

Conjugated organic radicals and polyradicals: electronic structure and photophysics

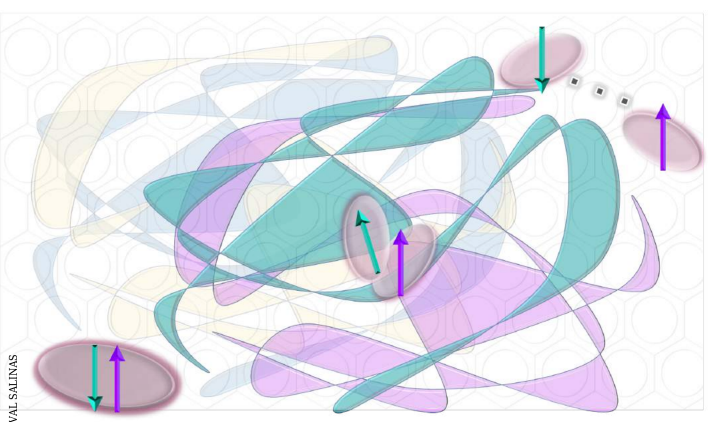
María Eugenia Sandoval-Salinas

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CONJUGATED ORGANIC DIRADICALS AND POLYRADICALS: ELECTRONIC STRUCTURE AND PHOTOPHYSICS



María Eugenia Sandoval-Salinas







Universitat de Barcelona Facultat de Química

Programa de Doctorat en Química Teòrica I Modelització Computacional

Conjugated organic radicals and polyradicals: electronic structure and photophysics

Presents:

María Eugenja Sandoval-Salinas

PhD supervisor:

Dr. David Casanova Casas

PhD tutor:

Prof. Pedro Alemany I Cahner

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Abstract

The principal aim of this thesis is the understanding of the electronic structure of organic diradical and polyradical molecules. Unveiling the properties that give them the electronic, magnetic, and optical properties to be applied as main components in optoelectronic devices. Specifically, the objectives that have been achieved are i) the characterization of organic compounds with diradical and polyradical character, and their electronic, magnetic, and spectroscopic properties; ii) the detailed description of the singlet fission mechanism, as well as the proposal of a new system that, theoretically, is able of carrying out efficiently the singlet fission process; and iii) the use of quantum mechanical methods (specifically RAS-SF) and computational tools to get a proper description of the electronic structure of the ground state of systems in which non-dynamic correlation plays an important role.

In the first place, organic compounds whose optical and magnetic properties make them interesting in the field of optoelectronic materials were studied. The relationship between molecular structure and the radical character was found by the study of linear and cyclic acenes and small triangular fragments derivatives of graphene. While the diradical and tetraradical character increase together with the size of the linear and cyclic compounds, the triangular structures (TGNF, the acronym for Triangulene Graphene Nano Fragments) are open-shell systems with high-spin ground state multiplicity. Furthermore, a manner to tune the spin multiplicity in the ground state of TGNFs is proposed through heteroatom doping or hydrogenation, which offers a way to design larger graphene nanofragments with well-defined spin-multiplicity. Towards larger systems, the increment of the size is associated with the increase of the polyradical character. This thesis presents the rationalization of the electronic structure of organic macrocycles with high polyradical character. Concretely, from triradicals to decaradicaloids (up to 10 radical centers). The properties triggered by the open-shell character of the ground state are as diverse as surprising. For instance, AWA systems (annulenes-within-annulenes) have been characterized in collaboration with experimental groups for the first time. The global aromaticity exhibit by these macrocycles responds to the radical interaction in each of the annulenes and is governed by both aromaticity rules, Hückel's, and Baird's simultaneously.

On the other hand, the singlet fission process (SF) was expanded from the clas-

sical model, which involves five electronic states, to a model that includes double excitations (D states), a seven-state model. Using a simple model, it is estimated that the D states can play an active role in SF, as well as the necessary conditions to maximize their participation as an initial or intermediate state in the process. The feasibility of spiro systems carrying out SF is exposed.

Dedication

To...



Acknowledgements

I want to thank...



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Chapter 1

How to (and why) read this thesis

It is not my intention to imitate Cortázar's masterpiece "Rayuela" (Hopscotch) providing a table of instructions to read this thesis. However, while there is not a hidden mystery in the organization of its chapters, it seems appropriated to introduce them in a naturally evolutionary way, exposing the objectives pursued at each step.

The main aim of the research presented in this thesis consists in the understanding of the electronic structure of organic diradical and polyradical molecules.

Through the study of organic systems with different radical natures, this work also intends to help in the design of new materials with improved tunable magnetic properties for applications in the wide field of organic electronics.

From a more personal perspective, the final aim of my Ph.D. in the group of Dr. Casanova is twofold, namely acquiring expertise in the study of electronic structure properties of complex systems and developing competences in the use of appropriate quantum chemistry methods and computational tools for the rationalization of experimental measurements

In Chapter 2, I provide the general background theory and concepts about diradicals, and how to characterize them from a theoretical point of view. Following that, I present in Chapter 3 a very broad overview of the theoretical methods and computational tools used to achieve the specific objectives set for the study of the systems considered in this thesis.

In results Chapters 4-8, the diradical character of the organic systems investigated is described and discussed from a structure-electronic properties relationship perspective in the context of their potential applications. While the systems considered purposely differ in terms of topology and associated electronic/magnetic features, all chapters share common objectives:

- Determine the diradical (polyradical) character of the ground state of the systems under study.
- Identify the structure-diradical character relationships.

- Propose an interpretation of the experimental observations (when experimental data is available) at the electronic and molecular levels.
- Assess the robustness and quality of computational methods.

Among the zoo of existing graphene nanostructures, phenalenyl, triangulene and acenes are well-known representative examples of small organic molecules with open-shell character. In Chapter 4, the modulation of their magnetic properties as a function of structural and chemical modifications, namely size, shape, heteroatom substitution, and connectivity, is theoretically investigated. Particular objectives of this chapter are:

- Compare the open-shell nature of the ground state in hydrocarbons with different structure.
- Determine the role that topology plays in the nature of their radical character.
- Identify the origin of the synthetic challenge they represent.
- Explore a way to modulate the radical character in small triangular graphene nanostructures.

Results included in this chapter are published in:

- M. E. Sandoval-Salinas, A. Carreras, D. Casanova, "Triangular Graphene Nanofragments: open-shell character and doping", *Phys. Chem. Chem. Phys.*, 2019, 21, 9069-9076.
- A. Perez-Guardiola⁺, M. E. Sandoval-Salinas⁺, D. Casanova, E. San Fabián,
 A. J. José Pérez-Jiménez, J. C. Sancho-Garcia, "The Role of Topology in Organic Molecules: Origin and Comparison of the Radical Character in Linear and Cyclic Oligoacenes and Related Oligomers", Phys. Chem. Chem. Phys., 2018, 20, 7112-7124.

In Chapter 5, the rise of diradical character by means of reduction/oxidation of a family of closed-shell oligomers was studied, as well as the modulation of diradical character of a poly-p-phenylene via external stimuli. In both cases, experimental data was available and all the calculations have the final purpose to explain the spectroscopic features triggered by the change in the diradical character of the systems, and particularly:

- Describe the electronic structure of the ground state of organic oligomers to explain the origin of the magnetic properties determined experimentally.
- Interpret the spectroscopic features in light of electronic structure calculations of the electronic excited states.

• Propose an explanation of the relative stability between conformers.

Results included in this chapter are published in:

- D. Yuan, S. Medina, P. Mayorga, L. Ren, M. E. Sandoval-Salinas, S. J. Grabowski, D. Casanova, X. Zhu, J. Casado, "Thieno[3,4-c]pyrrole-4,6-dione Oligothiophenes Have Two Crossed Paths for Electron Delocalization", *Chem. Eur. J.*, 2018, 24, 13523-13534.
- S. Medina, P. Mayorga, M. E. Sandoval-Salinas, T. Li, F. J. Ramírez,
 D. Casanova, X. Wang and J. Casado, "Isomerism, Diradical Signature, and
 Raman Spectroscopy: Underlying Connections in Diamino Oligophenyl Dications", ChemPhysChem 2018, 19, 1465.

Beyond diradical (or tetraradical) character, systems with a large number of radical centers are particularly appealing candidates for use in spintronic devices and molecular magnets. In Chapter 6, series of macrostructures with high polyradical character are studied in collaboration with experimental groups. The objectives in this chapter are:

- Describe the radical-radical interaction in macrostructures.
- Identify the origin of their inherent features at the electronic level.
- Prove the performance of common methods in the description of complex (big and largely correlated) systems.

Results included in this chapter are published in:

- Y. Ni, M. E. Sandoval-Salinas, T. Tanaka, H. Phan, T. S. Herng, T. Y. Gopalakrishna, J. Ding, A. Osuka, D. Casanova, J. Wu, "[n]Cyclo-parabiphenylmethine polyradicaloids: [n]annulene analogues and unusual valence tautomerization", *Chem*, 2019, 5, 108-121.
- C. Liu, M. E. Sandoval-Salinas, Y. Hong, T. Y. Gopalakrishna, H. Pan, N. Aratani, T. S. Herng, J. Ding, H. Yamada, D. Kim, D. Casanova, J. Wu, "Macrocyclic Polyradicaloids with Unusual Super-ring Structure and Global Aromaticity", Chem, 2018, 4, 1-10.
- J. Wang, G. Kim, M. E. Sandoval-Salinas, H. Phan, T. Y. Gopalakrishna, X. Lu, D. Casanova, D. Kim, J. Wu, "Stable 2D anti-ferromagnetically coupled fluorenyl radical dendrons", *Chem. Sci.*, 2018, 9, 3395-3400.

The first three results chapters focused on the description of the radical character of molecular systems in relation to their shape, size and other structural and chemical features. In order to go further, the two final chapters address a concrete photophysical process, namely singlet fission, through the study of a spiro-conjugated system able to hold radical character. Notably, the singlet fission process has been extensively investigated as an alternative to increase the efficiency of solar cells. In Chapter 7, a revision of the fundamental aspects of singlet fission theory is presented, as well as an extension of the widely accepted five-(electronic)state model by inclusion of doubly excited states.

- Understand the principles of singlet fission mechanism.
- Establish the structural and energetic conditions that facilitate doubly excited state to take part in the singlet fission mechanism.
- Estimate the impact of the doubly excitations in the singlet fission processes.

Results included in this chapter are published in:

• M. E. Sandoval-Salinas, D. Casanova, "The doubly excited state in singlet fission", *ChemPhotoChem*, accepted, DOI: 10.1002/cptc.202000211.

Finally, Chapter 8 presents the in-depth investigation of spiro-conjugated systems within the context of singlet fission. The specific objectives of this chapter are:

- Describe the electronic excited states of pro-spiro chromophores.
- Examine the effect of the spiroconjugation on the excited states presence and ordering.
- Evaluate the suitability of studied systems to carry out singlet fission.
- Understand the electronic structure of the nature of the spiro conjugation.
- Propose the singlet fission mechanism that spiro systems can undertake.
- Identify the effect of the structural deformations triggered by representative normal mode vibrations.
- Explore the suitability of spiro-systems to carry out ISC.
- Analyse the effect of the spiro conjugation on the radical character of the spiroconjugated system.

Results included in this chapter are published in:

• M. E. Sandoval-Salinas, A. Carreras, J. Casado, D. Casanova, "Singlet fission in spiroconjugated dimers", *J. Chem. Phys.*, 2019, 150, 204306.

- M LV⁺, Y. Yu⁺, M. E. Sandoval-Salinas⁺, J. Xu, Z. Lie, D. Casanova, Y. Yang, J. Chen, "Engineering the charge-transfer state to facilitate spin-orbit charge transfer intersystem crossing in spirobis[anthracene]diones. *Angew. Chem. Int. Ed.*, 2020.
- S. Rivero⁺, R. Shang⁺, **M. E. Sandoval-Salinas**⁺, H. Hamada, K. Nakabayashi, S. Ohkoshi. D. Casanova, E. Nakamura, J. Casado, "A spiro-driven tetraradical", submitted.

At the end of this dissertation, the general conclusions of this work are summarized.

CHAPTER 1.	HOW TO (AND WHY) READ THIS THESIS

Chapter 2

Introduction

2.1 Diradicals

In the Chemistry world, the terms radical and diradical are commonly associated with the idea of instability and high reactivity. Nevertheless, they have been extensively explored for more than one century and synthetic routes, stabilization techniques and characterization methods are nowadays well known. Organic diradicals are promising systems to be applied as main components in optoelectronic, photovoltaic, and magnetic devices. (1; 2; 3; 4)

2.1.1 What is a radical? and a diradical?

A radical or free radical is defined as a chemical entity with one unpaired electron. Depending on the atom that holds the unpaired electron, radicals are named as carbon-, nitrogen-, oxygen- or metal-centered. Also, they can be classified as σ - or π -radicals if the singly occupied orbital has considerable σ or π character.(???) The formation of radicals is usually associated with reaction intermediates or unstable products of bond cleavage. Due to the presence of the unpaired electron, radicals are highly reactive species with a very short lifetime. The understanding, detection, and characterization of radicals are crucial in organic synthesis. For instance, polymerization reactions are carried out mostly via radicals. They also play a key role in (several) biological processes such as the cellular oxidation, among others.

The first successful isolation of a radical molecule was achieved more than 100 years ago when Gomberg reported the observation of the triphenylmethyl radical (TM, Figure 2.1).(5; 6; 7) In this molecule, three phenyl moieties are connected through one sp^2 carbon (radical). Despite the sterical protection provided by the phenyl groups to the radical center, TM is a highly reactive species. Thenceforth, constant modifications of the TM backbone have been performed to improve its stability and potentiate its practical applications. Gomberg's pioneering work paved

the way for an extensive field of research that is still being explored nowadays.

Figure 2.1: Structure of the triphenylmethyl (TM) radical.

However, even before the observation of the monoradicals, the high reactivity in some alternated compounds with even number of electrons has been explained considering the idea of two unpaired electrons. (8??) This kind of systems, where two radicals interact across the molecular scaffold, are better known as diradicals or biradicals. (???) Although both terms have been used indistinctly through the years, they are not equivalent: while in a diradical there is a measurable interaction between unpaired electrons that determines the spin multiplicity of their ground state (singlet or triplet), there is no interaction in a biradical and unpaired electrons act as two independent doublets.(2) In this thesis, I focus my interest on diradicals. They are described as molecules with two unpaired electrons occupying two degenerate orbitals, (9; 2; 10) and are extremely important in the understanding of chemical reactivity, the nature of chemical bonding, or in biological processes.

2.1.2 Classes of diradicals

In the literature, several ways exist to classify diradicals, all of them having some points in common. Hereafter, I follow the classification used by M. Abe.(2)

2.1.2.1 Localized and delocalized diradicals

Intuitively, localized diradicals are those in which the unpaired electrons can be found over specific atoms. They can be generated by homolytic bond-cleavage and then change their spin multiplicity (singlet \rightleftharpoons triplet) by intersystem crossing. Localized triplet diradicals have long lifetimes(2) and their chemistry has been more extensively investigated than the one of singlet localized diradicals. In Figure 2.2a some examples of localized diradicals are shown.

On the contrary, delocalized diradicals are part of a π -conjugated system and the unpaired electrons cannot be associated to a particular atom (Figure 2.2b). These are classified as triplet antiaromatic or Kekulé and non-Kekulé diradicals.

• Antiaromatic delocalized diradicals:

 $4n\pi$ -annulenes are antiaromatic in their ground state according to the Hückel's rule but aromatic in their first excited triplet state, as explained by Baird in

1972(?). This aromaticity allows the delocalization of the two electrons of the triplet (Figure 2.2e).

• Kekulé delocalized diradicals:

Kekulé delocalized diradicals are those presenting two resonance structures: aromatic and quinoidal (Figure 2.2c). While the Lewis structure of the aromatic form has to be drawn with two unpaired electrons (spin-triplet ground state), the bond alternation in the quinoidal form permits all electrons to be paired in a "closed-shell" (CS) configuration (spin-singlet ground state). Normally, the ground state holds a singlet spin multiplicity, but the preferred formation of aromatic rings brings about the presence of unpaired electrons. The typical interconversion energy between quinoidal-aromatic (singlet-triplet) conformers is $\Delta E_{ST} = 8-30$ kcal/mol, making these compounds quite reactive even when they possess a singlet ground state.

• Non-Kekulé delocalized diradicals:

In these systems, no resonance structures without unpaired electrons can be drawn (Figure 2.2d). More specifically, in a localized orbital representation, if both frontier orbitals can be represented over the same group of atoms, they are termed as non-disjoint non-Kekulé diradicals and the electron repulsion between both electrons favors the triplet state over the singlet. If there is no shared atom in the radical orbitals, they are disjoint non-Kekulé diradicals.

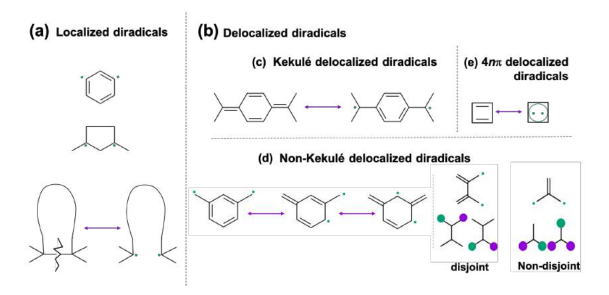


Figure 2.2: Classes of diradicals. Localized (a), delocalized (b), Kekulé delocalized (c), non-Kekulé delocalized, and disjoint and non-disjoint delocalized (d) diradicals.

2.1.3 Electronic states of diradicals

To properly describe the electronic structure of diradicals and rationalize their properties, it is necessary recall their definition: "two electrons occupying two degenerated orbitals". In a model of two electrons in two degenerate orbitals based on the previous definition, it is possible to form six electronic configurations (Figure 2.3), which can be used to generate the three microstates (M_s) of one triplet and three singlet states.

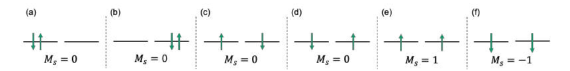


Figure 2.3: Electronic configurations and the projection of the total spin, M_s , that can be obtained by the distribution of two electrons in two degenerate orbitals.

Configurations (a) and (b) of Figure 2.3 are two closed-shell singlets; (e) and (f) are the $M_s = 1$ and $M_S = -1$ components of the triplet, respectively; and the linear combination of (c) and (d) gives rise to the $M_s = 0$ spin projection of the triplet and the open-shell singlet.

From these determinants, 6 wave functions are written, corresponding to singlet (Ψ_n^s) and triplet (Ψ_n^t) states (n=1-3) as:

$$\Psi_1^s = \frac{1}{2} [\phi_1^2 - \phi_2^2] (\alpha \beta - \beta \alpha), \qquad (2.1)$$

$$\Psi_2^s = \frac{1}{2} [\phi_1^2 + \phi_2^2] (\alpha \beta - \beta \alpha), \qquad (2.2)$$

$$\Psi_3^s = \frac{1}{2} (\phi_1 \phi_2 + \phi_2 \phi_1) (\alpha \beta - \beta \alpha),$$
(2.3)

$$\Psi_1^t = \frac{1}{2} (\phi_1 \phi_2 - \phi_2 \phi_1) (\alpha \beta + \beta \alpha), \qquad (2.4)$$

$$\Psi_2^t = \frac{1}{\sqrt{2}} (\phi_1 \phi_2 - \phi_2 \phi_1)(\alpha \alpha), \tag{2.5}$$

$$\Psi_3^t = \frac{1}{\sqrt{2}} (\phi_1 \phi_2 - \phi_2 \phi_1)(\beta \beta), \qquad (2.6)$$

where $\phi_i \phi_j$ is the spatial part of the wave function of electrons 1 and 2 in the *i* and *j* orbitals and α/β the spin of each particle. Equation 2.1 and 2.2 are the "closed-shell" singlets, in which the wave functions are written as the contribution of both (a) and (b) determinants.

The description of the ground state of diradical systems is given by the energy relationship between these singlet and triplet states. The energy of the Ψ_3^s (herein named as open-shell singlet) can be expressed as:

$$E_{os} = h_1 + h_2 + J_{12} + K_{12}, (2.7)$$

where h_i is the monoelectronic energy in the orbital i, J_{ij} is the Coulomb integral $(J_{ij} = \int \int \phi_i^2(1) \frac{e^2}{r_{12}} \phi_j^2(2) dr_1 dr_2)$, that counts for the repulsion between electrons 1 and 2 localized in the same orbital, and K_{ij} is the exchange integral, $K_{ij} = \int \int \phi_i(1)\phi_j(1) \frac{e^2}{r_{12}}\phi_i(2)\phi_j(2) dr_1 dr_2$.

