{\sum_i \lambda_i^2}, \quad (20)$$

quantifies the number of transitions between NTOs necessary to describe the excited state. This quantity was initially introduced as a collectivity measure in Ref. <sup>69</sup> and its physical meaning has recently been elucidated in the context of quantum entanglement.<sup>70</sup> Its upper bound is the number of nonzero singular values ( $N_{SVD}$ ) of  $\mathbf{D}^{0,J}$ , and it is equal to this number in the case when all nonzero singular values have the same magnitude. Oppositely, the lowest value of PRNTO, reached in the case when the excited state can be represented by only one transition between NTOs, is 1.<sup>63</sup>

### 3. Computational Details

We created sets of 500 initial conditions (ICs) (geometries and momenta) using a harmonic-oscillator Wigner distribution,<sup>71</sup> based on normal modes in the ground states of [8]CPP and [10]CPP. The vertical excitation energies and oscillator strengths for the transitions to the six lowest-lying states were computed with the TD-DFTB method with the third-order expansion of density<sup>32</sup> and 3ob-1-1 SK set<sup>72</sup> (Cartesian coordinates of the ground states of [8]CPP and [10]CPP are given in the Supporting Information, SI-I). The dispersion interactions were included via a Lennard-Jones potential<sup>21</sup> with parameters adopted from the Universal force field (UFF).<sup>20</sup> The sampled geometries are used in the simulations of the photoabsorption spectra with the nuclear ensemble approach<sup>73</sup> employing TD-DFTB excitation energies and oscillator strengths. The simulated spectra of both molecules are featured by broad bands peaked in both cases at  $\approx 2.95$  eV (Supporting Information, SI-II).

Nonadiabatic excited-state dynamics simulations were performed on TD-DFTB potential energy surfaces. The sets of ICs for dynamics propagation were selected from the initial set of 500 ICs starting from the bright  $S_6$  and  $S_5$  states applying a very narrow energy window ( $2.95 \pm 0.05$  eV) around the maxima of the absorption spectra. This procedure produced 72 ICs in the  $S_5$  state and 35 ICs in the  $S_6$  state for [8]CPP. We selected 35 ICs from the  $S_5$  and 17 ICs from the  $S_6$  state for dynamics propagation. In the case of [10]CPP, 64 ICs in the  $S_5$  state and 32 initial conditions in the  $S_6$  state are filtered, from which we selected 32 ICs in the  $S_5$  and 16 ICs in the  $S_6$  state for dynamics propagation. We limited the number of trajectories to  $\approx 50$  per molecule having in mind that the 3-ps propagation of larger number of trajectories would require substantial computational time.

The TD-DFTB dynamics (GGA functional) of [8]CPP lasted  $\sim 10$  CPU-minutes per time step (on Intel(R) Xeon 3.0 GHz processor), from which the electronic structure and gradients computations elapsed  $\sim 90\%$  of the time and  $\sim 10\%$  of the time was spent in nonadiabatic couplings computation. For the comparison, the same simulations with the TD-PBE/6-31G method require  $\sim 110$  CPU-minutes for the electronic structure and gradient computations and  $\sim 2$  minutes for nonadiabatic coupling computations on the same processor. Nonadiabatic couplings were computed in both simulations with the overlap derivative method. Thus, the construction of excited state functions in the valence-only approximation leads to  $\sim 50\%$  time saving for computations of nonadiabatic couplings of [8]CPP with the OD method. In the case of determinant derivative methods the time saving would be even more pronounced due to the 5<sup>th</sup> order scaling with the number of occupied orbitals used in the Slater determinant constructions.

Nonadiabatic events between excited states were taken into account by FSSH corrected for decoherence with the SDM method.<sup>51</sup> Since TD-DFTB as a single-reference method suffers from the same problems as

TD-DFT in the description of  $S_1$ - $S_0$  NAC topology,<sup>74</sup> we introduced a condition according to which the internal conversion between  $S_1$  and  $S_0$  states is assumed, and trajectories are stopped, whenever the energy gap between these states drops below 0.15 eV. Newton's equations of motion were integrated using the velocity Verlet algorithm<sup>75</sup> with the time step 0.5 fs. The TDSE was integrated using the fifth order Butcher's algorithm<sup>76</sup> with the time step 0.025 fs, applying interpolated electronic properties between classical steps. The nonadiabatic couplings between excited states were evaluated numerically using the orbital derivative method.<sup>33, 54</sup> The dynamics was propagated for 3 ps.

The excited states occurring during dynamics were analyzed based on their transition density matrices. For this purpose, the molecules are fragmented into phenylene rings. CT, PR, and PRNTO values for the active excited state during the trajectories were computed using the TheoDORE program.

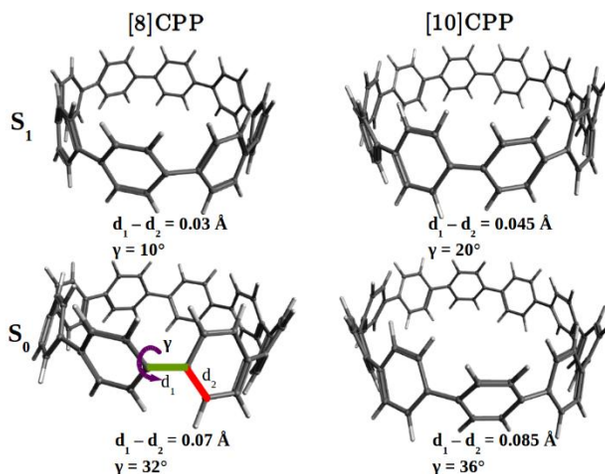
For the comparison with the TD-DFTB results, we also computed vertical excitation energies of both molecules with the long-range corrected TD-LC-DFTB method implemented in the DFTB+ code and with TD-DFT based on three different functionals (PBE,<sup>77</sup> B3LYP,<sup>78-79</sup> and CAM-B3LYP<sup>80</sup>), using the GAMESS-US code.<sup>81-82</sup>

## 4. Results and discussion

### 4.1. Ground and excited-state geometries

The optimized geometries of the ground and the first excited states of [8]CPP and [10]CPP are shown in Figure 2. The molecules are highly symmetric in their ground states, having the symmetries of the  $D_{4d}$  ([8]CPP) and  $D_{5d}$  ([10]CPP) point groups. Their characteristic geometrical features are influenced by three different effects: backbone strain, steric effects, and competition between  $\pi$ -conjugation and quinoidalization.<sup>40</sup>

Their ground states are featured by alternant dihedral angles between subsequent phenylene rings with the values  $\pm 32.1^\circ$  in [8]CPP and  $\pm 35.5^\circ$  in [10]CPP. These values are in excellent agreement with the ones obtained in TD-CAM-B3LYP optimization ( $32^\circ$  and  $36^\circ$ ).<sup>43</sup>



**Figure 2. Structures of [8]CPP and [10]CPP in the ground and the first excited states optimized with (TD-)DFTB.**

The optimized symmetric local minima of the  $S_1$  states are characterized by planarization of phenylene rings with decreased alternant dihedral angles obtaining the values  $\pm 9.9^\circ$  and  $\pm 20.2^\circ$  in [8]CPP and [10]CPP, respectively. Another important distinction between  $S_0$  and  $S_1$  geometries is in the differences between double intra-ring and single inter-ring C-C bond lengths. These differences decrease in the  $S_1$  states in comparison to the ground states—in [8]CPP they are 0.07 in  $S_0$  and 0.032 Å in  $S_1$ ; in [10]CPP they are 0.085 and 0.045 Å. These features indicate that quinoidalization, which is accompanied by ring planarization, is favored in the  $S_1$  states compared to the ground states, where  $\pi$ -conjugation has a larger effect on the structures. Quinoidalization is more pronounced in the smaller molecule because of the influence of larger ring strain, which is reflected in decreased dihedral angles and differences between the lengths of single and double C-C bonds. These findings are consistent with the experimental results obtained with Raman spectroscopy<sup>40</sup> and geometrical observations from TD-CAM-B3LYP optimizations.<sup>43</sup>

#### 4.2. Vertical excitation energies

The vertical excitation energies and oscillator strengths of the first six excited states of [8]CPP and [10]CPP are compared with the corresponding TD-DFT values obtained with the PBE, B3LYP, and CAM-B3LYP functionals in Table 1.

Among the first six excited states at the ground-state geometries, both molecules have two pairs of doubly-degenerate states,  $S_2/S_3$  and  $S_4/S_5$ . One of the pairs, which corresponds to transitions from HOMO to LUMO+1 and LUMO+2, is characterized by large oscillator strengths. The other one, corresponding to transitions from HOMO-1 and HOMO-2 to LUMO, is characterized by small or vanishing oscillator strengths. TD-DFTB, TD-PBE, and TD-B3LYP predict the same state order in [8]CPP, according to which

bright degenerate states are placed above dark ones. TD-CAM-B3LYP and LC-TD-DFTB invert this order, predicting the bright states to have lower energies. In the case of [10]CPP, TD-DFTB and TD-PBE predict intense transitions to degenerate  $S_4$  and  $S_5$  states, whereas the  $S_2$  and  $S_3$  states have low oscillator strengths. This order is inverted with the range-separated functionals and also with the hybrid functional, which predicts that the energies of the close-lying pairs of degenerate states differ only by 0.03 eV.

**Table 1. Computed and experimental vertical excitation energies and oscillator strengths of excited states of [8]CPP and [10]CPP.**

Molecule	TD-DFTB		TD-PBE		TD-B3LYP		TD-CAM-B3LYP		TD-LC-DFTB		Exp.
	$E$ (eV)	$f$	$E$ (eV)	$f$	$E$ (eV)	$f$	$E$ (eV)	$f$	$E$ (eV)	$f$	$E$ (eV)
[8]CPP	2.443	0.000	2.336	0.000	2.879	0.0	3.506	0.000	3.663	0.000	2.64 <sup>39</sup>
	2.842	0.006	2.775	0.000	3.563	0.007	4.177	1.595	4.282	1.152	3.67 <sup>39</sup>
	2.842	0.006	2.775	0.000	3.563	0.007	4.177	1.595	4.282	1.152	
	3.215	0.860	3.214	0.946	3.675	1.322	4.758	0.020	5.002	0.000	
	3.215	0.860	3.214	0.946	3.675	1.322	4.758	0.020	5.002	0.000	
	3.270	0.000	3.273	0.000	3.957	0.000	4.792	0.000	5.138	0.000	
[10]CPP	2.669	0.000	2.496	0.000	3.114	0.000	3.731	0.000	3.949	0.000	
	2.907	0.008	2.758	0.001	3.618	1.415	4.191	2.245	4.355	2.106	3.65 <sup>39</sup>
	2.907	0.008	2.758	0.001	3.618	1.415	4.191	2.245	4.355	2.106	
	3.139	0.855	3.067	0.987	3.647	0.275	4.767	0.000	4.905	0.000	
	3.139	0.855	3.067	0.987	3.647	0.275	4.767	0.000	4.905	0.000	
	3.168	0.000	3.069	0.000	3.869	0.000	4.838	0.000	5.263	0.000	

In comparison to the experimental absorption peak positions, the most reliable values of vertical excitation energies are obtained with the B3LYP functional. The TD-CAM-B3LYP excitation energies overestimate excitation energies by  $\approx 0.5$  eV in comparison to the absorption band peaks. It has already been noticed that range-separated TD-DFT methods tend to blue shift excitation energies of conjugated molecules compared to the experimental absorption peak positions.<sup>43, 83</sup>

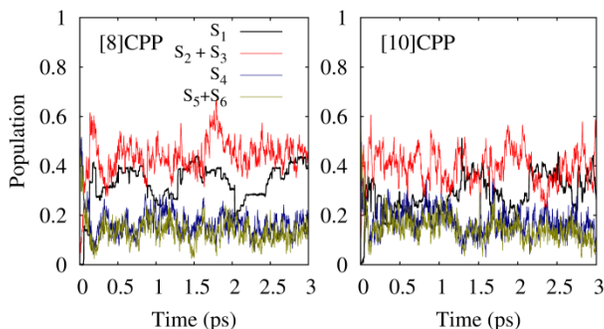
Compared to the experimental position of the tail of absorption spectra of [8]CPP<sup>39</sup> (which arises from the excitation to the  $S_1$  state), the TD-DFTB underestimates the excitation energy by  $\sim 0.2$  eV. The TD-DFTB energies corresponding to the highest oscillator strength transitions to  $S_4$  and  $S_5$  states are underestimated by 0.45 ([8]CPP) and 0.52 eV ([10]CPP), compared to the positions of the band peaks in the experimental absorption spectra. In the case of [8]CPP, TD-DFTB predicts the same order of the excited states as one obtained with TD-B3LYP. In the case of [10]CPP, it inverts the order of the close-lying double-degenerate states. The TD-DFTB energies are consistent with TD-PBE, and the TD-LC-DFTB energies are in line with the corresponding TD-CAM-B3LYP values.