The energy of the triplet, expressed in the same terms, is:

$$E_t = h_1 + h_2 + J_{12} - K_{12}. (2.8)$$

Notably, the only difference between Equation 2.7 and 2.8 is the sign of the exchange integral K_{12} . Due to the symmetry (antisymmetry) of the spacial wave function of the singlet (triplet), the probability of having two electrons together rises (vanishes) and the electronic energy increases (decreases) by K_{12} , which is always a positive term. Accordingly, the energy gap between the two lowest states of a diradical is dictated by:

$$E_{os} - E_t = 2K_{12}, (2.9)$$

where the energy of the triplet is lower than the OS by $2K_{12}$. This is in good agreement with the Pauli exclusion principle that prevents two electrons with the same spin from getting close and lowers their repulsion eventually resulting in the Hund's rule favoring the triplet stabilization.

On the other hand, the energy of the closed-shell states (Equation 2.1 and 2.2) is expressed as:

$$E_{cs_{\pm}} = h_1 + h_2 + \frac{J_{11} + J_{22}}{2} \pm K_{12}, \tag{2.10}$$

in which the energy splitting between both CS states is determined by the contribution of K_{12} . The addition or subtraction of the last term on the right side of Equation 2.10 corresponds to the energy of Ψ_1^s and Ψ_2^s , labeled as cs_+ or cs_- , respectively. The energy gap is $E_{cs_+} - E_{cs_-} = 2K_{12}$. The energy relation between the lowest closed-shell (cs_-) and open-shell singlet is:

$$E_{os} - E_{cs_{-}} = J_{12} - \frac{J_{11} + J_{22}}{2} + 2K_{12}. \tag{2.11}$$

Consequently, the final ordering of states depends strongly on the magnitude of K_{12} . However, it is important to remark that this condition is established for perfectly degenerate orbitals and considering that they are exactly the same in both states. In the presence of small (moderate) perturbations, Equations 2.9 and 2.11 are not entirely satisfied and the order of states can change, allowing the observation of both singlet and triplet ground state diradicals.

2.1.3.1 Diradicaloids: between closed-shell and diradicals

When the degeneracy of i and j orbitals in Equations 2.1 - 2.6 is lifted, their occupation is not equally probable anymore. The probability of having two electrons

in the energetically higher orbital is low and the latter is then referred to as the Lowest-Unoccupied-Molecular-Orbital (LUMO), while the lower-lying orbital have a higher probability of being populated and is referred to as the Highest-Occupied-Molecular-Orbital (HOMO). Hence, in the wave function of both closed-shell singlets, one configuration can be neglected, assuming that both electrons are either in the HOMO or in the LUMO, and state energies can be expressed as:

$$E_0 = 2h_H + J_{HH}, (2.12)$$

$$E_D = 2h_L + J_{LL}, (2.13)$$

where the i and j indices have been replaced by H and L referring to HOMO and LUMO orbitals, respectively. The subscripts correspond to the usual name for the ground state (S_0) and to the doubly excitation to the LUMO orbital (here labeled as D). Under the same approach of non-degenerated orbitals, the open-shell singlet and the triplet wave functions preserve their form and E_{os} and E_t can be written as:

$$E_{os} = h_H + h_L + J_{HL} + K_{HL}, (2.14)$$

$$E_t = h_H + h_L + J_{HL} - K_{HL}. (2.15)$$

According to the orbital splitting, two situations can be described. If the energy gap between orbitals is large, $h_L >> h_H$, the wave functions are not mixing and E_{os} and E_t are higher than E_{S_0} . Then, the overall picture corresponds to a closed-shell system with closed-shell ground state, the lowest excited triplet lying $2K_{hl}$ below the first excited singlet (HOMO-LUMO excitation) and the doubly excited state being higher in energy. However, if the energy gap between HOMO and LUMO is quite small, the probability of finding one or two electrons in the LUMO increases significantly and both closed-shell configurations must be considered to determine the properties of the ground state.

Rewriting Equation 2.1 as a configuration interaction (CI):

$$\Psi_1^s = C_1 \phi_H^2 - C_2 \phi_L^2, \tag{2.16}$$

where C_1 and C_2 are the coefficients that represent the weight of each configuration in the ground state wave function. Large C_1/C_2 are indicative of the closed-shell character of the system; as it decreases, the diradical character increases up to a value of 1, for which the system is a perfect diradical. The triplet state is then the first excited state, but lies very low in energy due to the small HOMO-LUMO gap, and might be degenerate with the singlet or, even, the ground state.

Systems in the situation between closed-shell and perfect diradicals are named diradicaloids (or diradical-like), and can be defined as molecules with two electrons occupying two quasi-degenerate orbitals. Interestingly, most of the organic systems with diradical character studied as candidates for optoelectonic or photovoltaic applications are indeed diradicaloids.

2.1.4 Characterization of diradicaloids

Diradical character is not an observable and thus cannot be quantified directly, neither theoretically or experimentally. Several quantities have been defined to characterize diradicals and diradicaloids: i) the singlet-triplet energy gap (ΔE_{ST}), in order to establish the multiplicity of the ground state and the interaction between both radicals ($2K_{12}$); ii) the HOMO-to-LUMO energy gap, to determine the degeneration of frontier molecular orbitals; and iii) the effective occupation of these orbitals.

 ΔE_{ST} can be approximated by twice the exchange integral $(2K_{12})$, which is the measurement of the electron-electron repulsion between two radicals. If $2K_{12}$ is equal to zero, the system is twice a radical (like two non-interacting doublets), which is the case of biradicals, as aforementioned. When $2K_{12}$ is different from zero, the radical interaction can be classified as antiferromagnetic (singlet ground state) or ferromagnetic (triplet ground state) if $2K_{12} < 0$ or > 0, respectively. While, in a perfect diradical there is one electron occupaying each orbital, there are 2 and 0 electrons in HOMO and LUMO orbitals, respectively, in the CS configuration.

Reliable characterization of diradicals needs robust electronic structure methods. Nevertheless, some useful rules can be used to describe diradicals and diradicaloids in a very basic way through graphical representation such as the Ovchinnikov's rule and the Clar's sextet rule.

2.1.4.1 Clar's sextet rule

In benzene rings, two equivalent forms with three localized double bonds are in resonance (cyclohexatriene), identified as Kekulé resonance structures. By superposition of both structures, a Clar's sextet is formed, *i.e.*, the benzene ring in which 6π -electrons are delocalized (Figure 2.4 a). Benzene is more stable than cyclohexatriene owing to the effect of aromaticity.(11) The formation of Clar's sextets explains the effect of the aromaticity in extended benzenoid systems (named as alternant polycyclic aromatic hydrocarbons or simply PAHs)(12). Clar's rule establishes that the resonance structure with the largest number of aromatic rings is the one that contributes the most to the electronic structure character of the system.(11) The extra stabilization in such structures is the result of having an "extra" aromatic ring.

For a given PAH, Kekulé and non-Kekulé structures (forms with absence and presence of unpaired electrons, respectively) can be drawn. In the case of radicals and diradicals, the non-Kekulé forms are the ones with the largest number of Clar's sextets. Following the Clar's rule, they have a critical role in the description of these systems(13), in which the instability induced by the unpaired electrons is compensated by the aromaticity gain. For instance, Figure 2.4 shows the resonance structures with their respective Clar's sextets for phenanthrene (b) and anthracene

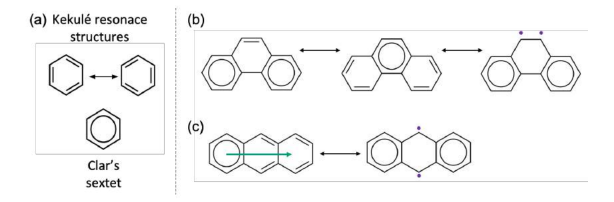


Figure 2.4: Benzene resonance structures and the Clar's sextet, which is represented as a circle inside the six-membered ring (a). Resonance structures of phenanthrene (b) and anthracene (c) with their Clar's sextets. The green arrow represents a Clar's sextet that can be localized in either one of the three benzene rings.

(c), PAHs with three fused benzene rings. Two Clar's sextets can be drawn in each structure, localized in the rings at the molecular end, which leads to having two unpaired electrons in the central benzene ring of each PAH. In phenanthrene, the presence of two radicals is largely unfavorable, then these electrons can form a π -bond and have a closed-shell structure, stabilizing the system. On the other hand, radicals in anthracene can not be bonded together or with adjacent atoms. The three equivalent closed-shell structures have the biggest influence in its electronic structure, but a fairly small diradical character can be expected. In this way, the (potential) radical character of PAHs can be predicted by the Clar's rule.

2.1.4.2 Ovchinnikov's rule

In alternant systems, as PAHs, simple models like Ovchinnikov's(14) rule can be used to predict the ground state spin multiplicity. According to this rule, atoms of a system can be classified in two types, labelled generically N and N^* (Equation 2.17), in such an arrangement that N atoms are surrounded only by N^* atoms and vice versa. Preferred ground state spin is predicted as the half of the difference between the number of carbon atoms belonging to N or N^* type (opposite lattices).

$$S = \frac{|N - N^*|}{2}. (2.17)$$

A special situation occurs when the number of N and N^* is not the same (Figure 2.5 b), which results in systems holding a high-spin ground state. This rule has been employed to predict the spin multiplicity of a lot of aromatic systems, as well as in the enginnering of structures with high spin-multiplicity state.

More formally, Lieb's theorem assets that the ground state-multiplicity of a bipartite lattice system is obtained from the Equation 2.17. (15)In the organic chemistry context, the Ovchinnikov's rule is equivalent to the Lieb's theorem.

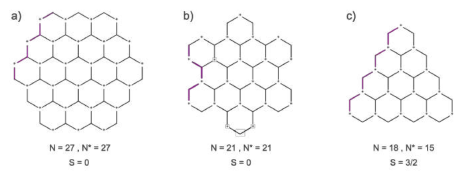


Figure 2.5: PAHs with different shapes and edges. N^* and N are the number of carbon atoms in each sublattice within the Ovchinnikov's rule. a) Hexagonal PAH with singlet ground state; b) armchair-edged hexagonal PAH with singlet ground state; and c) triangular PAH with quartet-spin ground state.

2.1.5 Triradicals and beyond

Molecules with an odd number of electrons inevitably have unpaired electrons. The diradical concept for the case of two electrons in two degenerate orbitals can be generalized to tri-, tetra-, pentaradicals, etc, and so on, for systems with a larger number of unpaired electrons. While there is an extensive number of examples of organic monoradicals and diradicals in the literature, the chemistry of triradicals (and beyond) has been less explored due to the increased instability associated with the increase of the number of unpaired electrons. Nevertheless, in the past few years fascinating examples of organic polyradicals have been designed, synthesized and characterized. For a couple of decades, Rajca and his group developed the concept of magnetic clusters, systems in which a core of high spin-multiplicity is growing in branches that couple diradicals in a polymeric way. With this strategy, the maximum spin-multiplicity reached was $S \approx 5000.(16)$ These systems with very large polyradical character are particularly attractive for applications as supermagnets and insulating spin glasses.

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Chapter 3

Electronic structure methods and computational tools

The underlying physical laws necessary for the mathematical theory of a large part of physics and the whole of chemistry are thus completely known, and the difficult is only that the exact solution of these laws leads to equations much too complicated to be soluble.

(P.A.M. Dirac, 1929)

This chapter sets the general quantum chemistry framework for the study of the electronic structure of molecular systems. In the following, I introduce and briefly describe an incomplete list of electronic structure approximations for the calculation of electronic states and their properties employed in the rest of chapters of this thesis.

3.1 The Schrödinger Equation

Quantum chemistry has served to describe all the physical and chemical properties of atoms and molecules. In a raw approximation, there is a mathematical object from which any desired information can be obtained from a chemical system. This is the Schrödinger equation, which without relativistic effects is expressed as:

$$\hat{H}\Psi(\mathbf{r},t) = i\hbar \frac{\delta\Psi(\mathbf{r},t)}{\delta t},\tag{3.1}$$

where \hat{H} is the well-known Hamiltonian operator, whose eigenvalue is the energy of the system. Normally, the wave function Ψ depends on the position (\mathbf{r}) of the particles and time (t) evolution, it is named as the *time-dependent Schrödinger* equation. If the Hamiltonian does not have any dependence with time, Equation 3.1 can be written as:

$$\hat{H}\Psi(\mathbf{r})F(t) = \Psi(\mathbf{r})i\hbar \frac{\delta F}{\delta t},$$
(3.2)

where F is the only term in which the time is involved, and $\Psi(\mathbf{r})$ depends only on the spatial coordinates. By keeping only the spatial term, the Schrödinger equation is written as:

$$\hat{H}\Psi(\mathbf{r}) = E\Psi(\mathbf{r}),\tag{3.3}$$

which is called as time-independent Schrödinger equation.

In quantum chemistry and molecular physics \hat{H} in equation 3.3 corresponds to the molecular (or atomic) Hamiltonian. Under the Born-Oppenheimer approximation, atomic and electronic coordinates are treated separately and the electronic Schrödinger equation is expressed as:

$$\hat{H}_e(\mathbf{r}, \mathbf{R})\phi(\mathbf{r}; \mathbf{R}) = E_e(\mathbf{R})\phi(\mathbf{r}; \mathbf{R}), \tag{3.4}$$

where \mathbf{r} and \mathbf{R} are the electronic and atomic coordinates, respectively, the electronic wave function $\phi(\mathbf{r}; \mathbf{R})$ and its energy $E_e(\mathbf{R})$ depend parametrically on the atomic positions \mathbf{R} .

In principle, all the information necessary to fully describe the electronic structure of a molecular system is provided by solving the electronic Schrödinger equation. In practice, this is a complicated task due to the large number of variables, and equation 3.4 can be solved exactly only for a couple of systems (Hydrogen and hydrogen-like).

3.2 Electronic Structure Approximations

Nowadays, seeking to solve the Schrödinger equation has lead to the development of several quantum chemistry methods based either on the wave function (Hartree-Fock and post-Hartree-Fock methods, e.g. Møler-Plesset perturbation theory (MP), configuration interaction (CI), multiconfigurational self-consistent field (MCSCF), couple cluster (CC), etc) or based on the electronic density (density functional theory, density matrix functionals).

3.2.1 Density Functional Theory

Within the Density functional theory (DFT) the Schrödinger equation can be solved in terms of the electronic density (ρ) avoiding the construction of the wave function of a N-electron system. The main premise in DFT is that the energy and all properties of a given system can be determined as a functional of ρ . This affirmation is known as the 1st theorem of Hohenberg and Kohn, establishing also that there is a unique external potential (v) determined by each ρ . At the same time, the electronic density depends (only) on the spatial coordinates \mathbf{r} , hereafter the density will be written as $\rho(\mathbf{r})$. Hence, within the DFT framework, the energy, and all properties of the system, are expressed as a functional of the density (a functional is a mathematical rule that express the dependence of a function on a function) and can be written as:

$$E[\rho] = T[\rho(\mathbf{r})] + V_{ne}[\rho(\mathbf{r})] + V_{ee}[\rho(\mathbf{r})]$$
(3.5)

$$V_{ne}[\rho] = \int \rho(\mathbf{r})v(\mathbf{r})d\mathbf{r}.$$
 (3.6)

The total energy of a system (Equation 3.5) includes the kinetic energy $(T[\rho(\mathbf{r})])$, electron-electron $(V_{ee}[\rho(\mathbf{r})])$, and nuclei-electron $(V_{ne}[\rho(\mathbf{r})])$ interaction energies, all of them as function of the $\rho(\mathbf{r})$.

The second Hohenberg and Kohn's theorem ensures the existence of an "exact" ρ (ρ_0) that minimizes the electronic energy of the ground state of the system. Despite the importance of this theorem, the general expressions of $T[\rho(\mathbf{r})]$ and $V_{ee}[\rho(\mathbf{r})]$ in Equation 3.5 are unknown, and in practice the description of the ground state electron density relies on further approximations.

Kohn and Sham established the most used approach in the quantum chemistry field to solve the Equation 3.5. Their method states that the density of a real system can be mapped to a single determinant wave function constructed by the Kohn-Sham orbitals $\psi_i(\mathbf{r})$ (orbitals of a system of non-interacting electrons), in such way that the density of the real system is:

$$\rho(\mathbf{r}) = \rho_s(\mathbf{r}) = \sum_{i=1}^{N} |\psi_i|^2(\mathbf{r}), \qquad (3.7)$$

and the electronic energy of the real system is defined as:

$$E[\rho(\mathbf{r})] = V_{ne}[\rho(\mathbf{r})] + T_s[\rho(\mathbf{r})] + J[\rho(\mathbf{r})] + E_{XC}[\rho(\mathbf{r})],$$

(3.8)

where, $T_s[\rho]$ is the kinetic energy of a system of non-interacting electrons, $J[\rho]$ is the Coulomb interaction energy, and $E_{XC}[\rho]$ is the exchange and correlation energy (the key quantity of DFT), which is defined as the difference between the "real" energy of the system and the KS one.

DFT-KS is formulated as an exact method, which with the correct terms yields the exact description of an N-electrons system. Despite all the terms in Equation 3.8 can be known, the practical limitation is that the exact exchange-correlation functional is unknown. In this sense, since the birth of KS-DFT numerous attempts in the search for the exact E_{XC} have been done. These approximations can be classified as the those based on the local density (LDA), generalized gradients of the density (GGA), and its derivatives of high order (metaGGA), or hybrid functionals that combine pure density functionals with exact exchange. Due to the large quantity of available functionals in quantum chemistry, a large sort of problems can

been addressed by the applications of DFT in the chemistry, physic, biology, etc. areas. But that does not mean that DFT offers all the time the "correct" description for the correct reasons, the use of of parameters to counterbalance the lack of exact functional can lead to cancellation of errors, and as a consequence can lead to meaningless results.

3.2.1.1 Constrained-DFT

Sometimes, it might be desirable to solve the KS equations under some physical constraints to the electron density. For instance, it might be interesting to restrict the net spin density of the system over some specific atoms, or maybe to enforce different amounts of electron density on different spatial regions, such as over electron donor and withdrawing groups in push-pull systems. The constrained-DFT (CDFT)(1) formalism corresponds to the minimization of the total energy under a constraint, in which a Lagrange multiplier λ is used to build a functional W as:

$$W[\rho, \lambda] = \min_{\rho} \max_{\lambda} [E[\rho(\mathbf{r})] + \lambda (\int_{\Omega} \rho(\mathbf{r}) d^3 r - N)]. \tag{3.9}$$

For a given restriction with N value in a Ω space, the energy of the whole system is computed by adding the energy of the constrain to the energy of the system without constrains $(E[\rho(\mathbf{r})])$. The density satisfies the constrains but is also the one that minimize the total energy of the system. Under the Becke's partition(???) the charge or spin population on atom A is weighted as a function of the distance from the nucleus of atom A by w_A (1 close to the nucleous of atom A and 0 close to the nucleus of other atom), and defined as:

$$N_A = \int w_A^{Becke}(\mathbf{r})\rho(\mathbf{r})d^3r. \tag{3.10}$$

Therefore, several constrains over both, charge and spin, can be fulfilled simultaneously in the ground state. Despite its limitations and range of applicability, CDFT has shown to be a valuable tool to describe charge-transfer states and spin localization.(2)

3.2.2 Time-dependent density functional theory

Time-dependent density functional theory (TD-DFT) is the DFT analog to deal with excited states. (3; 4) The Runge-Gross theorem is the equivalent to the first Hohenberg and Kohn's theorem, it relies in the fact that exists a one-to-one mapping between time-dependent electron density and the time-dependent external potential of an N-electron system that corresponds with a time-dependent Schrödinger equation. (5; 6)

The excitation energies (ω) in TD-DFT can be generally obtained as the solution of:

$$\begin{pmatrix} \mathbf{A} & \mathbf{B} \\ \mathbf{B} & \mathbf{A} \end{pmatrix} \begin{pmatrix} \mathbf{X} \\ \mathbf{Y} \end{pmatrix} = \omega \begin{pmatrix} \mathbf{1} & \mathbf{0} \\ \mathbf{0} & -\mathbf{1} \end{pmatrix} \begin{pmatrix} \mathbf{X} \\ \mathbf{Y} \end{pmatrix}, \tag{3.11}$$

in the linear-response framework (LR-TD-DFT)(7), this equation is known as the Casida's equation and represents one of the most used methods to obtain the TD-DFT energies. Since the matrix (**A-B**) is positive definite for real orbitals can be simplified as

$$(\mathbf{A} - \mathbf{B})^{1/2}(\mathbf{A} + \mathbf{B})(\mathbf{A} - \mathbf{B})^{1/2}\mathbf{Z} = \omega^2 \mathbf{Z},$$
(3.12)

where, $\mathbf{Z} = (\mathbf{A} - \mathbf{B})^{-1/2}(\mathbf{X} + \mathbf{Y})$ contains the information about occupied orbitals rotating into virtual orbitals. Matrices \mathbf{A} and \mathbf{B} depends on the exchange-correlation functional used.