According to the TD-DFTB results, the energy of  $S_4/S_5$  excitation is independent of the system size, and the  $S_0/S_1$  transition is blueshifted when the ring size is increased. These two features agree with the experimental results.<sup>39, 46</sup>

The initial conditions created using a Wigner distribution have broken symmetries compared to the ground state equilibrium geometry. For this reason, transitions to the  $S_6$  state, which are forbidden at the equilibrium geometry, become possible in the ensemble of geometries. The oscillator strength of the  $S_6$  state becomes comparable to the oscillator strengths of the degenerate  $S_4$  and  $S_5$  states. These three states are close-lying, and they can exchange order at different geometries. Thus, we selected initial conditions starting from the  $S_5$  and  $S_6$  adiabatic states including all six states in the dynamics simulations. The algorithm for the initial condition selection in the Newton-X code computes the probabilities for selecting an initial condition based on the oscillator strength for the selected state, which assures that at all selected geometries the  $S_5$  and  $S_6$  states are bright states.

### 4.3. Excited-state population dynamics

After the photoexcitation, the molecules undergo ultrafast internal conversion from the initially populated  $S_6$  and  $S_5$  states via the  $S_4$  state to the lower excited states. This process is shown in Figure 3, which reports the fraction of trajectories in each electronic state in each time step. For both molecules, after this initial period taking about 100 fs, the population remains mostly shared between  $S_1$  (~40%) and  $S_2+S_3$  (~40%). The occupations of  $S_4$  and  $S_5+S_6$  remain at about 10% each until the end of the simulations. The ground state is not reached in any of the trajectories during the 3-ps simulations.



**Figure 3. Fractional occupations of the excited states of [8]CPP (left) and [10]CPP (right) during dynamics.**

In general, the trajectories tend to relax to the  $S_1$ . However, due to the lack of dissipative modes to the environment, once the molecules relax to their  $S_1$  states, a part of the population is transferred back to higher excited states, when they reach regions of strong nonadiabatic couplings between  $S_1$  and  $S_2$ . Furthermore,

the  $S_3$  state can be easily repopulated from the close-lying  $S_2$  state. This recurrent back transfer explains the oscillatory behavior of the average population of these states during the dynamics (Figure 3).

Surface hopping dynamics for  $[n]$ CPPs have been reported by Adamska et al.<sup>43</sup> for several ring sizes  $n$  between 9 and 16. Their simulations were based on configuration interaction singles (CIS) on top of a semi-empirical Hamiltonian, propagated for 500 fs. Like our results, they also found that internal conversion to the  $S_1$  state is reached on the ultrafast scale. However, in their case, once the  $S_1$  is populated, the trajectory remains there, and the  $S_1$  state reaches 100% occupation within 200 fs. The main reason for this difference is that Adamska and co-workers propagated the trajectories using constant-temperature Langevin dynamics with solvation,<sup>7</sup> while we employed a microcanonical ensemble in the gas phase. The nuclear equations of motion in their case include a velocity-dependent friction term, which emulates the kinetic energy transfer to the solvent.

There is still a third set of dynamics simulations for [6]CPP, [8]CPP, and [10]CPP reported in Ref.<sup>44</sup>. These simulations, based on wave-packet propagation for 200 fs, were performed involving four excited states computed on a 26-dimensional model Hamiltonian parameterized at the TD-B3LYP level.

#### 4.4. Decay mechanisms

After the  $S_1$  state is reached during dynamics, three kinetically competitive decay processes from  $S_1$  to the ground state may occur—fluorescence, internal conversion, and intersystem crossing to the  $T_1$  state<sup>41</sup> followed by phosphorescence to the  $S_0$  state. Available experimental and computed quantum yields, rate constants, and lifetimes are summarized in Table 2.

Fujitsuka et al. estimated fluorescence quantum yields, reaction rates, and lifetimes from fluorescence intensity decay experiments<sup>39</sup> and determined triplet lifetimes from phosphorescence decay experiments.<sup>41</sup> Moreover, they estimated quantum yields and rate constants of  $S_1/T_1$  intersystem crossing and  $S_1/S_0$  internal conversion from experiments of triplet quenching with triplet oxygen. Based on the phosphorescence spectra, they confirmed that this reaction produces singlet oxygen, and from the triplet oxygen phosphorescence decay profile they determined the quantum yield of the singlet oxygen generation. They assumed that the quantum yield for singlet oxygen generation is equivalent to the quantum yield for intersystem crossing ( $\Phi_{ISC} = \Phi_{O_2}$ ). The rationale behind this hypothesis was that, in the range of applied triplet oxygen concentrations, the reaction of triplet quenching with oxygen is several orders of magnitude faster than competing phosphorescence process, meaning that the efficiency of triplet quenching is 100%.<sup>41</sup> The quantum yield of internal conversion was then estimated as  $\Phi_{IC} = 1 - \Phi_{ISC} - \Phi_{fl}$ .

From the quantum yields of these three processes, one can conclude that internal conversion ( $\Phi_{IC} = 0.48$ , Table 2) and intersystem crossing ( $\Phi_{ISC} = 0.44$ ) are the main deactivation mechanisms in [8]CPP, whereas fluorescence has negligible quantum yield ( $\Phi_{fl} = 0.08$ ). In [10]CPP, fluorescence becomes the dominant relaxation mechanism with a quantum yield of 0.46, whereas the quantum yield of intersystem crossing is significantly decreased ( $\Phi_{ISC} = 0.17$ ) compared to its value in [8]CPP. Internal conversion remains an important deactivation pathway in [10]CPP, with a quantum yield of 0.37.

**Table 2 Experimental quantum yields ( $\Phi$ ), decay rates ( $k$ ), and lifetimes ( $\tau$ ) of fluorescence (fl), internal conversion (IC), intersystem crossing (ISC), and phosphorescence (ph) of [8]CPP and [10]CPP.**

Molecule	$\Phi_{fl}$	$\Phi_{IC}$	$\Phi_{ISC}$	$k_{fl}$ ( $10^7$ s $^{-1}$ )	$k_{IC}$ ( $10^7$ s $^{-1}$ )	$k_{ISC}$ ( $10^7$ s $^{-1}$ )	$\tau_{fl}$ (ns)	$\tau_{ph}$ ( $\mu$ s)
[8]CPP	0.084 <sup>a</sup>	0.475 <sup>b</sup>	0.440 <sup>b</sup>	0.48 <sup>a, b</sup>	2.7 <sup>c</sup>	2.5 <sup>b</sup>	208.3 <sup>d</sup>	60 <sup>b</sup>
[10]CPP	0.46 <sup>a</sup>	0.368 <sup>b</sup>	0.171 <sup>b</sup>	7.00 <sup>a, b</sup>	5.5 <sup>c</sup>	2.6 <sup>b</sup>	14.3 <sup>d</sup>	58 <sup>b</sup>

<sup>a</sup> Ref. <sup>39</sup> (The absorption spectra of CPPs were recorded in THF.)

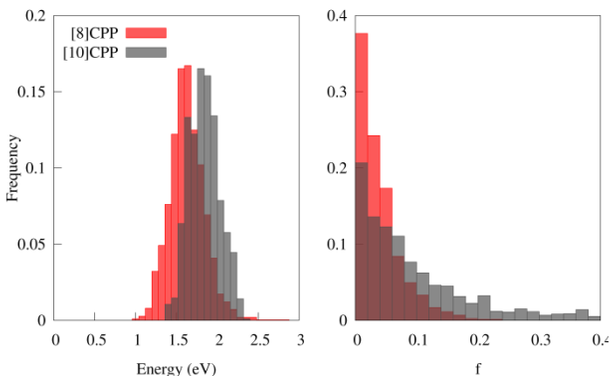
<sup>b</sup> Ref. <sup>41</sup>.  $\Phi_{ISC}$  is assumed to be the quantum yield of singlet oxygen (see text). (The experiments are performed in THF.)

<sup>c</sup> Computed as  $k_{IC} = (k_{fl} + k_{ISC}) / (\Phi_{IC}^{-1} - 1)$  applying the experimental rate constants for fluorescence and intersystem crossing and experimental quantum yield for internal conversion.

<sup>d</sup> Internal fluorescence lifetimes computed as  $\tau_{fl} = 1 / k_{fl}$ .

A recent wave packet dynamics simulation<sup>84</sup> of relaxation process in cycloparaphenylenes within triplet manifold, which included T<sub>3</sub>-T<sub>1</sub> states, revealed the ultrafast internal conversion to the T<sub>1</sub> state. Strong Jahn-Teller interactions occurring between doubly-degenerate T<sub>3</sub> and T<sub>2</sub> states, which induce this ultrafast relaxation, promote indirect intersystem crossing from S<sub>1</sub> to T<sub>1</sub> state (S<sub>1</sub>→T<sub>3</sub>/T<sub>2</sub>→T<sub>1</sub>).

Even though the intersystem crossing and phosphorescence were proven to be important, we do not consider this reaction channel, because our dynamics did not involve triplet states, and we limit the discussion to fluorescence and internal conversion.



**Figure 4. Distributions of the S<sub>1</sub> excitation energies (left) and oscillator strengths (right) of [8]CPP and [10]CPP during dynamics in single trajectories.**

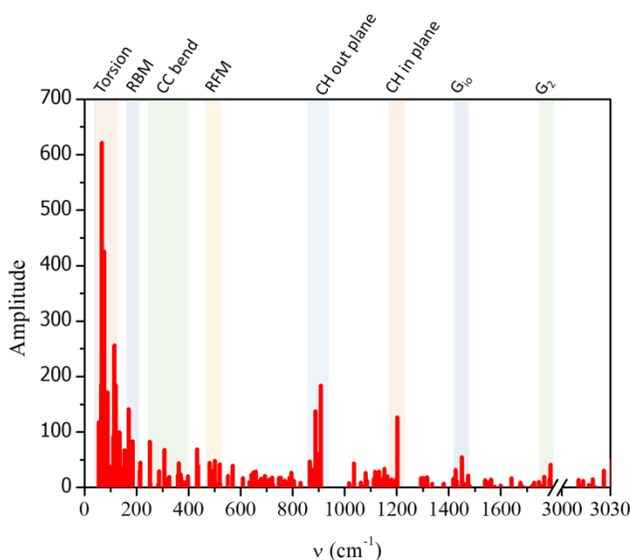
The fluorescence lifetimes are estimated from the measured fluorescence rate constants<sup>39</sup> (as  $\tau_{fl} = 1/k_{fl}$ ). The fluorescence lifetime of [8]CPP (204.3 ns) is ~15 times longer than of [10]CPP (14.3 ns) (Table 2). This shortening with the molecular size can be rationalized based on the dynamics results. The distributions of the excitation energies and oscillator strengths in the  $S_1$  states of both molecules are shown in Figure 4. **Error! Reference source not found.** The  $S_1$  excitation energy ( $E_v$ ) distribution is centered at 1.5 eV in [8]CPP and it increases to 1.8 eV in [10]CPP, indicating the local minima on the  $S_1$  surfaces. The  $S_1 \rightarrow S_0$  oscillator strength ( $f$ ) in [10]CPP is distributed towards higher values compared to the one in [8]CPP. Their mean values also increase with the molecular size, from  $f_{[8]CPP} = 0.04$  to  $f_{[10]CPP} = 0.10$ . These two trends contribute to decreasing the fluorescence lifetime ( $\tau_{fl}$ ) in [10]CPP in comparison to the [8]CPP, because  $\tau_{fl}$  is proportional to  $f^{-1}E_v^{-2}$ .<sup>73</sup> Using the mean values for the excitation energies and oscillator strengths, we may estimate that the ratio between fluorescence lifetimes is 3.6. This value is underestimated compared to the experimental value (~15), but correctly indicates the qualitative trend. The disagreement between these two values could be partially attributed to environmental effects, since the experimental rate constant measurements were done in tetrahydrofuran, whereas the simulations were performed in gas phase. The fluorescence lifetime shortening from [8]CPP to [10]CPP should directly lead to an increase of the fluorescence quantum yield, as it hinders competing processes. The experimental ratio between the fluorescence quantum yields is 5.5 (Table 2).