Further approximation to the Casida's equation is the Tamm-Dancoff approximation (TDA) in which the Equation 3.12 is simplify to $\mathbf{AX} = \omega \mathbf{X}$ neglecting the de-excitation contributions to the energy. Although, the simplification that TDA means in TD-DFT, the results are qualitatively compared with the LR-DFT ones.

As DFT formulation, TD-DFT is formally an exact theory, which with the universal exchange-correlation functional can describe all excitations of a system. However, in practice, the explicit form of the functional is unknown. Under the adiabatic approximation(8) TD-DFT cannot deal with multiple excitations, and non-adiabatic effects, e.g., conical intersections, are wrongly predicted. (9) Additionally, standard functionals are unable to recover the correct $\frac{1}{r}$ asymptotic behavior of electronic interactions,(10) which prevents the proper description of electronic transitions with weak hole/electron overlap, like in charge transfer excitations or Rydberg states.(? 2; 11) Schemes to address this problem have been proposed by modeling the decaiment of the exchange-correlation potential at short and long ranges separately.(12) In this approximation the short-range exchange is treated with a local potential, while the long-range expression is represented by the "exact" exchange term.(11)

3.2.3 Spin-flip methods

An appropriated method to describe the ground state of strongly correlated systems, such as diradicals and polyradicals, must be able to deal with degeneracies and near-degeneraies. In principle, non-dynamic (strong) correlation induce the needed for a multi-configurational ansatz that can be treated by multireference methodologies, where the ground state is described as the combination of two or more (semi-)equally important configurations (from the definition of diradicals these kind of methods offer an adequate scheme to characterize them). The complete active space-self consistent field (CASSCF) method is one of the methodologies most extensively used

to deal with non-dynamic correlation. Instantaneous (dynamic) electron correlation effect can be added a posteriori by means of multireference perturbation theory, e.g., CASPT2. The main "problem" with these methodologies is its large computational cost, which makes them prohibitive in the study of medium to large size molecules.

As an alternative that allows the inclusion of both, dynamic and non-dynamic correlation, a family of methods based on the spin-flip operator were proposed to obtain a multi-configurational description through a single-reference ansatz. (13) The spin-flip (SF) approach was formally developed in 2001 (??) within the equationof-motion (EOM) formalism to address the multi-configurational problems arising in the proper description of homolytic bond cleavage. In a given molecule with singlet ground state that is splitted in two fragments, each fragment takes one electron of the broken bond. At the equilibrium distance, one configuration might be enough to describe the system, that is, the closed-shell singlet configuration with two electrons in the bonding orbital. On the other hand, when the bond is elongated, the electronic population of the antibonding orbital gains importance in the description of the ground state of the system. At dissociation, (at least) two configurations are necessary to describe the electronic structure of the lowest singlet. On the other hand, these correlation effects are less important in the triplet state, which well described by a single configuration along the entire bond dissociation. (14; 15) According to this, SF methodologies use a well-behaved high-spin configuration as a reference to construct correlated wave functions. From the high-spin reference, e.g., the $M_s = 1$ of triplet, the target states are obtained by the action of an excitation operator flipping the spin of one electron, $\alpha \to \beta$:

$$|\Psi\rangle = \hat{R}^{SF} |\phi_0\rangle, \qquad (3.13)$$

where \hat{R}^{SF} is the SF excitation operator connecting the ϕ_0 (reference) state to the target state Ψ . Excitation operator \hat{R}^{SF} can be expanded as a linear combination of creation and annihilation operators, as:

$$\hat{R}^{SF} = \sum_{l} \hat{R}_{l}^{SF} = \sum_{ia} r_{i}^{a} a_{\beta}^{\dagger} i_{\alpha} + \frac{1}{2!} \sum_{ijab} r_{ij}^{ab} a_{\beta}^{\dagger} b_{\sigma}^{\dagger} i_{\alpha} j_{\sigma} \dots$$
 (3.14)

where $\sigma = \alpha, \beta$. Truncation of Equation 3.14 to different levels of excitations give rise to an hierarchy of SF methods. From application of the SF operator over the high-spin reference, a triplet for instance $(M_s = +1)$, final states are those with $M_s = 0$, *i.e.*, singlets and triplets in Figure 3.1.

Since its formulation in 2001, SF methods have been developed on top of well established wave function methodologies and in the DFT framework. (16) Spin-flip methodologies have been extensively used in the rationalization of bond-breaking process, determine the diradical (polyradical) character, non-linear optical properties, and photophysical process as singlet fission.

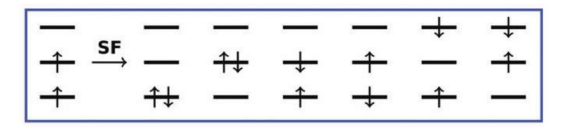


Figure 3.1: Target states obtained from the application of the SF operator (Equation 3.14) over a triplet reference of a system with two electrons in three orbitals. Figure adapted from reference (13).

3.2.3.1 Restricted active space - spin flip

The expansion in Equation 3.14 can be rearranged in series of excitation levels. In this way an active space where all possible configurations involving n_e electrons can be defined. Similarly to CASSCF methods, electrons included in the active space are those with higher reactivity, or more important in the description of the molecular electronic structure. By extension, two more spaces can be defined, one containing double occupied orbitals with less reactive electrons called as occupied space; while the other one groups the virtual orbitals, and is referred as the virtual subspace. Within the restricted active space (RAS) these spaces are considered as the RASI, RASII (active, A) and RASIII, see Figure 3.2 a). In these terms the spin flip operator is expressed as:

$$\hat{R}^{SF} = \sum_{\lambda \in A} \hat{R}_{\lambda}^{\Delta M_s = -n} + \sum_{\lambda \in h} \hat{R}_{\lambda}^{\Delta M_s = -n} + \sum_{\lambda \in p} \hat{R}_{\lambda}^{\Delta M_s = -n} + \sum_{\lambda \in hp} \hat{R}_{\lambda}^{\Delta M_s = -n} + \dots, (3.15)$$

configurations in $\sum_{\lambda \in h} \hat{R}_{\lambda}^{\Delta M_s = -n}$ are excitations from one double occupied orbital in the hole space to the active space, and configuration in $\sum_{\lambda \in p} \hat{R}_{\lambda}^{\Delta M_s = -n}$ corresponds to excitations from the active space to one empty orbital of the particle space.

In this manner the SF operator defines the RAS-SF method. (? 17) In general, the active space is determined by the high-spin reference and the number of spins-flipping, then in a RAS-nSF calculation with (2n+1)-tuplet high spin reference the minimum active space should contain 2n electrons and 2n spin-orbitals. RAS-SF is a single-reference method, which can leads to the excitations illustrated in Figure 3.1, as double ones (or higher depending on n). Within the hole and particle approximation in RAS-nSF, Equation 3.15 is truncated after the third term of the right side. The first one represents the full configuration interaction within the active space (3.2b), second and third ones allows monoexcitations out of the active space as illustrated in Figure 3.2c). As a consequence, the spin completeness of the method is ensured, which in general is a problem for SF methodologies. (13)

 RAS_n -SF represents a good compromise dealing with non-dynamic correlation

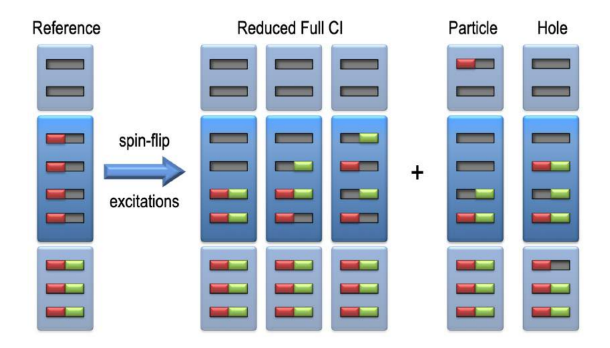


Figure 3.2: Schematic RAS-2SF subspaces partitioning, with 4 orbitals in the active space and a quintet as high spin reference (a). Resulting configurations in the active (b), and hole and particle spaces (c). Figure adapted from reference (?).

in (quite) large systems with an affordable computational cost. As demonstrated in this thesis (and in a large quantity of examples in the literature (18; 17)), RAS-nSF method is successfully used to characterize diradicals (polyradicals) and different excited state processes as singlet fission. Despite the advantages that RAS-SF method (and RAS-CI methos in general), one of its better-recognized deficiencies is the lack of dynamic correlation. In 2018 D. Casanova(?) proposed an hybrid method (RAS-srDFT) as a way to include dynamic correlation in RAS-CI methods, which consists in splitting the Coulomb operator (electron-electron repulsion, \hat{V}_{ee}) in long ($\hat{V}_{ee}^{lr,\mu}$) and short-range ($\hat{V}_{ee}^{sr,\mu}$) components. While the long-range is approximated by the wave function, the short-range term is provided by DFT. The energy of RAS-srDFT is obtained as:

$$E^{\text{WFT-}sr\text{DFT}} = \min_{\Psi^{\mu}} [\langle \Psi^{\mu} | \hat{T} + \hat{V}_{ne} + \hat{V}_{ee}^{lr,\mu} | \Psi^{\mu} \rangle + E_{Hxc}^{sr,\mu}[\rho]], \qquad (3.16)$$

where is the RAS-SF wave function is mixed with DFT short-range exchange-correlation term to include dynamic correlation in RAS-CI methods. (?)

3.2.3.2 Spin flip time-dependent density functional theory

The spin flip methodology has been also introduced in the frame of time-dependent density functional theory (SF-TD-DFT) addressing some of the drawbacks of LR-TD-DFT, as the inability to treat states in which the adiabatic coupling is important.

The best performance of SF-TD-DFT predicting singlet-triplet energy gaps in diradicaloids is obtained with functionals with large amount of HF exchange as BHHLYP(19) or PBE50(?) (with 50% HF exchange).

3.3 Computational tools for the characterization of electronic states

In addition to the computation of electronic states of molecules and their relative energies, the rationalization of their physical and chemical properties requires a detailed description of their electronic structure nature. In addition to the calculation of properties through well-defined operators, some useful descriptors can be established to describe not observable properties, e.g., the radical character, which is the core of this thesis. In the follow, I explain the indices I will use to describe the radical nature of the systems explored in the next chapters.

3.3.1 Diradical and polyradical character indeces

Probably the most extensively used index to quantify the di- and polyradical character of electronic states are the Yamaguchi indeces. (20) Which are based on the electronic occupation of the natural orbitals, defined as:

$$y_i = n_{\text{LUNO}+i}. (3.17)$$

Since HONO-i + LUNO+i = 2, y_i represents a measure of the electrons missing in HOMO-i to be double occupied. Then, y_i can take values between 0 and 1, and when $y_i = 0$, $\forall i$ the system is closed-shell, while $y_i = 1$ represents a perfect (di or tri, tetra or penta, etc) radical. Concretely, for compounds with even number of electrons y_0 is the index related to the diradical character, y_1 to the tetraradical character, y_2 to the hexaradical character, and so on. Polyradical character in systems with odd number of electrons are triradical (y_0) , pentaradical (y_1) , heptaradical (y_2) , etc.

3.3.2 Unpaired number of electrons

In open-shell systems the number of unpaired electron is a relevant quantity offering clues related with the presence of diradicals or beyond. For instance, a singlet state with two unpaired electrons can be classified as a singlet diradical.

In any electronic state the number of electrons can be computed by means of the occupation number of the natural orbitals (in our case they are obtained from the

one-particle density matrix extracted from RAS-SF calculations). The linear Head-Gordon equation (21) to quantify the number of unpaired electrons in a system is:

$$N_u = \sum_i (1 - abs(1 - n_i)),$$
 (3.18)

where, n_i is the natural occupation number of the natural orbital i. n_i can take values between 0 and 2; in the extremes, $n_i = 0$ (2), the orbital i is totally empty (doubly occupied) and does not contribute to N_u . If $n_i = 1$ there is just one electron in such orbital that can be related with an orbital holding a radical.

The spatial distribution of the unpaired electrons can be obtained through the graphical representation of N_u named as the fractional orbital density (FOD, as the graphical tool developed within the DFT framework(22; 23)).

$$\rho^{FOD}(\mathbf{r}) = \sum_{i}^{M} (\delta_1 - \delta_2 f_i) |\phi_i(\mathbf{r})|^2, \tag{3.19}$$

which by integration through all the space recover the N_u value (Equation 3.18).

3.3.2.1 Fragment-based wave function analysis

In the study of photophysical processes in dimers or aggregates is important to determine the electronic character of the excited states and the contribution of monomers to the collective properties. To describe the adiabatic excited states of the complex in terms of the diabatic states in the monomers, I used an approach developed by Luzanov, Casanova and Krylov (24; 25), in which the wave function of the dimer or (oligomer) is computed and then deconstructed in terms of the individual chromophores under a super-molecule scheme. In the case of RAS-SF, the canonical orbitals obtained from the SCF procedure are localized in fragments before the correlated calculation takes place. The total wave function is written as:

$$|\Psi\rangle = |\Psi^{LE}\rangle + |\Psi^{ME}\rangle + |\Psi^{CT}\rangle.$$
 (3.20)

Every term in the right side of the Equation 3.20 is determined by means of charge and spin cumulants identifying unambiguously local excitations (LE), charge transfer states (CT), and multiple excitation (ME).

Localized excitations in a dimer can be written in term of specific contributions from each monomer as:

$$|\Psi^{LE}\rangle = c_A |A^*B\rangle + c_B |AB^*\rangle, \qquad (3.21)$$

where A^*B are all possible combinations of A in an excited state and B in the ground state (or the opposite AB^*). In the same way charge resonance configurations are weighted as:

$$|\Psi^{CR}\rangle = t_{AB} |A^+B^-\rangle + t_{BA} |A^-B^+\rangle. \tag{3.22}$$

Finally, states in which both monomers are simultaneously excited giving rise to multiexcitations are described as:

$$|\Psi^{ME}\rangle = d_{AB} |A^*B^*\rangle. \tag{3.23}$$

By the combination of Equations 3.21, 3.22 and 3.23 in Equation 3.20 every adiabatic state can be expressed in terms of diabatic states, where $|c_A|^2 + |c_B|^2 + |t_{AB}|^2 + |t_{BA}|^2 + |t_{BA}|^2 + |t_{AB}|^2 = 1$ includes all the possible contribution of diabatic states to the adiabatic wave function.

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Chapter 4

Polycyclic aromatic hydrocarbon diradicals

Ever since its first experimental isolation in 2004(?), graphene has attracted immense interest among the scientific community, touted as a miracle material owing to its extraordinary physical and mechanical properties. (1; 2) Graphene can be viewed as an extended sheet of fused benzene rings. In this way, finite compounds with honeycomb-like structures can be considered as graphene derivatives, such as buckyballs, tubes, sheets and multidimensional graphene-fragments (0D, 1D and 2D; and 3D structures obtained as stacked graphene layers). In the nanoscale (1-100 nm), 2D-graphene fragments are known as graphene nanofragments or nanoflakes (GNFs), also referred as alternant polycyclic aromatic hydrocarbons (PAHs). The wide variety of shapes that graphene derivatives adopt, in particular GNFs together with their emerging molecular properties make them a fancy research topic. Its value in diverse fields has been demonstrated: optoelectronics, photocatalysis, energy storage, medical and biological applications, environmental sciences, and quantum computing. Most of these applications are related to electronic and magnetic properties, and are dictated by the topology and edge shape together with the size of the nanostructure. For instance, in GNFs, the stability of the ground state is directly related to the geometrical arrangement of the benzene rings. And the shape of their edges, zigzag or armchair, (Figure 4.1) modulate their electronic structure. GNFs with an armchair edge are closed-shell systems. While zigzag-edged GNFs have open-shell configuration with potential to exhibit high-spin ground state. Open-shell GNFs are very interesting as the main component of optoelectronic devices, but as a consequence of their radical (radicaloid) nature their synthesis is cumbersome and they have weak stability.

Through this chapter, I present the detailed electronic description of representative 1D and 2D GNF systems. The arising of radical character and its dependence with topology is rationalised in depth, in the first place by means of simple electronic structure analysis like Hückel model, Ovchinnikov's rule, and Clar's sextet rule(3)

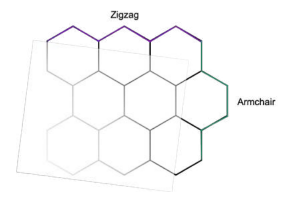


Figure 4.1: Armchair (green) and zigzag (purple) edges of a GNFs.

related to their classification as Kekulé and non-Kekulé structures. But mainly, by powerful electronic structure calculations, namely, restricted active spin-flip approach (RAS-SF), that allows the appropriated description of the radical nature and correlation effects of these systems.

In the first section, the intricacies of the electronic properties of smallest members of triangular graphene nanofragments (TGNFs) family in their ground and low-lying excited states are fully unravelled. Radical and diradical nature of these systems is described and rationalized in detail. Additionally, the modulation of their electronic and magnetic properties is explored by means of selective boron or nitrogen substitution at carbon sites as well as by hydrogen saturation. The second part of the chapter explores the nature of electron-correlation effects in carbon nanobelts and linear oligoacenes.

4.1 Triangular Graphene Nanofragments

TGNFs consist in benzene rings fused following D_{3h} symmetry, with all edges exhibiting a zigzag pattern that allows them to hold open-shell character. Analysis of the Clar's resonance structure of the small TGNFs shows the presence of at least one unpaired electron (Figure 4.2). Consequently, they are considered as the smallest open-shell non-Kekulé systems, and it is expected that all members of this family possess a high-spin ground state. In the following, the smallest members of the family, phenalenyl (1) and triangulene (2) are considered to discuss the implications of their triangular shape and resultant topology over radical and diradical character. The modulation of their electronic structure is then addressed via insertion of heteroatoms and hydrogenation.

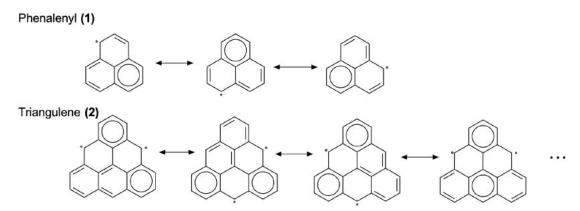


Figure 4.2: Clar's resonance structures in smaller triangulene graphene nanofragments: phenalenyl (1) and triangulene (2).

4.1.1 Pristine phenalenyl and triangulene

The smallest structure of the TGNF family, phenalenyl (1), is formed by three fused benzene rings sharing a central carbon atom and exhibits an odd number of electrons (13 π -electrons), thus inevitably resulting in a radical system. Ovchinnikov's rule predicts a spin-doublet ground state (Figure 4.3 a). This prediction is supported by the Hückel model, which shows one unpaired electron in the 1^2A_1 ground state (Figure 4.3 b).

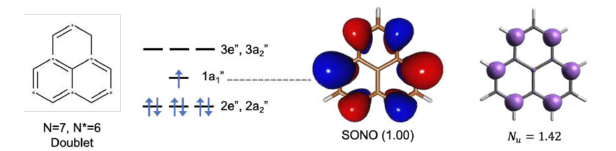


Figure 4.3: Characterization of the ground state of 1 a) Ovchinnikov's rule prediction; b) Hückel diagram; c) Semi Occupied Natural Orbital and occupation number; and d) FOD and number of unpaired electrons for the ground state. c) and d) were computed at RAS-SF level.

The next non-Kekulé triangular PAH, triangulene (2), differs from 1 by the addition of an extra row of benzene rings. While the molecular symmetry is not altered, 2 possesses an even number of carbon atoms and 22 π -electrons. The Ovchinnikov's rule and Hückel model predict a spin-triplet ground state (1³ A'_1 , Figure 4.4 a and b). The high-spin multiplicity of the ground state of 2 has been investigated theoretically and observed experimentally in its derivatives.(4; 5; 6?)

It is interesting to notice that both Hückel MO energy diagrams present degeneration degrees beyond the one dictated by the molecular D_{3h} symmetry, *i.e.*, there

are up to three degenerated π -orbitals, when the maximum dictated by molecular symmetry is two. Such triple degeneration is triggered by the symmetry of the Hückel Hamiltonian. By construction, only interactions with first neighbors are considered, therefore as a result the Hamiltonian symmetry is higher than the molecular one. (???) Nevertheless, orbital degeneracy is lifted when performing ab initio calculations.