The increase in the oscillator strength of the  $S_1$  state in [10]CPP in comparison to [8]CPP has been explained by decreasing the energy gap between  $S_1$  and  $S_2$  states, which increases the vibronic coupling between them<sup>44</sup> and by localization of excitations in the  $S_1$  states in larger cycloparaphenylenes, which is accompanied by asymmetrical geometrical distortions leading to an increase in the oscillator strength<sup>43</sup>.

#### 4.5. Vibrational dynamics

We have analyzed the vibrational relaxation using normal mode analysis.<sup>85-86</sup> The geometries at the  $S_2/S_1$  hopping point for all trajectories of [8]CPP were collected, and each of them was then aligned with the [8]CPP  $S_0$  minimum. The alignment of structures is achieved by removing the translational degrees of freedom by coinciding their centers of masses. The structures are then superimposed by rotation within their center of masses systems. The optimal rotation matrix that maximizes the overlap between two geometries is computed applying the Kabsch's algorithm.<sup>87</sup> After being aligned, the Cartesian difference between the  $S_2/S_1$  hopping geometry and the reference geometry was computed and projected on the normal mode basis calculated for the ground state. A map of the active normal modes at the  $S_2/S_1$  hopping point was built by averaging the computed activations of normal modes over all  $S_2/S_1$  hopping points.

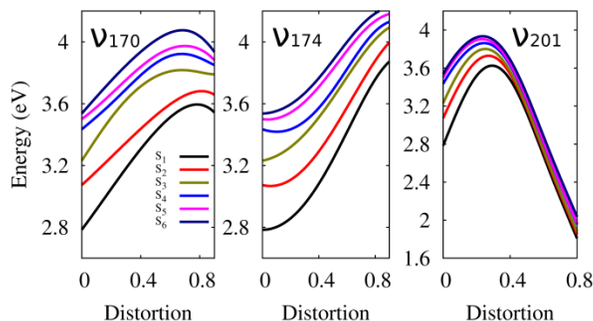
The relative activation of normal modes at the hopping points with respect to the ground-state geometry are shown in Figure 5. They identify the main vibrational motions driving the excited state relaxation. The most active are low-wavenumber torsional modes ( $\nu < 120 \text{ cm}^{-1}$ ) and radial breathing modes (RBM,  $150 < \nu < 220 \text{ cm}^{-1}$ ). Several normal modes in the region between  $230 \text{ cm}^{-1}$  and  $400 \text{ cm}^{-1}$  feature small activation levels. They involve bending of  $C_{ipso}-C_{ipso}$  bonds with respect to the ring planes. The radial flexural modes (RFM,  $\nu \approx 500 \text{ cm}^{-1}$ ) also show small activation levels. They correspond to ring deformations induced by  $C_{ipso}-C_{ipso}$  vibrations outward the ring and  $C_{ortho}-C_{ortho}$  vibrations inward the ring. Modes with the wavenumbers between  $800 \text{ cm}^{-1}$  and  $900 \text{ cm}^{-1}$  have medium activation levels. They are mostly ring deformation modes with out-of-plane C-H bending. The modes with wavenumbers about  $1200 \text{ cm}^{-1}$ , also with medium activation level, are in-plane C-H bending modes. In the region between  $1430$  and  $1790 \text{ cm}^{-1}$ , a series of C-C stretching modes show small activation. They are characterized in more details below.



**Figure 5. Relative activation of different normal modes at the geometry near the  $S_2/S_1$  conical intersection of [8]CPP.**

Cycloparaphenylenes, as well as carbon nanotubes, have three different types of C-C stretching modes:  $G_1$ ,  $G_2$ , and  $G_{10}$  vibrations.<sup>88</sup> The  $G_1$  and  $G_2$  modes correspond to the circumferential stretching of the *ortho-ortho* and *ipso-ipso* C-C bonds. In the  $G_1$  modes all phenylene rings experience identical in-phase motions with the same magnitudes, whereas in the  $G_2$  modes, the adjacent phenylene rings oscillate out of phase. The  $G_2$  vibrations have higher frequencies compared to the  $G_1$ . The  $G_{10}$  modes correspond to *ipso-ortho* C-C bond vibrations and have lower frequencies compared to  $G_1$  and  $G_2$ . The  $G_{10}$  vibrations also have two

components corresponding to in-phase and out-of-phase (we denote them as  $G_{io}$ ) motions in adjacent rings.  $G_{io}$  (1430-1480  $\text{cm}^{-1}$ ) and  $G_2$  ( $\sim 1790$   $\text{cm}^{-1}$ ) modes show small activation levels at the  $S_2/S_1$  crossing.



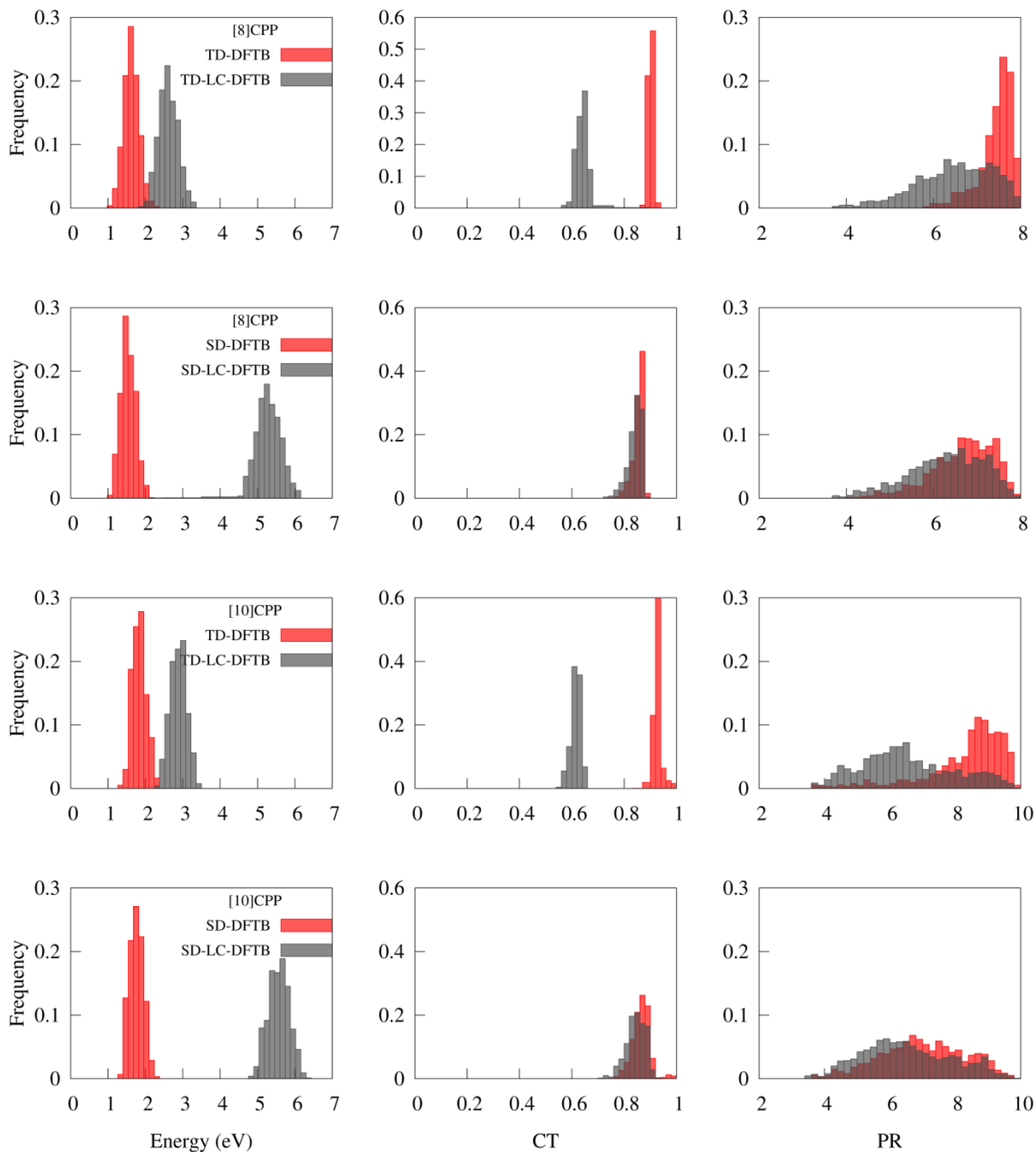
**Figure 6.** Excitation energies of  $S_1 - S_6$  states of [8]CPP at the geometries obtained by distorting the molecules along three C-C stretching modes,  $\nu_{170}$  ( $G_{io}$ , 1432.5  $\text{cm}^{-1}$ ),  $\nu_{174}$  ( $G_{io}$ , 1448.7  $\text{cm}^{-1}$ ), and  $\nu_{201}$  ( $G_2$ , 1786.0  $\text{cm}^{-1}$ ). The x-axis represents a fraction of a normalized normal mode for which the geometry is distorted with respect to the equilibrium geometry.

Rigid excited state potential energy profiles along all  $S_2/S_1$  active normal modes discussed above are computed with TD-DFTB at geometries obtained by elongation along the normal modes. For most of the modes, the excited states remain approximately parallel to each other along the displacement coordinate (see Supporting Information, SI-III). Nevertheless, there are three exceptions: the energy profiles along two  $G_{io}$  modes ( $\nu_{170} = 1432.5$   $\text{cm}^{-1}$  and  $\nu_{174} = 1448.7$   $\text{cm}^{-1}$ ) and the  $G_2$  mode ( $\nu_{201} = 1786.0$   $\text{cm}^{-1}$ ). These three C-C stretching modes induce a decrease in the  $S_2-S_1$  gap (see Figure 6). For example, the geometrical distortion along mode  $\nu_{174}$  for  $0.5\nu_{174}$  with respect to the equilibrium geometry (where  $\nu_{174}$  represents the normalized normal mode) reduces the  $S_2-S_1$  gap from 0.3 to 0.15 eV. The small  $S_2-S_1$  gaps indicate the vicinity of a conical intersection between these two states, which could be obtained by relaxing the other coordinates along the stretching modes. The activation of  $\nu_{170}$  and  $\nu_{174}$  also induces  $S_3/S_4$  and  $S_5/S_6$  crossings. These series of state crossings induced by  $G_{io}$  and  $G_2$  imply that these modes play a central role for the relaxation of the excited states down to  $S_1$ , and for the eventual repopulation of the higher excited states.

#### 4.6. Charge transfer and exciton (de)localization in the $S_1$ state

The excitation energies, charge transfer (CT) numbers, and participation ratio (PR) computed with TD-DFTB along a single trajectory of [8]CPP and [10]CPP, considering only time steps when the molecules are in the  $S_1$  states, are shown in Figure 7. To verify how reliable TD-DFTB is in the description of energies, charge transfer, and localization properties of the excited states

during the dynamics, we recomputed the excited states, CT, and PR numbers with TD-LC-DFTB at the same geometries. These values are also plotted in Figure 7 **Error! Reference source not found.**.