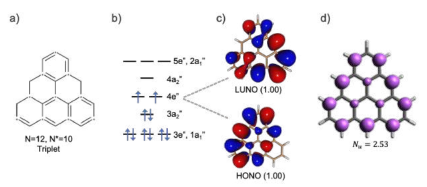


Figure 4.4: Characterization of the ground state of **2** a) Ovchinnikov's rule prediction; b) Hückel diagram; c) Frontier Natural Orbital and occupation numbers; and d) FOD and number of unpaired electrons for the ground state. c) and d) were computed at RAS-SF level.

The $2p_z$ orbital on the central carbon atom of **1** and **2** correspond to the A_2'' irreducible representation (D_{3h} point group). Then, this atom can only contribute to π -orbitals with that symmetry, *i.e.*, a_2'' molecular orbitals. In both systems, the electronic occupation of Hückel π -orbitals thus predicts the lack of unpaired electron density on the central atom, while $1a_1''$ and 4e'' π -orbitals are singly occupied in **1** and **2** respectively (Figure 4.3 b) and 4.4 b).

RAS-SF calculations provide further insights on the ground state radical nature of 1 and 2. Singly occupied natural orbital (SONO) of 1, depicted in Figure 4.3 c), belongs to the A_1'' irreducible representation, as predicted by Hückel model, and is localized over the molecular edges. Qualitatively, the same picture is offered by the FOD of the $1^2A_1'$ state (Figure 4.3 d). Similarly, the main contribution to the FOD, and consequently to N_u , of 2 is given by HONO and LUNO shapes and occupation numbers (Figure 4.4 c). Actually, both E'' irreducible representation are localized mostly over the edges of 2 with small contribution of inner carbons, apart from the central one.

Straightforward comparison of FOD distribution of 1 and 2 with their respective Ovchinnikov's representation pointed out the bilattice-like motif of TGNFs, where carbon atoms on the sublattice with more C atoms contribute to the unpaired electron density.

The lowest excited states of 1 correspond to $1^2E''$ and $1^2A_2''$ with vertical excitation energy of 2.95 and 3.29 eV, respectively (Table 4.1), in excellent agreement with MR-CI energies.(7) Doubly degenerated $1^2E''$ state corresponds to the promotion of

the edge electrons $(2e'' \to 1a_1'')$ whereas the $1^2A_2''$ state contains a partial displacement of central electrons to the molecular contour, as obtained from the $2a_2'' \to 1a_1''$ electronic excitation.

Considering now compound 2, the single electron occupation of the 4e'' orbitals in 2 results in a four dimensional space with A'_1 , A'_2 and E' symmetries. The antisymmetric spatial representation component of the direct product $E \otimes E$ corresponds to the ground state triplet $(1^3A'_2)$, while A'_1 and E' are linked to spin-singlet excited states (Table 4.1).

The energy gap to the lowest excited singlet of $2 (1^1E')$ is computed at 0.57 eV, again in line with values obtained by means of other correlated methods (0.56-0.84 eV) (???). This doubly degenerated singlet is mainly obtained as the $(3a_2'')^2(4e'')^2(4a_2'')^0$ configuration ($\sim 71\%$) with both states resulting from the single electronic occupation of 4e'' orbitals. The fractional occupation density is mainly localized on the molecular edges (Figure 4.5), resembling qualitatively the one exhibited by the triplet ground state (Figure 4.4).

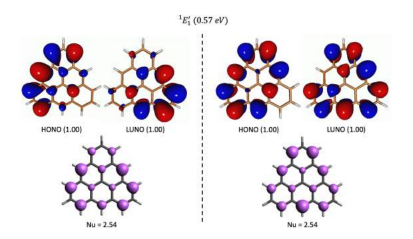


Figure 4.5: Frontier Natural Orbitals with their occupation numbers, and FOD and N_u of ${}^1E'_1$ state of 2.

The triplet-triplet transition energy to the first excited triplet of $\mathbf{2}$ (1³E' state) is computed at 2.82 eV, and involves electronic configurations with the partial occupation of the $3a_2''$ and $4a_2''$ orbitals, which means that the localization of the FOD on the central carbon is not symmetry forbidden for this state.

4.1.2 B-doped and N-doped TGNF

In light of the above characterization of 1 and 2, doping of these systems by substitution of carbon atoms with electron-donating and electron-accepting elements can be envisioned as an efficient tool to tune their electronic and magnetic properties.(??) Notably, experimental methods devoted to the controlled substitution of carbon

mol.	state	ΔE	N_U
1	$1^2 A_1''$	_	1.33
	$1^{2}E''$	2.95	2.17
	$1^2A_2^{\prime\prime}$	3.29	1.87
2	$1^{3}A'_{2}$	-	2.53
	$1^1E'$	0.57	2.54
	$1^3E'$	2.82	3.44

Table 4.1: Low-lying states of **1** and **2**, energy gaps (in eV) and unpaired electron numbers (N_U) computed at the RAS-SF/6-311G(d,p) level.

atoms by other species have been developed since the early studies on graphene, especially in the synthesis of N- and B-doped graphene. (8; 9; 10; 11; 12) It has also been evidenced that the electronic structure characteristics of these compounds depend not only on the dopant concentration, but also on the doping symmetry pattern. (13)

Next, I explore the changes in the electronic properties of **1** and **2** upon replacing 1, 2, or 3 C sites with electron-donating (N atoms) or electron-accepting (B atoms) elements (Figure 4.6). In order to preserve the chemical composition at the edges, all substitutions were performed over the inner carbons.

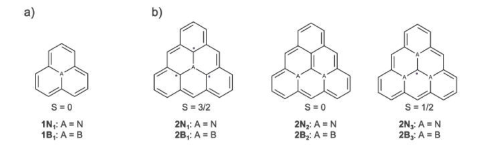


Figure 4.6: N- and B-doped derivatives of a) 1 and b) 2 systems with the spin predicted by Ovchinnikov's rule.

4.1.2.1 Phenalenyl derivatives

 $1\mathbf{B}_1$ and $1\mathbf{N}_1$ are the monodoped derivatives of phenalenyl. While both systems conserve the D_{3h} symmetry of their pristine equivalent, the electronegativity difference of B and N atoms with respect to C results in slightly longer bond lengths in $1\mathbf{B}_1$ and shorter ones in $1\mathbf{N}_1$ (B-C > C-C > N-C).

The heteroatom substitution of the central carbon modifies the number of π electrons with respect to **1**. In the application of the Ovchinnikov's rule to $\mathbf{1B}_1$ and $\mathbf{1N}_1$, the central site is not considered in the evaluation of carbon atoms belonging to

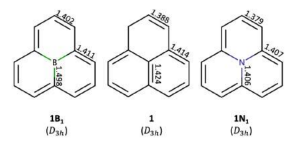


Figure 4.7: Bond lengths in Å in $1B_1$ (left), 1 (center) and $1N_1$ (right) systems.

opposite sublattices and ground state spin-singlet systems are predicted. However, this simplification cannot account for the electronic structure details of the doped systems, such as the diradical (or diradicaloid) character of the molecule that is dictated by the gap between occupied and virtual orbitals.

Since the $2p_z$ orbital of boron (nitrogen) is higher (lower) in energy than the carbon's counterpart, the a_2'' frontier π MOs are destabilized (stabilized) in $\mathbf{1B}_1$ ($\mathbf{1N}_1$), as obtained by the Hückel model (Figure 4.8). The energy shift of a_2'' orbitals might eventually result in small HOMO-to-LUMO energy gaps inducing open-shell character to the ground state wave function.

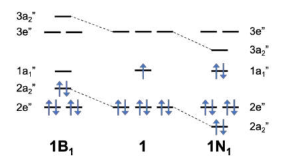


Figure 4.8: Hückel diagram of π -orbitals of $1N_1$ and $1B_1$.

Electronic structure calculations confirm the ground state spin-singlet nature of ${\bf 1B_1}$ and ${\bf 1N_1}$, and predict singlet-triplet energy differences smaller than 1 eV (Table 4.2). The quite small energy gap to the lowest triplet state suggest a partial open-shell character of molecular ground state singlets, both doped derivatives showing moderate diradical (or diradicaloid) character that approximately corresponds to half unpaired electron as measured by N_u values of 0.63 and 0.55 for ${\bf 1B_1}$ and ${\bf 1N_1}$, respectively. Differently to 1, the unpaired electron density emerges, for the most part, from the fractional occupation of a_1'' and a_2'' orbitals. The moderate amount of unpaired electrons in substituted systems is almost equally distributed between all contour carbons with contributions from the central doping atom, especially in ${\bf 1N_1}$ (Figure 4.9).

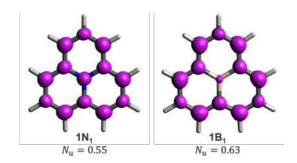


Figure 4.9: FOD of the ground state of $1N_1$ and $1B_1$.

The lowest excited singlet state of the doped phenalenyls is mainly obtained as the HOMO-to-LUMO single electron excitation $(a_1'' \to a_2'')$ for $\mathbf{1B}_1$ and $a_2'' \to a_1''$ for $\mathbf{1N}_1$) and belongs to the same spatial symmetry as the lowest triplet (A_2') representation. The computed energy of $1^1A_2'$ is systematically lower than $1^3A_2'$, which is the inverse of the common singlet/triplet state ordering but can be rationalized as the result of two effects:

i) The weak HOMO/LUMO exchange, K_{hl} , expressed as:

$$K_{hl} \simeq \frac{E_S - E_T}{2}. (4.1)$$

For both 1 derivatives, $\frac{\Delta E_{ST}}{2} \sim -0.06$ eV is computed at RAS-SF level. A better approximation of K_{hl} is obtained by the uncorrelated Single Excitation Configuration Interaction (CIS) method(????). At CIS/6-31G* level of theory, $K_{hl} = 0.14$ eV and 0.16 eV for $\mathbf{1B}_1$ and $\mathbf{1N}_1$, respectively. These small values are due to the poor spatial overlap between a_1'' and a_2'' orbitals as illustrated in the Figure 4.10.

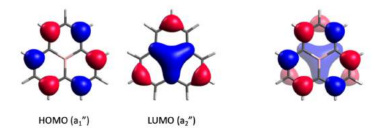


Figure 4.10: Frontier molecular orbitals (a_1'') and a_2'' of $\mathbf{1B}_1$ (left) and their superposition (right).

ii) Mixing of other configurations: the main configuration of $1^1A_2'$ and $1^3A_2'$ states correspond to the single electron occupation of a_1'' and a_2'' orbitals ($\sim 85\%$). However, the wave function the singlet state is mixed with other open-shell electronic configurations involving 3e'' orbitals, helping to stabilize it with respect to the $1^3A_2'$ state (Table 4.2).

mol.	state	ΔE_{ST}	N_u	configuration	contributions (%)
1B₁	$1^{1}A_{2}^{\prime}$	0.66	2.40	$(2e'')^4(2a''_9)^1(1a''_1)^1(3e'')^0(3a''_9)^0$	84.13
1	1 112	0.00	_,10	$ \begin{vmatrix} (2e'')^4(2a_2'')^1(1a_1'')^1(3e'')^0(3a_2'')^0 \\ (2e'')^3(2a_2'')^1(1a_1'')^1(3e'')^1(3a_2'')^0 \\ (2e'')^3(2a_2'')^1(1a_1'')^1(3e'')^1(3a_2'')^0 \\ (2e'')^4(2a_2'')^1(1a_1'')^1(3e'')^0(3a_2'')^0 \end{vmatrix} $	2.24
				$(2e'')^3(2a_2'')^1(1a_1'')^1(3e'')^1(3a_2'')^0$	2.24
	$1^{3}A_{2}'$	0.79	2.36	$(2e'')^4(2a_2'')^1(1a_1'')^1(3e'')^0(3a_2'')^0$	85.46
$\mathbf{1N}_1$	$1^{1}A_{2}'$	0.85	2.40	$ \begin{vmatrix} (2e'')^4(2a_2'')^2(1a_1'')^1(3a_2'')^1(3e'')^0 \\ (2e'')^4(2a_2'')^1(1a_1'')^1(3a_2'')^1(3e'')^1 \\ (2e'')^3(2a_2'')^2(1a_1'')^1(3a_2'')^1(3e'')^1 \\ (2e'')^4(2a_2'')^2(1a_1'')^1(3a_2'')^1(3e'')^0 \end{vmatrix} $	84.25
				$(2e'')^4(2a_2'')^1(1a_1'')^1(3a_2'')^1(3e'')^1$	2.10
				$(2e'')^3(2a_2'')^2(1a_1'')^1(3a_2'')^1(3e'')^1$	2.09
	$1^{3}A_{2}'$	0.96	2.36	$(2e'')^4(2a_2'')^2(1a_1'')^1(3a_2'')^1(3e'')^0$	85.76

Table 4.2: Low-lying states of **1** and **2**, energy gaps (in eV), unpaired electron numbers (N_u) and the main configuration contributions, computed at the RAS-SF/6-311G(d,p) level.

4.1.2.2 Triangulene derivatives

Triangulene allows different possible doping patterns preserving its D_{3h} symmetry or not, by replacing one, two or three carbon sites (Figure 4.6 b). The boron and nitrogen mono-substituted derivatives, $2\mathbf{B}_1$ and $2\mathbf{N}_1$, hold odd numbers of π -electrons. DFT results indicate a doublet ground state with adiabatic energy gaps to the quartet state (ΔE_{DQ}^{ad}) of 4.2 and 11.2 kcal/mol respectively, in disagreement with the Ovchinnikov's rule, which predicts a preference for the spin-quartet multiplicity.

Stabilization of the doublet state in $2\mathbf{B}_1$ and $2\mathbf{N}_1$ is driven by a Jahn-Teller distortion (14; 15?) that lowers the molecular symmetry to C_{2v} . The high-spin quartet of the D_{3h} geometry in both mono-doped species (predicted by the Ovchinnikov's rule as the ground state) is computed as the single electron occupation of a_2'' and e'' orbitals (${}^4A_2''$), and lies slightly above the two-folded doublet ${}^2E''$ state (Table 4.3). However, the D_{3h} arrangement is unstable upon certain structural deformations. The existence of a third order symmetry axis in TGNFs induces the presence of two-fold degenerate electronic terms E, which are prompt to instabilities with respect to nuclear displacements that reduce the symmetry and remove the degeneracies.(16)

The Jahn-Teller active distortion modes able to split the spin doublet state must be different from the totally symmetric representation and must be contained in the symmetric contribution of the direct self-product of the E'' representation. Therefore, Jahn-Teller distortions able to break the degeneracy of ${}^{2}E''$ states must follow an S(e') mode. Structure modification upon the action of the S(e') vibration breaks the degeneracy of e'' orbitals and allows e''/a''_{2} orbital mixing. As a result, the doublet configuration is stabilized in the C_{2v} geometry with the unpaired electron occupying a b_{2} -orbital (Figure 4.11 a). Moreover, the vertical gap to the spin-quartet

mol.	sym.	state	ΔE	N_u
$2B_1$	D_{3h}	$1^2E''$	-	1.58
		$1^4 A_2''$	0.15	3.03
$2B_1$	C_{2v}	$1^{2}B_{2}$	-	1.47
		1^4A_2	0.32	3.03
$2N_1$	D_{3h}	$1^2E''$	_	1.50
		$1^4 A_2''$	0.06	3.02
$\mathbf{2N}_1$	C_{2v}	$1^{2}B_{2}$	-	1.49
		$1^{4}A_{2}$	0.25	3.02
$2B_3$	D_{3h}	$1^2 A_2''$	-	2.99
		$1^4 A_2''$	1.40	3.25
$2N_3$	D_{3h}	$1^2 A_2''$	-	2.77
		$1^4 A_2''$	2.06	4.72

Table 4.3: Vertical energy gaps (in eV) between low-lying states with different spin-multiplicities and unpaired electron numbers (N_u) of B- and N-doped triangulene derivatives computed at the RAS-SF/6-311G(d,p) level.

state of $2\mathbf{B}_1$ and $2\mathbf{N}_1$ sensibly increases upon the S(e') distortion towards the C_{2v} geometry (Figure 4.11 b).

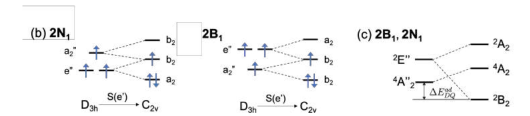


Figure 4.11: Qualitative representation of the S(e') distortion between the quartet (D_{3h}) and doublet (C_{2v}) optimized geometries of $\mathbf{2B}_1$ and $\mathbf{2N}_1$ (a), and state diagram of both monodoped triangulenes at each molecular symmetry (b).

Accordingly, while the unpaired electrons of the high-spin state symmetrically delocalize over the entire π -system, the unpaired electron density of the ground state doublet (1^2B_2) in the mono-substituted triangulenes does not preserve the three fold symmetry, and delocalize the spin density on one of the triangle edges of $2B_1$ and at one of the vertices in $2N_1$ (Figure 4.12).

Equivalently, the action of e'-symmetry Jahn-Teller distortion modes on **2** results in the splitting of the lowest ${}^{1}E'$ state. But, contrary to the situation in **2B**₁ and **2N**₁, such stabilization is not enough for the lowest spin-state to become the global minimum of the system ($\Delta E_{ST}^{ad} = -8.6 \text{ kcal/mol}$).

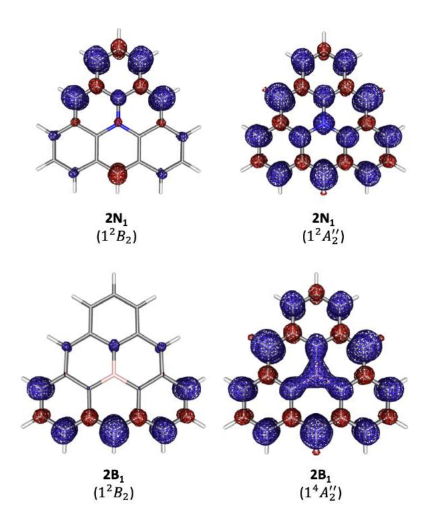


Figure 4.12: Spin densities of 1^2B_2 (C_{2v}) and $1^4A_2^{"}$ (D_{3h}) states of $2\mathbf{N}_1$ (top) and $2\mathbf{B}_1$ (bottom) in their respective optimized geometries.

The C_3 -symmetric substitution of three carbon atoms by boron or nitrogen in ${\bf 2B_3}$ and ${\bf 2N_3}$, respectively, stabilizes the low-spin doublet state. Contrary to the mono-substituted species, ${\bf 2B_3}$ and ${\bf 2N_3}$ keep the overall D_{3h} molecular symmetry, since the unpaired electron lies on a (non-degenerated) a_2'' -orbital. Characterization of their ground state radical character identifies one unpaired electron mainly localized on the central atom with small C_3 -symmetrical contributions from the doping atoms and the carbons at the center of the triangle edges, Figure 4.13. Computed energy gaps to the quartet state are shown in Table 4.3.

Finally, we consider the non-symmetric doping of $\mathbf{2}$ by replacing two inner carbon atoms ($\mathbf{2N_2}$ and $\mathbf{2B_2}$). As shown in Figure 4.6, this substitution pattern implies the reduction of molecular symmetry to C_{2v} . The connectivity between carbon atoms sensibly stabilizes the closed-shell configuration, resulting in a spin-singlet ground state with a weak diradical character identified by a quite small N_U value. Similarly to pristine triangulene, the unpaired electron density of $\mathbf{2N_2}$ locates over

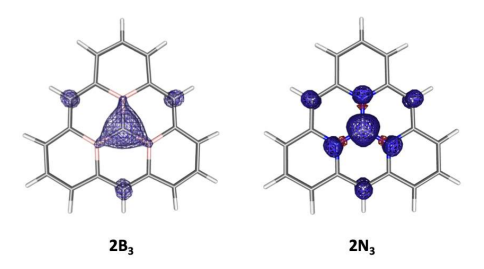


Figure 4.13: Spin densities of $2\mathbf{B}_3$ and $2\mathbf{N}_3$ doublet ground state.

the molecular edges, although it also exhibits radical character on the two doping atoms (Figure 4.14). The small diradical character of $2N_2$ is in agreement with the computed rather large gap to the lowest triplet state, as presented in Table 4.3.



Figure 4.14: Ground state fractional occupied density (FOD) of $2N_2$.

Therefore, we can conclude that doping of 1 and 2 with N and B results in the reduction of spin-multiplicity of the ground state. By extrapolation to bigger members of TGNFs, symmetric and asymmetric substitution with heteroatoms offers a way to obtain stable larger graphene nanofragments with zigzag edges.

4.1.3 Hydrogenated TGNF

Selective hydrogen saturation of peripheral carbon positions according to Ovchinnikov's rule has been shown to constitute a simple and effective strategy in order to design high-spin states of PAHs.(17) Here, we symmetrically hydrogenize three carbons of 1 and 2 molecules (Figure 4.15 a). The application of the Ovchinnikov's rule is done by removing the sp^3 carbon atoms and preserving only the conjugated system as illustrated in Figure 4.15 b.

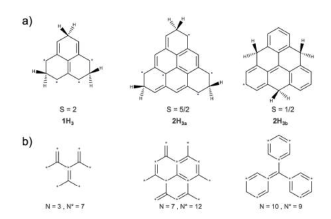


Figure 4.15: Hydrogenated derivatives of **1** and **2**. Multiplicity predicted by Ovchinnikov's rule is shown.

Saturation of the three corners of 1 results in a triallyl-methane type product in which the sp^3 atoms enforce molecular planarity with a D_{3h} symmetry. The topology of sp^2 sites in $1\mathbf{H}_3$ favors the high-spin stabilization, and its ground state corresponds to a spin-quintet $(1^5A'_1)$ state. The unpaired electron density at the ground state is distributed amongst the sp^2 atoms bonded to the CH_2 fragments and the central atom (Figure 4.16). The lowest doubly degenerated triplet and singlet states lie at 0.40 and 1.11 eV above the high-spin ground state, respectively, in very good agreement with accurate correlated *ab initio* and spin decontamination corrections of DFT energies.(17)

Similarly, hydrogenation of corner carbon atoms of $\mathbf{2}$ in $\mathbf{2H}_{3a}$ results in a high-spin ground state with five unpaired electrons, and with energy gaps to lower spin-multiplicities obtained from the spin-flip of one and two electrons of 0.26 and 0.66 eV, respectively. Contrarily, three-folded saturation of $\mathbf{2}$ at the center of the triangle edges $(\mathbf{2H}_{3b})$ reduces the imbalance between the number of carbons in each of the two sublattices, which stabilizes the low-spin (doublet) state, giving rise to a planar triphenylmethane-like radical structure (Figure 4.15 b).

Replacement of sp^2 by sp^3 carbon atoms in the periphery positions of **1** and **2** bears the rise of spin-multiplicity at ground state if done at the triangle's vertices. If the carbons in the center of the triangle edges are hydrogenated, the diminution of the ground state multiplicity is predicted, suggesting a selective way to modulate the

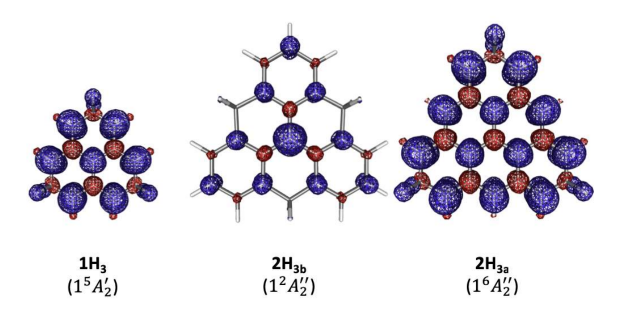


Figure 4.16: Spin density of hydrogenated derivative from 1 and 2.

mol.	sym.	state	ΔE	N_U
$1\mathrm{H}_3$	D_{3h}	$1^{5}A'_{2}$	_	4.16
	2 311	$1^{3}E'$	0.40	4.09
		$1^1E'$	1.11	4.05
$2\mathrm{H}_{3a}$	D_{3h}	$1^6 A_2''$	_	5.28
=11 3a	D 3n	$1^{4}E''$	0.26	4.79
		$1^2E''$	0.66	4.60
$\mathbf{2H}_{3b}$	D_{3h}	$1^2 A_2''$	_	1.29
=± 36	2 3n	$1^{2}E''$	3.16	1.82

Table 4.4: Vertical energy gaps (in eV) to the low-lying states with different spin-multiplicities and unpaired electron numbers (N_U) of H-saturated 1 and 2 derivatives computed at the RAS-SF/6-311G(d,p) level.

radical character of TGNFs by hydrogen saturation of peripheral carbon positions.

In summary, I have described the electronic structure of the doublet ground state of the phenalenyl molecule and the nature of its radical character arising from the single electron occupation of a a_1'' orbital delocalized at the molecular edges. Similarly, the radical character of ground state triplet of triangulene is located at the carbon edges with no participation of the central and vertex sites. Substitution of the carbon atoms by boron or nitrogen can be used to tune the electronic and magnetic

properties of TGNFs. The substitution symmetry pattern dictates the distribution of unpaired electrons and can be used to design TGNF ensembles with the desired properties. The fractional occupation of degenerated levels induces a Jahn-Teller distortion of mono-doped triangulene resulting in a stabilization of the doublet state, which cannot be taken into account by simple models such as Ovchinniko's rule. Selective saturation of carbon edge positions of 1 and 2 has a very strong impact on the ground state spin multiplicity, hence their magnetic properties.

4.2 Diradical character in cyclic and linear acenes

IUPAC Golden Book defines acenes as "polycyclic aromatic hydrocarbons consisting of fused benzene rings in a rectilinear arrangement". (????) Cyclic acenes or cyclacenes ([n]CC) can be obtained by fusing the ends of linear acenes ([n]Ac). In the context of the singlet-walled carbon nanotubes (SWCNT), [n]CC (also known as nanobelts) are important as they are consider as the smallest chemical unit to growing up size-specific singlet-walled carbon nanotubes (SWCNT).(18)

According to Ovchinnikov's rule, [n]CC and [n]Ac, presented in Figure 4.17 a), are predicted to hold spin-singlet ground state since they have same number of atoms in each sublattice. Even so, the presence of zigzag edges suggests the rise of open-shell properties, which is supported by the Clar's sextet rule. Only one one Clar's sextet ring can be drawn in closed-shell structures of [n]Ac, no aromatic rings can be formed without unpaired electrons in [n]CC (Figure 4.17 b)). These observations anticipate that linear and cyclic acenes have slight diradical character, at least. Furthermore, is predicted that the [n]Ac are most stable than [n]CC.

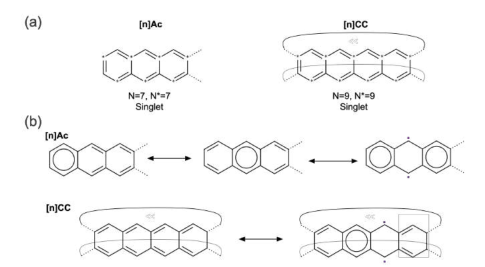


Figure 4.17: Ovchinnikov's rule applied to [n]Ac and [n]CC (a). Clar's resonate structures of [n]Ac and [n]CC (b).

At the beginning of the 50s decade cyclic acenes, nanobelts with zigzag edges,

were proposed and investigated theoretically. (???) The first attempt of synthesize a zigzag-cyclic acene was reported in 1987, when Stoddart et al. tried to obtain duodecacyclacene. (??) Since then the synthesis of [n]CC has been widely unsuccessfully attempted. (19) Recently, a carbon-nanobelt isomer of the [12] cyclophenacenehas has been isolated under standard conditions and characterized by X-ray diffraction by Povie et al. (20) Synthesis of linear acenes can be also challenging. The smallest pristine [n]**Ac** are stable under ambient conditions and commercially available (up to pentacene). Mainly tetracene and pentacene have been extensively used as component of organic thin film transistors, organic photovoltaics and solar cells. (21; 22; 23) Together with hexacene and its derivatives, they have been tested as singlet fission molecules. (24; 25) In medium size linear acenes, from hexacene to nonacene, the synthesis of stable compounds at ambient conditions have not been achieved yet but they are obtained under special conditions. (26; 27; 28; 29). Routes to prepare hexacene and heptacene consist mainly on the *in situ* generation of pristine acenes from its derivatives, which could be over metallic layers, (30) in solution, (31), under matrix isolation conditions, (????) or directly as part of a thin-film(32). Nevertheless, crystalline hexacene can be stable for few days under ambient conditions but in the dark, (???). Larger acenes can be photogenerated (33?) or obtained on-surface (? 34????) up to undecacene. Radical character has been associated to the unstable behavior of cyclic acenes and medium to large linear acenes.

With that in mind, a family of increasing size [n]CC, with n = 5 - 12, was thoroughly studied and compared with the parent [n]Ac (from pentacene to dode-cacene), so as to rationalise the relationship between topology and radical character.

Geometrically, the effect of different contour conditions, in this particular case if the acenes are closed or not, i.e., cyclic or linear acenes, significantly impacts their bond length alternation. While in [n]CC all rings are equivalent, benzene rings at the extremities of [n]Ac are smaller than those in the molecular center. If acenes are considered as two conjugated chains (up and down) linked by a bridge two kinds of C-C bonds can be defined, namely intrachain and interchain (or bridge) bonds. Intrachain bonds of [n]CC are close to C-C length in benzene and graphite (1.38) and 1.42 Å, respectively), (?) indicating the presence of two conjugated paths, top and bottom edges. Interchain distance between both paths is similar to the single bond length between two double bonds in butadiene, i.e., single bond connecting two sp^2 carbons (1.46 Å).(?) Therefore, [n]CC can be understood as two decoupled conjugated chains covalently bounded. On the other hand, distribution of C-C bond lengths in linear acenes is not uniform. Intrachain distances fluctuate between standard double bond and double bond in conjugated systems. While the bridging distances increase progressively from the edges to the center of the linear acene. Central ring of [n] Ac exhibit essentially the same bond lengths as the corresponding [n]CC compound. To exemplify, Figure 4.18 depicts differences between linear and cyclic undecacene, [11]Ac and [11]CC.

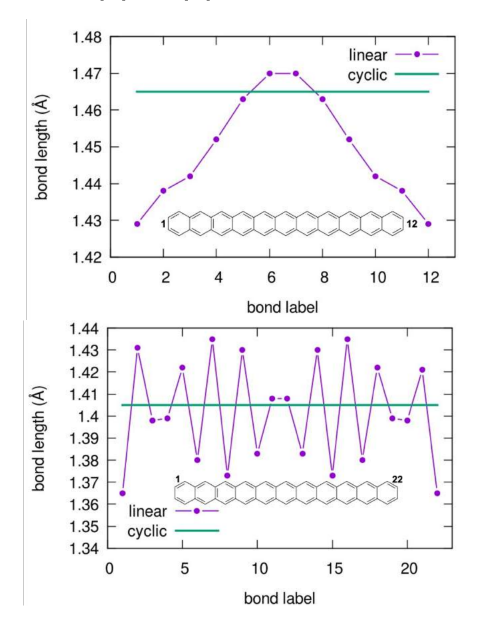


Figure 4.18: Consecutive C-C edge (top) and bridge (bottom) bond lengths for [11]Ac and [11]CC oligoacenes.

The Hückel model pointed out a difference in the electronic structure of both families thorugh their molecular orbital diagram, which helps to clarify the nature of radical character in both families of acenes (Figure 4.19). The HOMO-LUMO energy gap in [n]Ac decreases with the increase of the molecular size (*i.e.*, diradical character increases with n). Similarly, the energy gap between occupied and virtual orbitals of [n]CC decreases constantly with molecular size. However, the degeneracy of frontier molecular orbitals changes if the numeber of fused rings in these systems is odd or even. When n is odd, they hold a doubly degenerated HOMO and LUMO, at very large n can foresee the vanish of the HOMO-LUMO energy gap and the emergence

of the tetraradical character. On the other hand, even-[n]CC, two electrons located in two degenerated orbitals are predicted, suggesting a strong diradical character for these systems.

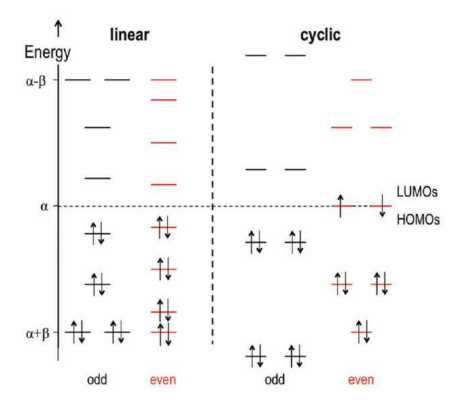


Figure 4.19: Hückel molecular orbital energy diagram around the Fermi energy level (E = α) for [n]Ac (left) and [n]CC (right) with an odd (n = 5) and even (n = 6) fused rings.

These elementary considerations evidence a strong relation between topology and the radical character of both linear and cyclic acenes. Deep electronic structure characterization of these systems through wave function analysis is summarised for each family in following subsections.

4.2.1 Linear acenes

Diadical character, characterised as the vertical singlet-triplet energy gap, $\Delta E_{ST} = E(T_1) - E(S_0)$, is summarized in Table 4.5, together with the occupation of frontier natural orbitals, HONO and LUNO, obtained by means of RAS-SF calculations. As predicted by the Hückel model, ΔE_{ST} diminishes from shorter to longer acenes. The occupation of HONO decreases by transferring population to the LUNO (and other higher virtual orbitals) as the size of oligoacenes increases. From [n]Ac with n = 10 - 12 both orbitals are occupied by nearly 1 electron, which is reflected by the number of unpaired electrons ($N_u = 2.44$, 2.48 and 2.42, respectively). The smooth increase of both LUNO occupation and N_u with the increment of n indicates

the appearance of diradicaloid character. A rough change occurs from nonacene to decacene, beyond this point diradical character is stabilized.

From decacene to dodecacene, they are close to be a triplet ground state with a diradicaloid character high enough to make it highly reactive at standard conditions and prevent their experimental isolation. Actually, from hexacene up to nonacene, linear acenes have not been synthesised yet as free molecules, but as stabilized over metal layers.

System	ΔE_{ST}	HONO	LUNO	N_u
Pentacene	1.275	1.844	0.157	0.62
Hexacene	1.015	1.826	0.175	0.67
Heptacene	0.753	1.772	0.229	0.79
Octacene	0.360	1.598	0.403	1.17
Nonacene	0.237	1.462	0.538	1.46
Decacene	0.108	1.018	0.982	2.44
Undecacene	0.111	1.013	0.988	2.48
Dodecacene	0.120	1.058	0.943	2.42

Table 4.5: Vertical singlet-triplet energy gaps (in eV), natural orbital occupation and unpaired electron numbers (N_u) at the RAS-SF/6-31G(d) level for linear oligoacenes of increasing size.

FOD distribution (graphical representation of N_u) in linear acenes is illustrated Fractional density is localized upon C atoms that are not part of interchain bonds. Mostly in the central benzene rings. This observation is in good agreement with bond length behavior since rings in molecular ends have aromatic-like structure, *i.e.*, are conjugated and with closed-shell character. In the middle of linear acenes interchain distance is maximum and the electron interaction is minimised.

The origin of radical character of linear oligoacens has been exposed. Based on the results here presented is clear the unsuccessful attempts to synthesize and isolate bigger acenes results from their high diradical character.

4.2.2 Cyclic acenes

Fundamental electronic differences between linear and cyclic acenes induced by topology were exposed by the Hücklel model. While frontier molecular orbitals in linear acenes display molecular orbital distribution with no degenerations with an energy gap between frontier orbitals that decreases with n, the orbital distribution of [n]CC is different if the number of fused rings is odd or even.

Analogous to linear compounds, only carbon atoms localized in the vertices of the zig-zag edges contribute to the fractional occupation density, as illustrated in -Figure X-. In contrast to linear acenes, in [n]CC the FOD distribution is uniform over upper and bottom chains, this is in good agreement with the higher molecular symmetry. The characterization of the radical character of the ground state is shown in Table 4.6. With the increase of the molecular size the ΔE_{ST} decreases and N_u increases. In Figure 4.20 both, ΔE_{ST} and N_u are plotted as a function of the number of fused benzene rings, exhibiting the electronic differences between odd and even- $[\mathbf{n}]$ CC.

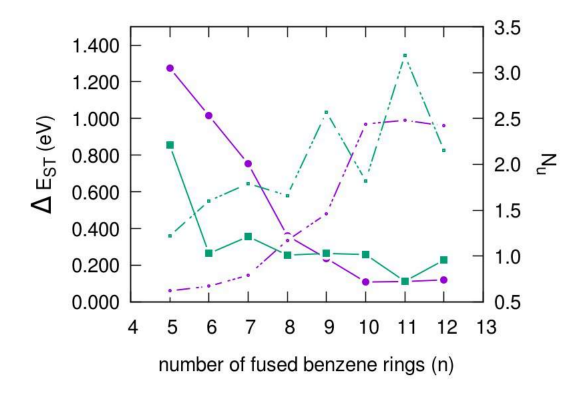


Figure 4.20: Singlet-triplet energy gap (continued lines) and N_u (discontinued lines) of $[\mathbf{n}]\mathbf{CC}$ (green squares) and parent $[\mathbf{n}]\mathbf{Ac}$ (purple circles).

These results point out that [n]CC with n odd and even are part of two separated families. This different behaviour of ΔE_{ST} and N_u for n odd or even can be related to the differences in their Hückel diagram which suggest that their electronic structure is qualitatively different. Actually, when n is odd, N_u tends to four, indicating they are tetraradicaloid systems, as indicated by the doubly degeneracy of the HOMO and LUMO in the Hückel model. Fractional occupation of frontier natural orbitals, from HONO-1 to LUNO+1, shows the simultaneous increment of population in LUNO and LUNO+1, while HONO and HONO-1 are depopulated. Another way to describe the tetraradicaloid character of these systems is by the energy gap between singlet ground state and lowest quintet state, $\Delta E_{SQ} = E_{Q_1} - E_{S_0}$.

The occupation numbers in [n]CC with even number of rings (HONO ~ 1.4 and LUNO ~ 0.6) and N_u (> 1.5) indicates strong radical ground state charac-

$\overline{[n]CC}$	ΔE_{ST}	ΔE_{SQ}	HONO-1	HONO	LUNO	LUNO+1	N_u
5	0.854	2.068	1.778	1.731	0.266	0.216	1.22
6	0.264	2.000	1.902	1.751 1.457	0.260 0.542	0.102	1.60
7	0.356	0.842	1.627	1.626	0.369	0.368	1.79
8	0.255	_	1.880	1.462	0.542	0.119	1.66
9	0.164	0.399	1.449	1.449	0.549	0.549	2.57
10	0.258	-	1.841	1.435	0.570	0.158	1.82
11	0.112	0.284	1.317	1.310	0.689	0.682	3.19
12	0.228	-	1.813	1.340	0.664	0.186	2.15

Table 4.6: Vertical singlet-triplet and singlet-quintet energy gaps (in eV), natural orbital occupation and unpaired electron number (N_u) at the RAS-SF/6-31G(d) level for [n]CC.

ter.Interestingly, [n]CC, n = 6, 8, 10 and 12 present a moderate decrease of ΔE_{ST} , from [6]CC = 0.264 eV to [12]CC = 0.228 eV, which means that the diradical character is increasing slowly with n.

Actually, the electronic structure of cyclic acenes with even number of benzene rings diverge from what is anticipated by the Hückel model, the degeneracy of the energy of HONO and LUNO is not observed. Similarly to observed in previous section, the symmetry of the Hückel Hamiltonian is higher than molecular symmetry. For example, HONO and LUNO of [8]CC (Figure 4.21 a) correspond to the B_{2g} and B_{2u} irreducible representations of the D_{8h} point group and are predicted to be degenerated by the Hückel model. In electronic structure calculations, when not only interactions with first neighbours are considered, this orbital degeneracy is lifted.

Inspection of frontier natural orbitals in Figure 4.21 reveals the opposite interaction between up and down edges when odd- or even-[n]CC. When n = 5, 7, 9 and 11 the HONOs (LUNOs) corresponds to out-of-phase (in-phase) arrangement between π -orbitals, which also is observed in linear acenes. On the contrary HONO (LUNO) of n-even [n]CC holds bonding (anti-bonding) interaction between unpaired electrons in top and down edges. It is worth noticing that NOs, as the FOD, are mostly localized on the carbons belonging to vertices of zizag pattern (C-bridging atoms are barely involved). This bonding (anti-bonding) interaction between up and down edges can be added in the Hückel Hamiltonian (γ in Figure 4.22), as a result the HOMO-LUMO gap decreases in n-odd cyclacenes. On the other hand, orbital degeneracy previously observed in even-[n]CC is lifted, and the HOMO-LUMO gap appears. These results are in good agreement with their rapid increment of tetraradicaloid character of odd-[n]CC and explains why even-[n]CC are diradicaloids but not perfect diradicals.