**Figure 7. Distributions of the excitation energies (left), charge transfer numbers (middle), and participation ratio numbers (right) in the  $S_1$  state of [8]CPP (top of the figure) and [10]CPP (bottom of the figure), during dynamics of a single trajectory considering the TD-(LC)-DFTB and SD-(LC)-DFTB methods.**

As expected from our analysis of the vertical excitation energies, the distributions of the excitation energies show significant differences between  $S_1$  energies computed with the two methods. The TD-DFTB energies are red shifted in comparison to the TD-LC-DFTB energies by  $\sim 1$  eV. Nevertheless, not all disagreement between the results is caused by TD-DFTB inaccuracy. We have already discussed that range-separated functionals tend to blue shift the excitation energies of CPPs with respect to the experimental absorption peak for approximately 0.5 eV, whereas the TD-DFTB tends to red shift these energies for similar magnitude (see discussion of Table 1).

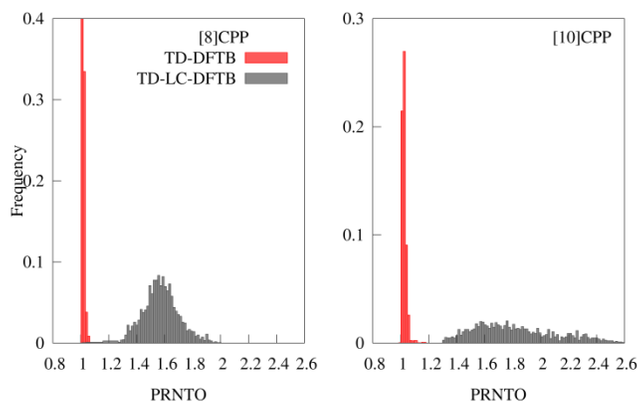
Moreover, TD-DFTB overestimates CT numbers of the  $S_1$  state of both molecules in comparison with the CT numbers obtained from TD-LC-DFTB computations. According to the TD-LC-DFTB, the  $S_1$  states are obtained by mixing of Frenkel and charge transfer excitons, with the CT distributions peaked at  $\sim 0.65$ . On the other hand, TD-DFTB predicts almost pure CT character of the  $S_1$  states with CT distributions peaked at  $\sim 0.9$  in both molecules. This indicates that TD-DFTB, similarly as TD-DFT at the level of GGA functionals, overestimates charge-transfer character of excited states. It has been confirmed, for instance, on a set of conjugated systems<sup>89</sup> that TD-DFT with the GGA-level functionals predicts spurious charge transfer character for the lowest excited state whereas predominantly local character is obtained at the ADC(2) level and with TD-DFT with hybrid and long-range corrected functionals.

The PR distributions calculated with TD-DFTB indicate that the  $S_1$  states of both molecules are delocalized over 8 fragments in [8]CPP and  $\sim 9$  fragments in [10]CPP (Figure 7). The inclusion of long-range correction to the functional leads to some localization of excitons in comparison to the TD-DFTB prediction, with the  $S_1$  state of [8]CPP mostly delocalized over 6 to 8 fragments and that of [10]CPP showing a broad distribution between 4 to 9 fragments and peaking at 6. (It has been already observed that the localization of excitons can be adequately described only with range-corrected hybrid DFT models.<sup>90-91</sup>)

According to the TD-LC-DFTB, the broad PR distribution for [10]CPP implies that in this molecule both localized and delocalized excitons occur in the first excited state. The maximum of the PR distribution peaked at 6 fragments indicates a more pronounced exciton localization than in the case of the smaller molecule. This result is in agreement with the dynamics results of Adamska et al.,<sup>43</sup> who noticed that localization of excitons, absent in [9]CPP occurs in larger CPPs ( $n \geq 12$ ), where they are localized on 5 to 7 rings. This exciton localization is attributed to its self-trapping induced by geometrical distortions. In addition, Nishihara et al.<sup>38</sup> observed, based on the temperature dependence of the fluorescence lifetime, that the excitons of larger CPPs ( $n = 12$ ) are localized at temperatures smaller than the room temperature and delocalized at higher temperatures.

It is worth mentioning here that, even though, based on energy criterium, the B3LYP functional provides the most reliable description on the excited states of cycloparaphenylenes, it does not correctly describe the delocalization properties of the exciton in the  $S_1$  state, which is an important feature of dynamics. We have optimized the geometries corresponding to the localized exciton in [10]CPP starting from several different geometries taken from dynamics at which the system was in the  $S_1$  state and the exciton was localized. The optimizations were performed with TD-B3LYP and TD-CAMB3LYP using 6-31G basis set and with a small optimization radius. TD-CAMB3LYP yielded non-symmetric structure and localized  $S_1$  state (PR~7), whereas TD-B3LYP always converged to the completely symmetrical structure and completely delocalized  $S_1$  state (PR~10). This finding indicates that TD-B3LYP predicts that localization of exciton in the  $S_1$  state is not the most favorable energetically, and probably it would be shown in dynamics performed based on TD-B3LYP energies and gradients. Most probably in the TD-B3LYP dynamics the system would converge to the symmetrical structures and delocalized excitons in the  $S_1$  state.

We wanted to verify how the single-determinant (SD) approximation of the excited-state wave functions<sup>10</sup> affects the description of their localization and charge transfer properties. We represented the  $S_1$  states in the form of single determinants selecting the one with the largest CI-coefficients obtained either with TD-DFTB (rendering SD-DFTB) or TD-LC-DFTB (rendering SD-LC-DFTB). The excitation energies in the SD approximation correspond to simple KS orbital energy gaps. CT and PR were computed with the renormalized SD wave functions. The three quantities for [8]CPP and [10]CPP are shown in Figure 7**Error! Reference source not found.** In both cases, the energies of the  $S_1$  state at the SD-DFTB approximation are very close to the TD-DFTB values, whereas the SD-LC-DFTB excitation energies are overestimated by approximately 3 eV in comparison to the LC-TD-DFTB values. CT and PR are both insensitive to range separation, and SD-DFTB and SD-LC-DFTB provide very similar distributions. In the SD approximation, CT has a sharp distribution peaked at ~0.85. (A result like that obtained with TD-DFTB). The results for PR, however, are somewhat surprising. For both molecules, PR distributions obtained with both SD approximations are in reasonable agreement with that from TD-LC-DFTB, our highest level.