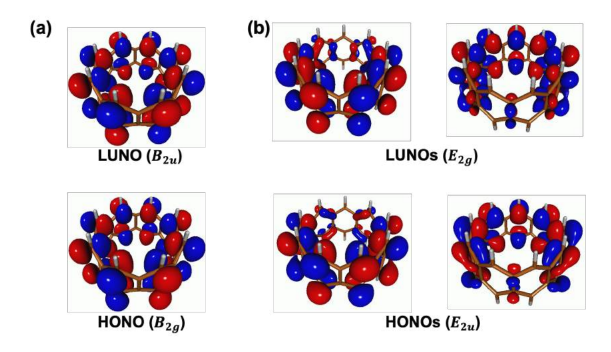


Figure 4.21: Frontier natural orbitals of [8]CC (a) and [9]CC (b).

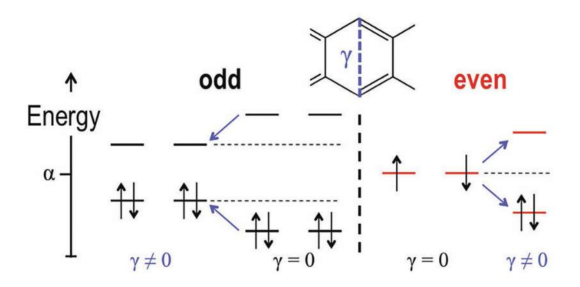


Figure 4.22: Effect of radical center interaction (γ) in the Hückel frontier orbitals for [n]CC with n-odd (left) and n-even (right).

Alternatively, the interaction between radical centers can be rationalised by the contribution of the ionic configuration to the wave function of the ground state. In a simple model, two electrons in two orbitals, where one electron (radical) is localized over each ring, up or down edge in [n]CC. The ability of two radical

centers to share electron density is measured by the ionic contributions that help to stabilize closed-shell configurations. As a consequence, strong interaction results in large HOMO-LUMO energy gaps and the diminution of the radical character. This inter-radical interaction can be described by deconstruction of the ground-state wave function of [n]CC, when n is even, in a fragment decomposition scheme (Subsection 3.3.2.1). Defined fragments are both radical cycles, up (ϕ_u) and down (ϕ_d) . The wave function of the full system in terms of covalent (cov) and ionic (ion) contributions is expressed as:

$$|\Psi\rangle = C_{cov}|\phi_u^{(1)}\phi_d^{(1)}\rangle + C_{ion}(|\phi_u^{(2)}\phi_d^{(0)}\rangle + |\phi_u^{(0)}\phi_d^{(2)}\rangle), \tag{4.2}$$

where (m) superindex indicates the number of electrons in each ring. The correlation between ionic contribution to ground-state wave function and ΔE_{ST} of [n]CC, n = 6, 8, 10 and 12 is displayed in Figure 4.23. In the simple model used here the values in the graph have to be considered only qualitatively since fragment decomposition has been done with a small active space and are not the same as reported in Table 4.6. Nevertheless, the correlation indicates the weak interaction between radical centers with ionic contributions to the wave function less than 4%. It is noticeable that the presence of radical-radical interaction between centers (γ in Hückel model) controls and tunes the diradical character.

To conclude, the relation between the crescent strength of the diradical character with the size of cyclic and lineal acenes and its dependence on the topology of both families was exposed here. Cyclic and longer lineal acenes constitute a challenging synthetic effort, while systems as tetracene, pentacene and hexacene are easy to functionalize and are being routinely exploited in many fields. Detailed analysis of the ground state of [n]CC allows us to unveil their radical nature is different if are constituted by an odd or even number of fused benzene rings. Even-n cyclacenes are diradicaloids, while odd ones can be regarded as tetraradicaloids.

4.3 Appendix

4.3.1 Computational details

Molecular geometries have been optimized in the framework of the density functional theory (DFT) with the M06-2X exchange-correlation functional. Ground and low-lying excited electronic states have been computed by means of the restricted active space configuration interaction (RASCI)(? 35) with the *hole* and *electron* approximation and the use of a spin-flip excitation operator. All π -orbitals were included in the calculation while core electrons orbitals were frozen, in this way excitations from σ -orbitals are not considered.

Radical and poly-radical character of electronic states have been evaluated through the representation of natural orbitals and the fractional occupation density. FODs

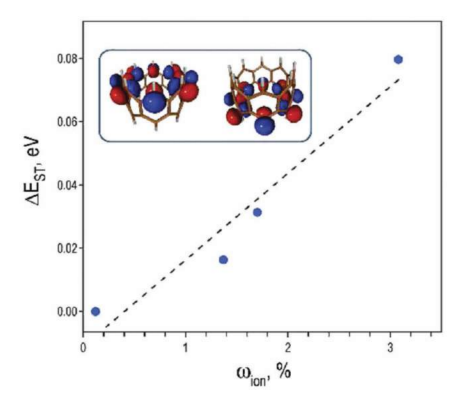


Figure 4.23: Correlation between ionic contributions ($\omega_{ion} = 100 \times |c_{ion}|^2$) and singlet–triplet energy gaps for the RAS-SF[2,2] ground state wave functions of [n]CC molecules with n-even. Inset shows fragment orbitals for the case with n = 8. Dashed line corresponds to the regression line only plotted for guidance.

have been plotted with a modified version of Avogrado (Version 1.2).(36) The number of unpaired electrons has been quantified by the Head-Gordon (N_u) metrics.(37) All calculations have been performed with the Q-Chem package.(38)

4.3.1.1 Triangular graphene nanofragments

All calculations for these systems were performed with the 6-311G(d,p) basis set. Detailed description of the orbital spaces employed for each system can be found in Table 4.7. Doped 1 and 2 derivatives have the same set of active orbitals in the RAS2 space than the pristine counterpart.

TGNF	Symmetry	Spin	(Active) RAS2	(Active) RAS2
		reference	electrons	orbitals
1	D_{3h}	Doublet	7	$2e'', 2a_2'', 1a_1'', 3e'' \text{ and } 3a_2''$
$\mathbf{1N}_1$	D_{3h}	Triplet	8	$2e''$, $2a_2''$, $1a_1''$, $3e''$ and $3a_2''$
$\mathbf{1B}_1$	D_{3h}	Triplet	6	$2e''$, $2a_2''$, $1a_1''$, $3a_2''$ and $3e''$
$\mathbf{1H}_3$	D_{3h}	Quintet	8	$2e''$, $1a_1''$, $4a_2''$, $3e''$ and $4e''$
2	D_{3h}	Triplet	10	$3e''$, $1a_1''$, $3a_2''$, $4e''$, $4a_2''$, $5e''$ and $2a_1''$
$\mathbf{2B}_1$	D_{3h}/C_{2v}	Quartet	3	$3a_2'', 4e'' \text{ and } 4a_2''$
$\mathbf{2B}_3$	D_{3h}	Quartet	7	$3e''$, $1a_1''$, $3a_2''$, $4e''$, $4a_2''$, $5e''$ and $2a_1''$
$\mathbf{2N}_1$	D_{3h}/C_{c2v}	Quartet	5	$3a_2'', 4e'' \text{ and } 4a_2''$
$\mathbf{2N}_3$	D_{3h}	Quartet	13	$3e''$, $1a_1''$, $3a_2''$, $4e''$, $4a_2''$, $5e''$ and $2a_1''$
$2\mathbf{H}_3a$	D_{3h}	Sextet	9	$3a_2'', 1a_1', 4e'', 5e'', 4a_2'', 2a_1'' $ and $5a_2''$
$2\mathbf{H}_3b$	D_{3h}	Doublet	5	$3e'', 3a_2'' \text{ and } 3e''$
$2N_2$	C_{2v}	Singlet	4	$3a''2, 4e'' \text{ and } 4a_2''$

Table 4.7: Active space parameter employed in the RAS-SF calculations.

4.3.1.2 Cyclic and linear acenes

Study over oligoacenes was carried out together with the theoretical group leaded by Prof. Juan Carlos Sancho-Garcia from the University of Alicante.

RAS-SF calculations were performed over a set of geometries provided by the group of Prof. Sancho-Garcia (optimized and confirmed as a minimum in the ground state PES at M06-2X/6-31+G* level, using Gaussian09 package (?)). The active subspace of RAS-SF included 8 electrons in 8 π -orbitals, and the 6-31G(d) was used in all calculations. The high-spin reference was the triplet ROHF for even-[n]CC and linear acenes. For [n]CC with odd n the lowest ROHF quintet was employed instead. Fragment decomposition of the RAS-SF ground state function of even [n]CC molecules have been done considering a RAS2 subspace with 2 electrons in 2 π -orbitals. Molecular orbital energies for the Hückel model were obtained with $\alpha < 0$, $\beta = \alpha/10$ and $\gamma = \beta/5$ for the Coulomb and resonance integrals.

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Chapter 5

Anionic and cationic oligomer diradicals

Conjugated organic polymers have been extensively investigated because of their promising magnetic, optical and electronic properties for applications in solar cells, charge carriers, energy storage, bioelectronics, or chemical sensors.(1; 2; 3? ?) Polyvinylenes, polythiophenes, poly-p-phenylene, polyfluorene, polypyrrole, among others(4; 3; 5) are paramount examples of conjugated polymers. They exhibit electronic resonance between aromatic and quinoidal structures, as Figure 5.1 shows.

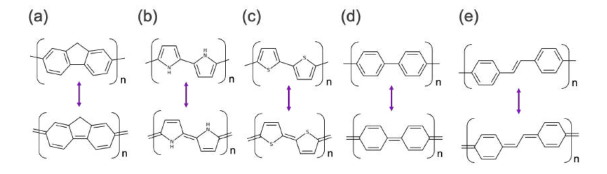


Figure 5.1: Polyfluorene (a), polypyrrole (b), polythiophene (c), poly-p-phenylene (d), polyvinylene (e) in aromatic (top) and quinoidal (bottom) structures.

A typical strategy to improve their electronic and magnetic properties consists in substituting the polymeric chains with electron donor and withdrawing groups. For instance, n-type (electron acceptor) and p-type (electron donor) polythiophenes have proved their excellent performance for decades (6; 7; 8) in organic field-effect transistors(?), organic light-emitting diodes, photovoltaic devices, batteries, solar cells(9), etc. (10) and catalyst to solar energy utilization(11), that can be modulated by structural modifications, e.g., functionalization of 3,4 positions, introduction of a copolymer between thiophene units.

Modelling of the optoelectronic properties of polymers is commonly addressed

using the so-called oligomer approach, which relies on the extrapolation of the properties of finite length oligomers to the polymer limit. From a fundamental point of view, oligomers provide accurate structure-property relationships that are beneficial for the knowledge and understanding of the photomechanisms of operation in polymers and for unraveling the design rules for the elaboration of new materials with enhanced functionalities.

Stable organic oligomers with closed-shell (CS) singlet ground state, *i.e.*, large HOMO-LUMO energy gap, can be transformed into diradicals by adding or removing a pair of electrons. Double oxidation of the system may lead to the formation of a dication-diradical. If the HOMO to HOMO-1 energy gap is rather narrow, as a result of removing two electrons a pair of HOMO and LUMO nearly degenerate can share electronic population. Similarly, if the LUMO and LUMO+1 energy gap is small enough, the double reduction can lead to the partial occupancy of both HOMO and LUMO orbitals, on the formed dianion. Thus a di-ionic diradicaloid species appears (Figure 5.2).

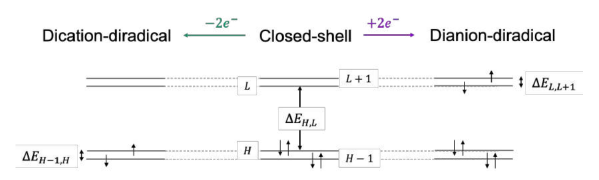


Figure 5.2: Dication (left) and diamon (right) diradical formation from a closed-shell structure (center) by double oxidation and reduction, respectively.

The present chapter is devoted to the exhaustive description of two organic diradical oligomers. The main goal is to unravel the origin of their diradical nature and control their structural, photophysical, and spectroscopic features. In the first part, oligothiophene derivatives will be investigated. In the second part the equilibrium between aromatic and quinoidal structures (and the associated variation of radical character) of a substituted poly-p-phenylene oligomer will be described in detail.

5.1 Thieno[3,4-c]pyrrole-4,6-dione oligothiophene

Thiophene based π -functional materials have been intensively investigated for their application in organic (opto)electronic devices. Recently, Zhu et al. developed a new low-bandgap copolymer combining the n-type (electron acceptor) thieno[3,4-c]pyrrole-4,6-dione oligothiophenes (**TPD**) units with the quinoid-enhancing thieno[3

,4-b]thiophene (TbT) moiety and a weakly electron-donating benzo[1,2-b:4,5b'] dithiophene (BDT),(12) which was successfully applied to polymeric solar cells demonstrating a power conversion efficiency over 9%.(??) In order to investigate the nature of TPD as an acceptor moitey, a new series of thieno[3,4-c]pyrrole-4,6-dione oligothiophenes, \mathbf{OTPD}_n (n=2-6, Figure 5.3), were synthesized and characterized spectroscopically, and found to be able to stabilize both anions and dianions in their reduced forms. The properties of anions and dianions of oligothiophenes have been scarcely investigated, since oligothiophenes are better suited for the stabilization of positive charges than negative ones. The study of \mathbf{OTPD}_n oligomers therefore might represent a significant step towards understanding the oligothiophene anionic species.

In the following, I present a comprehensive computational investigation of the main electronic features of \mathbf{OTPD}_n oligomers in their neutral, anionic and dianionic forms. Interestingly, these oligomers display a clear opposition between the peripheral dione groups and the central oligothiophene array, which establish two separated electronic domains: one among the dione units and one among the thiophene moieties, referred to as interdione (IDn) and interthiophene (IT) π -conjugation paths (Figure 5.3), respectively.

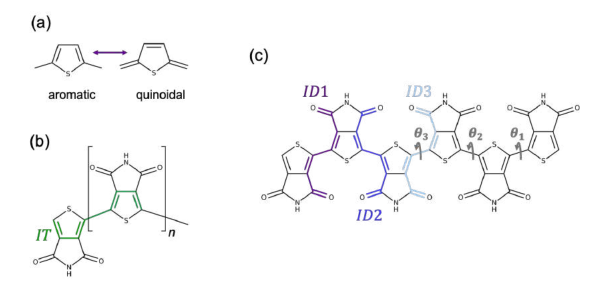


Figure 5.3: Aromatic and quinoidal forms of thiophene (a). Structure of \mathbf{OTPD}_n with n=1-5, interthiophene (*IT*) π -conjugation path indicated in green (b). Interdione (*IDn*) π -conjugation paths and the dihedral angle (θ) between adjacent thiophene units of $[\mathbf{OTPD}_n]$ oligomers (c).

5.1.1 Monoanionic species

The ground state molecular geometry of the $OTPD_n$ presents significant structural differences as a function of n (Table 5.1). Namely, the (quasi-)planarity the back-

bone, measured as the dihedral angle (θ_n) between adjacent TPD units as shown in Figure 5.3 (b), decrease as the system size increases, in agreement with what has been reported theoretically and experimentally for other substituted oligothiophenes. (?????) Despite the fluctuation of θ , conjugation along the thiophene backbone is not affected, as illustrated by the evolution of the bond length alternation of the IT path (BLA_{IT} in Table 5.1), for which the largest decrease is found from the dimer to the trimer. Such change is associated with the downshift of the band assigned to the collective stretching of C-C bonds in the Raman spectra(????) and can be understood as the result of the slight increase of the π -conjugation from $OTPD_2 \rightarrow OTPD_6$. On the other hand, the BLA of the interdione conjugation path does not vary with the length of the oligomer.

species	BLA_{IT}	BLA_{ID1}	BLA_{ID2}	BLA_{ID3}	θ_1	θ_2	θ_3
$\overline{\textbf{OTPD}_2}$	0.063	0.092			179.9		
-			-	-		-	-
$[\mathbf{OTPD}_2]^{-1}$	0.031	0.023	-	-	180.0	-	-
$[\mathbf{OTPD}_2]^{-2}$	-0.003	-0.061	-	-	170.7	-	-
OTPD_3	0.052	0.091	-	-	171.2	_	-
$[\mathbf{OTPD}_3]^{-1}$	0.028	0.048	-	-	171.2	-	-
$[\mathbf{OTPD}_3]^{-2}$	0.017	0.011	-	-	165.2	-	-
\mathbf{OTPD}_4	0.055	0.091	0.094	-	171.7	155.1	-
$[\mathbf{OTPD}_4]^{-1}$	0.029	0.065	0.049	-	171.8	164.0	-
$[\mathbf{OTPD}_4]^{-2}$	0.025	0.021	0.070	-	169.1	153.9	-
\mathbf{OTPD}_5	0.050	0.091	0.092	-	172.1	158.0	_
$[\mathbf{OTPD}_5]^{-1}$	0.029	0.073	0.057	-	171.8	165.4	-
$[\mathbf{OTPD}_5]^{-2}$	0.026	0.027	0.071	-	170.4	155.8	-
\mathbf{OTPD}_6	0.051	0.092	0.094	0.088	170.1	153.7	180.0
$[\mathbf{OTPD}_6]^{-1}$	0.034	0.078	0.065	0.060	171.1	166.1	178.8
$[\mathbf{OTPD}_6]^{-2}$	0.025	0.038	0.061	0.074	169.8	159.9	179.8

Table 5.1: Bond length alternation (in Å) of π -conjugation paths IT, ID1-3 and dihedral angles θ_1 , θ_1 , θ_3 as described in Figure 5.3.

Aromatic/quinoidal forms of the five-member thiophene rings (Figure 5.3 a) can be described by the double-single bond alternation of the C-C bonds in the ring, where C=C-C=C (C-C=C-C) alternation pattern corresponds to the aromatic (quinoidal) structure. By plotting the bond lengths of the π backbone, aromatic and quinoidal thiophenes are represented by the three C-C bond lengths within

each thiophene unit connected by a line, a Λ -shape line corresponds to an aromatic ring while a V-shape is indicative of a quinoidal structure. Figure 5.4 shows the bond length distribution of the π -conjugated backbone of \mathbf{OTPD}_6 , where thiophene units have almost equivalent aromatic rings connected by C-C bonds.

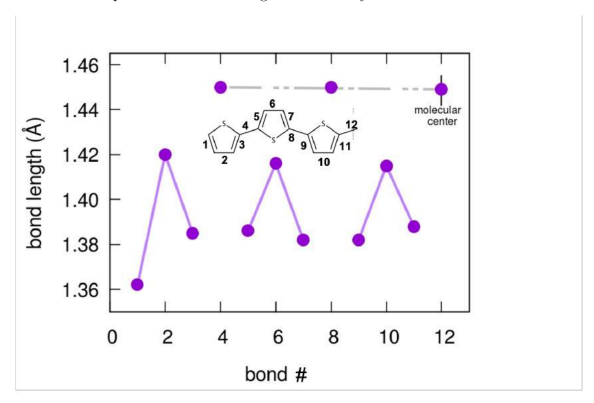


Figure 5.4: C-C bond lengths along the IT path of \mathbf{OTPD}_6 . Purple lines connect the three C-C bond lengths within each thiophene ring, while the dashed gray line connects the interthiophene bond lengths.

All \mathbf{OTPD}_n oligomers exhibit a large HOMO-LUMO energy gap in their singlet ground state (Table 5.2), illustrating their closed-shell character. The large adiabatic energy gap to the lowest triplet further supports the closed-shell nature of the ground state.

By means of cyclic voltammetry measurements the consecutive formation of the anion and dianion is observed with a very weak size-dependency of the reduction potentials, from -1.65 and -1.85 eV in the trimer to -1.38 and -1.55 eV in the hexamer (Figure 4 in reference (???)). This behaviour suggests that the dione group plays a main role in the stabilization of the injected charge by resonance between adjacent TPD units (IDn paths on Figure 5.3). It is noteworthy that the LUMO-LUMO+1 energy difference is rather small, suggesting a potential diradical character in the dianionic species.

Single reduction of \mathbf{OTPD}_n leads to the formation of the monoanion monoradicals $[\mathbf{OTPD}_n]^{-1}$. The distribution of the excess charge over each thiophene unit is presented in Figure 5.5. The additional electron is shared equitably between

$\overline{ ext{OTPD}_n}$	ΔE_{ST}	$\Delta E_{H,L}$	$\Delta E_{L,L+1}$
n=2	48.1	88.5	20.1
n = 3	42.7	76.6	8.8
n = 4	39.9	70.9	8.8
n = 5	38.9	67.8	7.5
n = 6	38.0	65.3	6.9

Table 5.2: Adiabatic energy gaps in kcal mol⁻¹ between closed-shell singlet and triplet states (ΔE_{ST}). $\Delta E_{H,L}$ ($\Delta E_{L,L+1}$) is the energy difference between HOMO (LUMO) and LUMO (LUMO+1) orbitals in the ground state geometry.

monomers in $[\mathbf{OTPD}_n]^{-1}$ with n=2-4. More specifically, atomic charge analysis for the smallest oligomer suggests the excess of the charge is located over oxygen atoms in dione group, and are in resonance with the dione group in the adjacent thiophene unit through an interdione path (IDn in Figure 5.3 c). As a result, there is an increment of the conjugation over the IDn path, i.e., the BLA_{IDn} decreases. When n=3, the additional electron delocalizes over the two equivalent interdione paths. The presence of another dione group in the extreme thiophene unit explains the asymmetrical charge distribution. In the tetramer, excess of charges in oxygen atoms are in resonance only through the ID path between central units, and the charge is equally distributed. Finally, a moderate charge localization over the molecular center is observed for n=5 and 6.