**Figure 8. Distributions of the PRNTO of the  $S_1$  states reached during dynamics in single trajectories of [8]CPP (left) and [10]CPP (right).**

Some insights into the differences between compositions of the TD-DFTB and TD-LC-DFTB wave functions that induce distinct charge-transfer and exciton localization properties can be gained by analyzing their PRNTO distributions (Figure 8 **Error! Reference source not found.**). The PRNTO distributions show that the TD-DFTB wave functions, built via Eq. (13), are represented mainly by a single transition, whereas the TD-LC-DFTB wave functions are represented by more than one important transition between electron/hole NTO pairs. Thus, it is clear that the LC correction not only changes the orbitals (via the action of the HF exchange) but also changes the distribution of determinants contributing to the state description. Based on this constatation, we are in a position to understand the relations between  $\Delta E$ , CT, and PR with the different levels.

TD-DFTB and SD-DFTB do not show a significant difference in the excitation energies because the TD-DFTB wave functions are dominated by one transition. In addition, the two-electron integrals in Eq. 10 (that are included in TD-DFTB but not SD-DFTB) are usually small compared to the HOMO-LUMO gap for local/semilocal functionals. On the contrary, when LC corrections are considered, the SD approximation is an inadequate representation of the TD-LC-DFTB result, composed of many determinants. For functionals involving a fraction of Hartree-Fock exchange the mentioned two-electron integrals are typically large. These two effects lead to the strong deviation in the excitation energy between the two levels.

The CT distributions computed based on TD-DFTB and SD-DFTB both predict almost pure charge transfer character of the  $S_1$  states, implying that the highest-weight transitions between electron/hole NTO pairs are the charge-transfer transitions. The inclusion of the LC correction in TD-LC-DFTB reduces the CT value through the multideterminantal contribution to the density. Nevertheless, the LC correction in SD-DFTB

does not change the CT values in comparison to SD-DFTB because, by construction, only one determinant is used.

The situation of PR is more subtle than that of  $\Delta E$  and CT. Although TD-DFTB state is dominated by a single determinant, other determinants still contribute to the final result. And these secondary contributions are responsible for donating the delocalized character to the state, reflected on the large PR. In SD-DFTB, however, only the main determinant is considered, resulting in more localized densities and smaller PR. The inclusion of LC correction in TD-LC-DFTB leads to the same effect—PR reduction—but due to a different reason, as now the reduced PR is obtained as result of a multideterminantal composition. As far as we can see, the good agreement of PR computed with TD-LC-DFTB and SD-DFTB was just coincidence. If for instance, the multideterminantal TD-LC-DFTB results were delocalized, the SD-DFTB would still regardless predict a small PR. Once more, SD-LC-DFTB and SD-DFTB show similar PR values because only a single determinant is analyzed.

## 5. Conclusions

In the present study, we implemented the decoherence-corrected fewest switches surface hopping based on LR TD-DFTB, enhanced by transition density analysis through interfaces between Newton-X, DFTB+, and TheoDORÉ programs. This implementation was applied to study the excited states and relaxation dynamics of two cycloparaphenylenes, [8]CPP and [10]CPP.

The vertical excitation energies of [8]CPP and [10]CPP computed with the TD-DFTB and LC-TD-DFTB were compared with the TD-DFT energies obtained with the PBE, B3LYP, and CAM-B3LYP functionals. TD-DFTB predicts the same ordering of the states as TD-B3LYP for [8]CPP, which gives the closest excitation energies to the experimental values. For [10]CPP, the order of the close-lying pairs of degenerate states obtained with TD-DFTB does not correspond to the one predicted with TD-B3LYP. From the comparison of energies, we can conclude that TD-DFTB systematically underestimates the excitation energies by approximately 0.5 eV, which is a typical feature of DFT methods at the GGA functional level. However, it describes two important features of the absorption spectra well: the most intense peaks are independent of system size, and the  $S_0$ - $S_1$  energy gap is blue-shifted with the increase of system size.

Based on the results of the TD-DFTB surface hopping dynamics and normal mode analysis, we can draw the following conclusions:

- (1) Both molecules relax to  $S_1$  within 100 fs. Up to 3 ps, there is no indication of further relaxation to  $S_0$ . In the gas phase, repopulation of higher excited states, especially  $S_2$  and  $S_3$ , does not allow for full population transfer to  $S_1$ ;
- (2) The experimentally observed trend of increasing the fluorescence rate constant in CPP[10] in comparison to CPP[8] is reproduced by the TD-DFTB dynamics. It originates from a simultaneous increase of the  $S_0$ - $S_1$  energy gap and oscillator strength in [10]CPP in comparison to its values in [8]CPP;
- (3) The vibrational dynamics driving the relaxation in the manifold of excited states is enabled by low-amplitude activation of C-C stretching modes, which induces a series of state crossings. These same modes may be responsible for the back transfer in the gas phase.

By performing the analysis of the  $S_1$  states based on their transition density matrices, we concluded that the correct description of the charge-transfer and localization properties of the  $S_1$  states can be achieved only with long-range corrected (LC) TD-DFTB. According to TD-LC-DFTB, the  $S_1$  states are mixtures of the Frenkel and charge-transfer excitons, with the CT numbers peaked at  $\sim 0.65$ . The TD-DFTB method tends to overestimate their charge transfer character, predicting the  $S_1$  states to have almost pure charge transfer character. According to the TD-LC-DFTB transition densities, the  $S_1$  state is delocalized in the smaller molecule, whereas it is localized in the larger one. The feature of density localization in larger [ $n$ ]CPPs ( $n \geq 10$ ), which has been earlier observed from the results of dynamics simulation<sup>43</sup> and also experimentally, based on the temperature dependence of fluorescence lifetime<sup>38</sup> (for  $n = 12$ ), could not be reproduced from the TD-DFTB transition densities.

We have also tested the performance of single-determinant (SD) approximations of the wave function with DFTB and LC-DFTB. The results show that SD-DFTB provides a description of the excited states with quality like that of TD-DFTB. SD-LC-DFTB, however, deteriorates the excitation energies. The density localization is well described in both cases. Nevertheless, because this localization is based on a single-determinant representation, it may fail in cases where the actual density is delocalized due to multideterminantal effects.

## Associated content

### Supporting information

Cartesian coordinates, simulated spectrum, potential energy curves. This information is available free of charge via the Internet at <https://pubs.acs.org>.

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### Notes

The authors declare no competing financial interest.

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