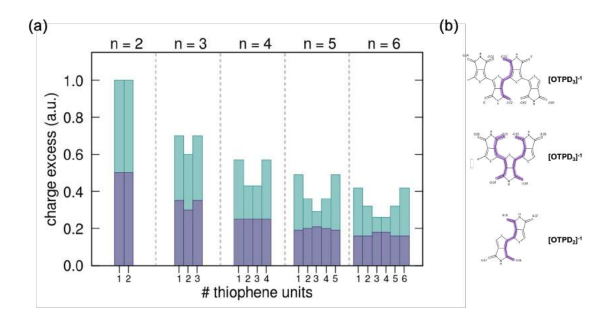


Figure 5.5: Distribution of the Mulliken charge over each thiophene unit in \mathbf{OTPD}_n monoanions (purple bars) and dianions (green bars) (a). The ID path for $[\mathbf{OTPD}n]^{-1}$ (n=2-4) and atomic charge of O atom are illustrated (b).

Structurally, a slight increase of the molecular planarity of the monoanionic oligomers is obtained, with an average dihedral angle (θ_2) of 165° (155° in neutral forms), while θ_1 and θ_3 barely change with respect to the neutral species (see Table 5.1). Structural stabilization of the additional charge translates into significant changes in the molecular bond lengths. Namely, BLA_{IT} of $[\text{OTPD}n]^{-1}$ considerably decreases as compared to the neutral form. The BLA_{ID} of monoanions also decrease, but such reduction strongly depends on the oligomer size. Interestingly, the central units are the ones showing the smaller value of BLA_{ID} , meaning that the bond lengths along this path are more homogeneous and the π -conjugation increases by delocalization of additional charge. This result is in good agreement with what was observed by cyclic voltammetry, *i.e.*, that the charge is mainly stabilized by the dione group and has a weak dependence on the oligomer length.

A more detailed picture can be seen from the bond length distribution along the IT path as shown for $[\mathbf{OTDP_6}]^{-1}$ in Figure 5.6. While thiophene units at the molecular ends are aromatic, this character gradually decreases towards the central rings, along with a decrease of the interring distance. Comparison to the neutral $\mathbf{OTDP_6}$ (Figure 5.4) illustrates that the main geometrical effect of adding an electron is the increment of the quinoidal character at the molecular center.

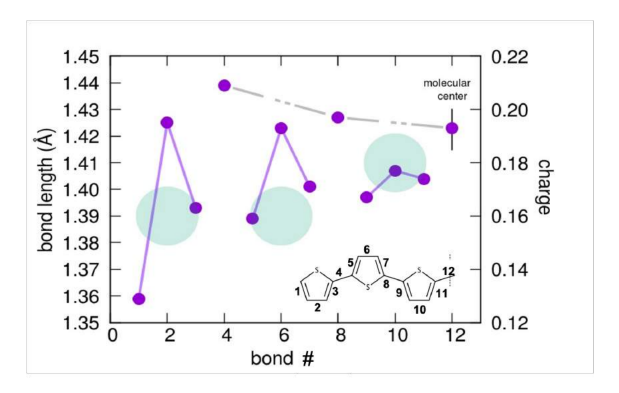


Figure 5.6: C-C bond lengths in the IT path of $[OTDP_6]^{-1}$. Purple lines connect the three C-C bond lengths within each thiophene ring, while the dashed gray line connects the interthiophene bond lengths.

All these features are a clear indication that charge stabilization leads to a dis-

tortion of the geometry that confines the unpaired electron (the additional charge). In 1985 Bredas (13) characterized this kind of charge localization in organic polymers and defined it as a polaron (as in solid materials): a radical-ion associated with a lattice distortion. Since then, the formation of polarons has been mainly described in cationic oligomers, but it has been theoretically proven that monocations and monoanions of pristine oligothiophenes localize the charge in very similar manners.(? ? ? ?) In anion-radicals $[OTPD_n]^{-1}$ (up to 6 units at least) a polaron is localized over the molecular center.

5.1.2 Dianionic species: Bipolaron vs polaron-pair

Following the reduction of $[OTPD_n]^{-1}$ to $[OTPD_n]^{-2}$ we envision two possible situations: a) a bipolaron, in which two negative charges are confined together and associated with a single but strong geometrical distortion, and a closed-shell like structure; or b) a polaron-pair, as a result of the formation of two interacting polarons (localized radical-ion) leading to a diradical-dianionic species.(13)

The charge distribution over TPD units for n=2 and 3 is simply twice the one observed in the monoanions, as a consequence of their small size. For larger dianionic oligothiophenes, namely $[OTPD_{4-6}]^{-2}$, both charges are symmetrically localized on the external units of the oligomer, as presented in Figure 5.5 (a).

Geometrical distortions of the dianions are summarized in Table 5.1. To offer a better interpretation of the results, I will discuss first the oligomers with n=4-6. In these larger oligothiophenes, the stabilization of a second electron reverts the planarization gained (θ_2) in the monoanionic structures, returning to the distortion exhibited by the neutral parents. BLA_{IT} is not significantly affected by the reduction of $[\mathbf{OTPD}_{4-6}]^{-1}$, supporting the idea that the thiophene π -backbone plays a minor role in the charge stabilization. Interestingly, BLA_{ID2-3} increase, meaning that in these paths the π -conjugation is lower than in monoanions, but the opposite effect is observed across ID1, where conjugation significantly increases with a very weak dependence on the size of the system. These results indicate two equivalent geometrical deformations localized in each extreme of the molecule.

A better insight on the role of the interthiophene path is provided by the evolution of the C-C bond lengths of $[OTPD_6]^{-2}$ π -backbone (Figure 5.7). Thiophene rings at the extremities clearly exhibit the pattern classically associated to aromatic thiophene-rings, while the second thiophene unit shows a more quinoidal-like structure, before a partial recovery of aromaticity is found in the central thiophene ring. Accordingly, the interring bond length becomes larger towards the molecular center (more single bond character).

On the other hand, $[OTPD_{2-3}]^{-2}$ are very challenging systems whose small size leads to a severe structural deformation driven by the addition of two electrons to the system.

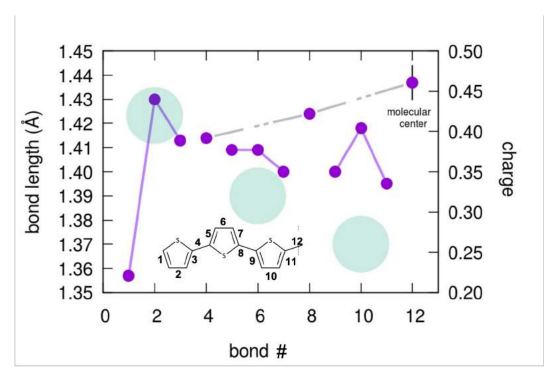


Figure 5.7: C-C bond lengths in the IT path of $[OTDP_6]^{-2}$. Purple lines connect the three C-C bond lengths within each thiophene ring, while dashed gray line connects the interthiophene bond lengths.

The small LUMO to LUMO+1 energy gap of the neutral oligomers suggests the formation of a dianion diradical by addition of two electrons. Indeed, the small $\Delta E_{L,L+1}$ in neutral systems allows the partial population of LUMO+1 when two electrons are added to the system, which induces the formation of two (quasi)degenerated HOMOs in $[OTPD_n]^{-2}$, see Table 5.2. Radical characterisation of oligothiophene dianions is presented in Table 5.3. The adiabatic energy difference between open and closed-shell solutions (ΔE_{OS-CS}^{ad}) shows that all systems hold an open-shell (OS) singlet ground state. When n=2, $\Delta E_{OS-CS}^{ad}=0.04~{\rm kcal~mol^{-1}}$ and ${\bf [OTPD_2]^{-2}}$ must be considered a closed-shell system. On the other hand, when n=6, the CS solution is destabilized by 4.2 kcal mol⁻¹ with respect to the OS one. This is reaffirmed by the adiabatic and vertical energy gaps between the open-shell singlet and the lowest triplet state, both decreasing substantially from n=2 to n=4. For n = 4 - 6, ΔE_{OS-T}^{ad} remains nearly constant. Based on these trends, it can be deduced that the lengthening of the oligomer will not result in an increase of the radical character (beyond the diradical). In fact, the quintet state, which represents an ideal tetraradical (4 unpaired electrons), is rather high in energy. Accordingly, the number of unpaired electrons gradually approximates to 2 (n = 4 - 6), meaning that they are not perfect diradicals, but can be appropriately named diradicaloids. Indeed, natural orbitals shown in Figure 5.8 evidence the presence of two weakly interacting radical moieties located over each molecular end, with electronic occupancy numbers approaching to 1 as the molecular length increases.

$\overline{[\mathbf{OTPD}_n]^{-2}}$	ΔE_{OS-CS}^{ad}	ΔE_{OS-T}^{ad}	$\Delta E_{OS-T}^{ad}{}^{[a]}$	$N_u^{[a]}$	ΔE_{OS-Q}^{vert}	ΔE_{int}
n=2	~ 0.0	13.6	21.3	0.47	77.0	92.04
n = 3	1.1	1.5	1.2	0.86	62.9	77.35
n=4	2.2	0.4	0.7	1.61	48.4	66.72
n = 5	3.5	0.3	0.5	1.80	46.5	58.50
n = 6	4.2	0.4	0.3	1.84	44.4	52.22

Table 5.3: Adiabatic (ad) and vertical (vert) energy gaps in kcal mol⁻¹. ^[a] Computed at RAS-SF level. ΔE_{int} is the energy difference between the dianion and twice the monoanion.

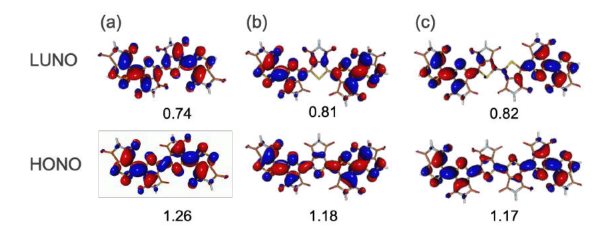


Figure 5.8: Frontier natural orbitals with their electronic occupation numbers for $[\mathbf{OTPD_4}]^{-2}$ (a), $[\mathbf{OTPD_5}]^{-2}$ (b), and $[\mathbf{OTPD_6}]^{-2}$ (c).

The interaction energy of the two radicals can be approximated as $\Delta E_{int} = \Delta E_{di}^{ad} - \Delta E_{2mono}^{ad}$, where the first term in the right side of the equation is the difference between the dianion and the neutral compounds ($\Delta E_{di}^{ad} = E_D^{ad} - E_N^{ad}$), i.e., the increment of energy from adding two electrons to the system. On the other hand, ΔE_{2mono}^{ad} is twice the energy gap between monoanion and neutral compounds ($\Delta E_{2mono}^{ad} = 2(E_M^{ad} - E_N^{ad})$). Thus, ΔE_{int} approximates the difference of having either a diradicaloid or two "independent" radicals (twice a radical). As n increases, ΔE_{int} decreases, reproducing the dependence on the distance of the classical repulsion between two negative charges $(1/n \propto 1/R$, Figure 5.9), in good agreement with the picture exposed by the natural orbitals.

Therefore, we conclude that the electronic structure of $[OTPD_{4-6}]^{-2}$ can be described as a polaron-pair with two favorable conjugation routes: the interdione path which stabilizes the total charge and the interthiophene-interdione path which modulates the diradical character. The small size and closed-shell nature of $[OTPD_{2-3}]^{-2}$

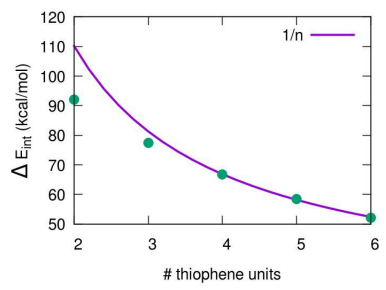


Figure 5.9: Adiabatic energy difference ΔE (kcal/mol) between dianion and twice the monoanion for n=2-6. Purple line corresponds to the energy adjusted to a 1/n for $4 \le n \le 6$.

is not appropriate to classify them as bipolaron or polaron-pair.

5.1.3 Spectroscopic features

So far, the appearance of polaron, bipolaron, and polaron-pair derived from the consecutive reduction of oligothiopenes have been described here by means of the analysis of geometrical and electronic features. There is a considerable lack of information about the experimental characterization of n-doped oligothiophenes or their polaronic species with only few reports in the literature. Salzner et al. (? ? ? ?) carried out a computational study to investigate the spectroscopic properties of a series of monoanionic and monocationic pristine thiophene oligomers concluding that the spectra of p- or n-doped species are virtually identical.

UV-VIS/NIR spectra of neutral, monoanion and dianion \mathbf{OPTD}_n have been measured experimentally (Figure 5 in reference (???)) and interpreted by means of TD-DFT calculations presented hereafter, the most relevant data being collected in Table 5.4. In neutral oligothiophenes, the maximum of the main band of the absorption spectra is continuously redshifted from colorless (dimer) to strongly yellow-orange (hexamer) which is similar to the behaviour observed in unsubstituted thiophene oligomers.

The measured bathochromic shift in neutral oligothiophenes when increasing n is nicely reproduced by our calculations with a gradual shift of the $S_0 \to S_1$ vertical transition as n increases, from 347 nm for n=2 to 475 nm for n=6. This transition is characterized by the electronic excitation from HOMO to LUMO, in line with what is expected for closed-shell π -conjugated systems.

	$\overline{ ext{OTPD}_n}$	$[\mathbf{OTPD}_n]^{-1}$		[OTP:	$[\mathbf{D}_n]^{-2}$
_n	S_1	S_1	S_2	S_1	S_2
2	347 (0.499)	981 (0.057)	414 (0.447)	-	846 (0.091)
3	402 (0.738)	1784 (0.076)	482 (0.658)	$1635 \ (0.062)$	438 (0.134)
4	441 (1.042)	$1734 \ (0.213)$	537 (0.954)	$1534 \ (0.103)$	601 (0.101)
5	459 (1.276)	$1849 \ (0.365)$	573 (1.196)	1879 (0.188)	542 (0.484)
6	475 (1.612)	$1975 \ (0.567)$	596 (1.432)	2089 (0.344)	557 (0.691)

Table 5.4: Wavelength (in nm) of the two lowest singlet-singlet vertical excitations for \mathbf{OTPD}_n , $[\mathbf{OTPD}_n]^{-1}$ and $[\mathbf{OTPD}_n]^{-2}$ (n = 2-6) computed at the B3LYP/6-31G(d,p) level. Values in parenthesis correspond to oscillator strengths.

Upon reduction of the \mathbf{OTPD}_{3-6} oligomers, the absorption spectra exhibit two important bands due to their radical nature, namely a main absorption in the visible and a second broad weak one in the infrared. This is the typical two-bands pattern for the open-shell doublet of anions and cations of linear π -conjugated oligomers.(?) This is fully supported by our calculations, which characterize the bands of these open-shell anion radicals as two excitations that involve the doubly occupied, semi-occupied, and empty frontier molecular orbitals. The first band corresponds to the HOMO \rightarrow LUMO transition, as in neutral compounds, but red-shifted from 347 nm to 414 nm when n=2, and from 475 nm to 596 nm for n=6. The second one appears as a broad band in the infrared that involves the excitations SOMO \rightarrow LUMO and HOMO \rightarrow SOMO, with a transition intensity (approximated by the oscillator strength) that increases with the system size.

Similarly, UV-VIS spectra of dianions show a two-band profile. $[\mathbf{OPTD}_{n=4-6}]^{-2}$ experimental band maxima were characterized at 580/1537 nm (n=4), 569/1590 nm (n=5), and 568/1707 nm (n=6), which are blueshifted to those of their respective anion radicals. This is typical of dicationic π -conjugated molecules in which the dication discloses a polaron-pair structure,[32] while their dianions are typical closed-shell systems that exhibit a single absorption band in the optical spectra.[30] This two-band pattern in the absorption spectra of dianions of π -conjugated molecules has been reported in the case of oligofluorenes[31] and assigned to the presence and stabilization of segregated polarons. The separation of the bipolaron structure into two symmetrical polarons (polaron-pair) might justify the similitude of the spectra of radical anions and polaron-pair dianions in $[\mathbf{OTPD}n]^{-2}$ compounds.

TD-DFT results identify the main absorption band as the $\pi - \pi^*$ transition from the highest doubly occupied orbital to the lowest totally empty orbital. This band resembles to the one in the monoanionic species although is slightly blue-shifted. The low-energy broad band associated with the presence of radical species is located in the IR region, slightly red-shifted with respect to the low-energy band

of the monoanions, and arises from an excitation that involves the two SOMOs. This band is the main spectroscopic characteristic of polarons, inline with the polaron-pair characterization of $[OTPD_{4-6}]^{-2}$.

Figure 5.10 shows the simulated absorption spectrum of $[OTPD_6]^{-2}$, with assignement of the polaron-pair band and the molecular orbitals involved, as well as the $\pi - \pi^*$ transition. In contrast, the smaller members (n = 2 and n = 3) just exhibit one absorption band (HOMO-to-LUMO) due to its closed-shell nature.

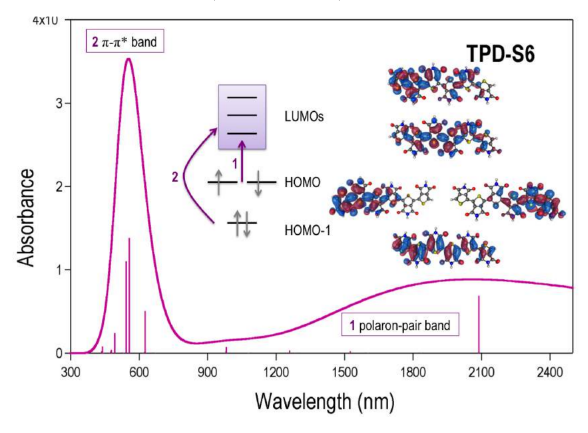


Figure 5.10: Absorption spectrum for the dianion $[\mathbf{OTPD}_6]^{-2}$. Main orbital-to-orbital transitions are shown in the inset

An important result of this study was the segmentation of the electronic structure of these \mathbf{OTPD}_n oligomers, which provoked the stabilization of dianions as segregated anions or polaron-pair dianions placed in parallel (*i.e.*, in contrast to polaron-pairs in series, as described in oligofluorenes). The existence of two paths of π -conjugation; 1) between neighbor diones and 2) through the thiophene spine was pointed out. They allow interspins and through-bond interaction, which result in the existence of a singlet diradical ground electronic state.

5.2 Diamino Oligophenyl Dication

The Thiele's, (14) Chichibabin's, (15) and Muller's (?) hydrocarbons are the smallest members among the p-phenylene substituted oligomers. They have a singlet-ground

state with a narrow singlet-triplet energy gap, and can be described by a superposition of open-shell (aromatic) and closed-shell (quinoidal) resonance structures (16), Figure 5.11. Typically, they are unstable molecules with no practical application. In order to understand their properties, stable derivatives have been synthesized and extensively studied. (16), doi:10.1021/acs.accounts.7b00229

Figure 5.11: Aromatic (radical) and quinoidal (closed-shell) resonant forms of Thiele's (n = 1), Chichibabin's (n = 2) and Müller's (n = 3) hydrocarbons.

Recently, the formation of a bis(triarylamine) unit in each extreme of the p-phenylen bridge by substitution of the carbon centers with nitrogen atoms, and posterior generation of the dication, has been proved to be an efficient way to obtain stable diradicals analogous to the Thiele's series.(??????) Concretely, the stable 4,4'-di(bis(1,4-methylphenyl)amino)-p-terphenyl dication (named as $\mathbf{1}^{+2}$ in the original work, herein \mathbf{PAT}^{+2}), see Figure 5.12, was synthesized in 2016 by Su et al.(17) Interestingly, its characterization by EPR spectroscopy, SQUID measurement and X-ray crystallography evidenced the equilibrium between two conformational isomers with singlet ground state, but different diradicaloid nature. The equilibrium between the two conformers is temperature dependent, which allows changing the diradical character of the system by heating/cooling. This feature is very promising in the design of stable materials that can "activate" their diradical character by external stimuli.

At low temperature (LT = 123 K), a distortion from planarity in the central ring of the bridge is observed. The associated singlet-triplet energy gap is determined by SQUID magnetic measurements to be 1.14 kcal/mol (400.57 cm⁻¹)(17), suggesting a strong radical character. This LT conformation can thus be referred to as a distorted diradical (**DD-PAT**⁺²). As the temperature is increased to 200 K (HT), a planar biradicaloid conformer is obtained, namely **PB-PAT**⁺². As its name indicates, a larger singlet-triplet energy gap of $\Delta E_{ST} = -2.37 \text{ kcal/mol (-872.84 cm}^{-1})$ is observed (SQUID experiments(17)) together with the full planarization of the terphenyl bridge.

In the following, the detailed description of the electronic structure of PAT^{+2} is presented, and the ddriving force of the interconversion mechanism between the

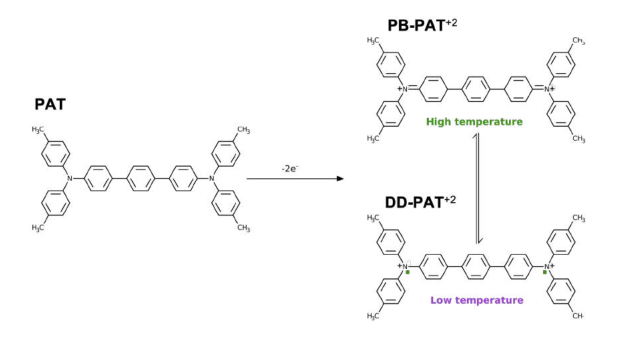


Figure 5.12: Neutral **PAT** with the resonance structures at high and low temperatures of PAT^{+2} .

two configuration that two unveiled.

5.2.1 Origin of the diradical character

PAT holds a closed-shell singlet ground state, in line with the large HOMO to LUMO energy gap (23.55 eV computed at CAM-B3LYP/6-31G(d) level), while the gap between HOMO and HOMO-1 is noticeably smaller (0.91 eV). The removal of two electrons from PAT results in (quasi)degenerated HOMO-LUMO pair of orbitals, which is the origin of the diradical character of PAT⁺². To get a balanced description between the diradical nature and accurate reproduction of experimental observations, PAT⁺² conformers are characterized using DFT, RAS-SF and SF-TDDFT methods.

Small singlet-triplet energy gaps reported in Table 5.5 are in good agreement with experimental measurements. Furthermore, the experimental relative diradical character (OS) is well reproduced as indicated by diradical index (y_0) . **DD-PAT**⁺² has a larger diradical character than **PB-PAT**⁺² (0.84 and 0.65, respectively), still the computed diradical character is significantly high for both conformers. **DD-PAT**⁺² possesses roughly two single-occupied orbitals (HONO occupation number is 1.16), and a large number of unpaired electrons. y_0 and N_u of **PB-PAT**⁺² are indicative of a lower diradical character, due to the moderate fractional occupancy of its orbitals (HONO occupation number is 1.36).

Interestingly, the distorted stronger diradicaloid form (DD-PAT⁺²) is obtained

Structure	SF-TDDFT	RAS-SF	\exp^a	N_u	y_0
$\mathbf{DD} extbf{-}\mathbf{PAT}^{+2}$	1.16	0.49	1.14	1.72	0.84
$\mathbf{PB} ext{-}\mathbf{PAT}^{+2}$	3.88	2.87	2.37	1.36	0.65

Table 5.5: Singlet-triplet energy gap (kcal/mol) for both conformers of PAT^{+2} . Number of unpaired electrons (N_u) and diradical index (y_0) were computed at RAS-SF level. Calculations were performed over the X-ray structures. ^a Experimental values obtained from (????)

at low temperature, while the planarization of the bridge (**PB-PAT**⁺²) is observed at higher temperature. This counterintuitive result indicates that the interconversion process and diradical formation is not entropy-driven. In order to understand the leading force of this transformation, first, the structural parameters of both conformers will be rationalized.

In Table 5.6, the most relevant bond distances of crystal structures are compared with those obtained by the geometrical relaxation at DFT level. Terphenyl moiety has two kinds of nonequivalent benzene rings referred to as A (at the two sides) and B (the central ring). BLA_X is the bond length alternation in each ring. N-Ar is the N-C bond length at the bisphenylamino (BPA) unit, and N-C_{terphenyl} is the distance between the p-terphenyl bridge and the BPA moiety. Overall, computational results agree well with the experimental data. The most important discrepancies are observed in the overestimation of quinoidal-like form of the B ring in **PB-PAT**⁺², as well as in the prediction of a larger θ angle by DFT. These differences can be explained by the absence in the calculations of the structural constraints the crystal environment.

In **PB-PAT**⁺² N-Ar and N-C_{terphenyl} length corresponds to single and double bonds, respectively. While BLA_A and BLA_B are virtually identical in DFT optimized geometries, rings A and B are clearly different in the crystal structure. At LT the N-Ar and N-C_{terphenyl} bond lengths of **DD-PAT**⁺² are quite homogeneous. In addition, both kind of rings possess an aromatic structure and are linked by a single bond (k, which compares well with the standard single bond between two carbons sp² in butadiene). The distortion of the central benzene is slightly overestimated probably due to the absence of the constraints imposed by the crystalline package, but in quite good agreement with the X-ray structure. Molecular length, measured as the distance between N atoms, is shorter in **PB-PAT**⁺² indicating a more quinoidized structure. For the sake of comparison, **PAT** was optimized at the same level of theory as the two dicationic conformers. The ground state minimum presents a distorted structure in which all the rings are rather aromatic (see Table 5.6). Geometrically, it is closer to **DD-PAT**⁺², than **PB-PAT**⁺².

Mulliken analysis for the two forms of PAT⁺² locates the positive excess charge

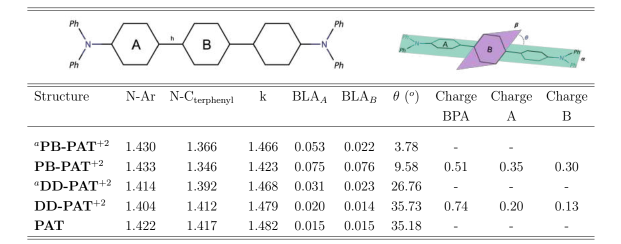


Table 5.6: Main geometrical parameters of **PB-PAT**⁺² and **DD-PAT**⁺² optimized geometries at UCAM-B3LYP/6-31G* level. BLA_A is the bond length alternation of the ring A whereas BLA_B is from ring B. Charges are computed under Mulliken approximation. BLA were calculated as is indicated in the appendix. a X-ray structural data from (?). All bond lengths, Å, are defined in the figure.

mainly on the BPA units, with small contribution over the A rings. This data suggests that the charge and unpaired electron density is stabilized mainly over BPA moieties and towards the A rings of the p-terphenyl bridge, see Table 5.6.

Frontier natural orbitals, Figure 5.13, are mostly localized over central nitrogen atom of the BPA moiety and the side ring of the terphenyl (A). Disruption of the π -conjugation should be attributed to the lost of planarity in the bridge. The interaction between radical centers is larger in **PB-PAT**⁺², as the distortion between A and B rings is smaller than in **DD-PAT**⁺².

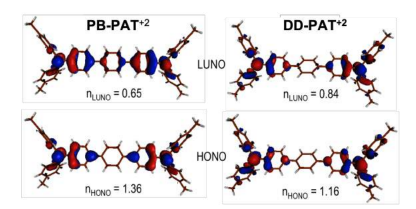


Figure 5.13: Frontier natural orbitals and their electronic occupancy of **PAT**⁺² conformers.

The singlet-ground state of PAT⁺² can be rationalize as a consequence of the antiferromagnetic interaction between radical centers, stronger at high temperature when the system is more planar, giving a moderate diradical character. The radical-

radical interaction at low temperature is weaker, the singlet-triplet gap is rather small and both states, singlet and triplet, are in equilibrium.

Quinoidal/aromatic-like rings, radical character, and charge distribution together offer a complete picture of the electronic structure nature of each conformer of \mathbf{PAT}^{+2} and explain their fundamental differences.

5.2.2 The PB-PAT $^{+2} \leftrightarrow \text{DD-PAT}^{+2}$ conversion

In order to explain the stability of both PAT⁺² conformers, it is important to take into account the steric strain within the phenyl rings of the p-terphenyl unit. Adiabatic singlet-triplet energy gaps at distorted and planar analogs with smaller oligopehnylene, p-phenyl (N=1) and p-biphenyl (N=2), were used to analyze the stability provided by the aromatic/quinoidal-like conjugation through the bridge. Adiabatic singlet-triplet energy gap when N = 1 shows the stabilization of the closed-shell (quinoidal-like) form over triplet conformation (aromatic structure) by 2.03 kcal/mol. By addition of one more ring, p-biphenyl bridge can adopt the planar and distorted form in both, aromatic (triplet) and quinoidal-like (CS), structures. The distortion dihedral angle (θ) between benzene rings in the bridge that minimize the energy in singlet and triplet states is $\sim 36^{\circ}$. Interestingly, the triplet state is more stable than the CS configuration in both, planar and distorted, configurations ($\Delta E_{ST,\theta=0^{\circ}} = -7.78 \text{ kcal/mol}$, and $\Delta E_{ST,\theta=36^{\circ}} = -8 \text{ kcal/mol}$). This result can be associated to the stabilization provided by the aromaticity of the p-biphenyl in the triplet state, since both benzene rings hold a Clar's sextet. Indeed, the significant hydrogen repulsion in the planar conformation can be counterbalanced by the aromaticity, so that the diradical structure (triplet state) is favoured over the closedshell form (singlet). Upon the distortion ($\theta_0 \to \theta_{\sim 36}$), the steric repulsion within the p-biphenyl benzenes lowers and the triplet state is further stabilized by $\sim 2 \text{ kcal/mol}$ with respect to the planar structure. The distorted closed-shell structure is the most unstable situation due to its tendency to break the double bond between benzene rings (similar to the ethylene torsion). In summary, the relative energy of the four conformations follows the trend: $E(CS_{\theta \sim 36^{\circ}}) \gg E(CS_{\theta = 0^{\circ}}) > E(T_{\theta = 0^{\circ}}) > E(T_{\theta \sim 36^{\circ}})$.

These results indicate that the distortion of the central ring of the p-terphenyl is the key structural parameter that governs magnetic properties in the equilibrium between the low and high temperature PAT^{+2} conformers. As mentioned beforehand, conversion between the two isomers is not entropy driven: what is then the driving force behind the interconversion mechanism? To address the interconversion between $PB-PAT^{+2}$ and $DD-PAT^{+2}$, the torsional potential for the dihedral angle θ is computed at the RAS-SF level for the two conformers, as presented in Figure 5.14. These energy profiles are obtained by rigid scans of the torsion, *i.e.*, single point energy calculation with all internal coordinates fixed to their equilibrium values besides θ , which is scanned between -25° and 45° by steps of 5°.

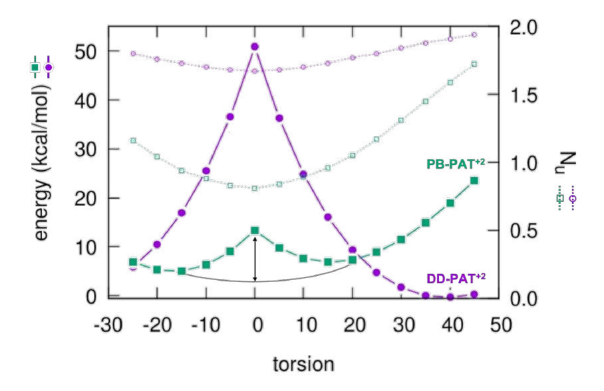


Figure 5.14: Energy profiles and N_u for the **PB-PAT**⁺² (green squares) and **DD-PAT**⁺² (purple circles) molecular torsion model. Vertical arrow represents the steric hindrance in **PB-PAT**⁺².

The most stable conformation is found for **DD-PAT**⁺² at $\theta \sim 35^{\circ}$. From this geometry, the gradual planarization of the p-terphenyl entails the loss of stability due to the increase of steric hindrance (the potential energy curve reaches a maximum when the bridge is perfectly planar). On the other hand, the minimum of **PB-PAT**⁺² is found at $\theta \sim 15^{\circ}$, almost 5 kcal/mol above the **DD-PAT**⁺² minimum, in good agreement with DFT calculations that predict an energy difference between the two minima of 5.4 kcal/mol.

The geometry in which the bridge is perfectly planar might be anticipated to be the minimum of **PB-PAT**⁺². Nevertheless, an energy peak is obtained for $\theta = 0^{\circ}$, which can be interpreted as the consequence of steric effect between the hydrogens of contiguous phenyls in the terphenyl unit (estimated about 5 kcal/mol as indicated in by the vertical arrow in 5.14). The crossing point between **PB-PAT**⁺² and **DD-PAT**⁺² curves locates approximately at $\theta = 23^{\circ}$. The path connecting both minima, from **DD-PAT**⁺² to **PB-PAT**⁺², presents a barrier towards the molecular planarization, *i.e.*, the energy increases due to steric effects up to $\sim 23^{\circ}$, where a pseudo-Jahn-Teller distortion might be correlated with the increment of energy by the gain of local symmetry (at the *p*-terphenyl core). Once passed the crossing point, to stabilize the planar biradicaloid conformation, the quinoidal structure

arises through the formation of a π -bond and the reduction of the diradical character, quantified in Figure 5.14 as the decrease of the N_u from $\theta \sim 23^{\circ} \rightarrow 15^{\circ}$ (**PB-PAT**⁺² minimum). The stability provided by the high symmetrical conformation of the **PAT**⁺² at high temperature is compensated, and exceeded, by the aromaticity recovered in the central ring and internal p-terphenyl steric strain at low temperature conformation.

The entropy should increase with temperature, enabling the population of rotational conformers, as already well described for the oligophenylenes family. (18) The so called "bond stretching isomerism" establishes that breaking of a σ -bond and subsequent stabilization of a diradical is obtained as the effect of a pseudo Jahn-Teller distortion (reduction of the local symmetry to stabilize the energy) when temperature decreases. In other words, bond formation is promoted by heating the system. In **PAT**⁺² conformers, the quinoidal-like configuration of **PB-PAT**⁺² can be understood as the partial formation of a π -bond (N-C_{terphenyl} bond holds a double bond character) as the temperature increases, in such a way that the radical character is decreased, as shown in Figure 5.14 by the diminution of N_u towards $\theta = 0^{\circ}$.

To summarize, the interconversion between the two isomers can be understood as a dynamic process fueled by thermal energy. At low temperature, steric congestion dominates and the equilibrium structure exhibits a p-terphenyl core distortion together with the associated sacrifice of a π -bond. On the other hand, by heating this bond can be formed as the energy gained is invested into keeping steric repulsion. Finally, the origin of the small interconversion barrier between conformers with different radical character was rationalized. Equilibrium between conformers of \mathbf{PAT}^{+2} can be tuned by design of the bridge: use of different linkers, changing the size, modifying its topology or electronic structure nature. Here singlet-triplet energy gap of \mathbf{PAT}^{+2} is thermally controllable in the solid state.

5.3 Appendix

5.3.1 Computational details

Geometry optimizations of \mathbf{OTPD}_n and \mathbf{PAT} have been performed within the density functional theory (DFT) framework with the unrestricted B3LYP and CAM-B3LYP exchange-correlation functionals, respectively. Basis set of double- ζ quality with polarization functions were used, namely 6-31G(d) for \mathbf{OTPD}_n and 6-31G(d,p) for \mathbf{PAT} . Optimized geometries were identified as true local minima with no imaginary frequencies within the harmonic approximation. $\mathbf{UV/VIS/NIR}$ spectra were calculated using TD-DFT method at the same level of theory.

5.3.1.1 $OTPD_n$

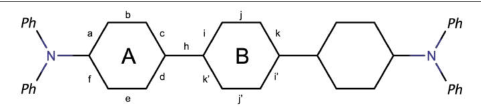
RAS-SF calculations were performed with 4 electrons in 4 orbitals in the active space with triplet high-spin as reference. The bond length alternation is computed as the average of simple bond length minus the average of double bond length trough the π -conjugated path.

5.3.1.2 PAT^{+2}

Raman spectra were simulated within the harmonic approximation by convolution of Lorentzian functions with a half-with at half maximum (HWHM) of 10 cm⁻¹. Computed vibrational frequencies were scaled by a constant factor of 0.96.

RAS-SF calculations were performed including 2 electrons in 2 π -orbitals as the active space (RAS2) with the ROHF triplet high-spin as reference.

The BLA of benzene rings of the *p*-terphenyl are defined as the difference between average of length of longitudinal bonds (a, c, d, f in ring A, and i, k in ring B) and transverse bonds (e and b in ring A, and j in B) in each ring. The bond lengths used to calculate the BLA in Table 5.6 are presented in Table 5.7.



Structure	Average a, c, d and f	Average b and e	Average i and k	Bond j
$^a\mathrm{PB-1}^{+2}$	1.415	1.362	1.396	1.374
$\mathbf{PB-1}^{+2}$	1.433	1.358	1.434	1.358
$^a \mathrm{DD} extbf{-}1^{+2}$	1.407	1.376	1.402	1.379
$\mathbf{DD-1}^{+2}$	1.402	1.382	1.399	1.387
1	1.405	1.390	1.406	1.391

Table 5.7: Average of transverse a, b, c and d (h and i) and longitudinal b and e (j) bonds of the A (B) ring for high and low temperature conformers at DFT and ${}^{a}X$ -ray(?) structure.

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Chapter 6

Organic macrostructures with polyradical character

As was proclaimed in previous Chapters, 4 and 5, small to medium size organic radicaloids have been extensively applied in the development of emerging technologies in miscellaneous fields, from light/energy generation, transport and storage to medical and biological sciences.(1; 2; 3; 4) Performance on this areas can be improved by heightening of radical character. In other words, materials with more than two radical centers (*i.e.*, polyradicals) could be valuable for applications based on magnetic, electronic, and optic properties.() The interaction between radical sites is related with the potential application of polyradicals, for instance ferromagnetic interaction is important for organic magnets,(?) while the anti-ferromagnetic one can help to improve electronic conjugation between radicals.(?)

Despite the constant efforts and development of synthetic routes, nowadays is not straightforward to achieve the production of stable polyradical materials. Nevertheless, small organic radicals can be experimentally obtained and applied thanks to special stabilization techniques, as described in Chapter 2. A well-known example is the polychlorotriphenylmethyl (**PTM**) radical, derivative of the **TM** monoradical, where phenyl rings are totally (or partially) chlorinated (Figure 6.1 a). Steric strain provided by the Cl atoms at *ortho* position force the phenyl rings to adopt a propeller-like conformation, which sterically protects the methyl radical site providing an improved stability and long life-times.(5; 6)

Another paramount example of carbon-radicals stabilized by protection with bulky groups is the cyclopentadienyl ring (\mathbf{Cp}), Figure 6.1. Stabilization of the unpaired electron can be improved by the expansion of the π -conjugation around the radical center. For instance, the benzannulation, as in fluorenyl radical (\mathbf{FR}), is a common way to effectively hyper-conjugate and delocalize the spin through the system. \mathbf{FR} is an extensively exploited frame in the design of radical macrostructures (7; 8), since it can be easily stabilized by kinetic blocking the 9-position with a bulky group.

Figure 6.1: Structure of tripehenylmethyl, **TM**, radical and its polychlorated derivative, **PTM**, (a). Structure of the cyclopentadienyl ring, **Cp**, and the extended aromatic system fluorenyl, **FR** (b).

Given their stability, these moieties are invaluable frames to build larger systems with high-spin ground state or polyradical character. These radical centers are connected by means of a rigid or flexible organic linkers to produce macroestructures with a large diversity of topological arrangements (linear, macrocyclic, polymeric, and the so-called spin clusters, that is branched polyradicals). (? ? ? ?) For instance, recently, 3D motifs have been constructed as the ensemble of multiple radical units.(? ?)

In this chapter, I will describe stable macrostructures with polyradicaloid character constructed as the assembly of **FR** and **PTM** radical moieties. The relation between connectivity and the nature of the radical interactions (ferromagnetic or antiferromagnetic) is studied here, rationalizing their strong open-shell nature. Finally, those properties induced from the radical character will be investigated from a computational point of view by means of correlated electronic structure methods, as demanded by the nature of the systems. In particular, I focus my attention to a series of systems synthesized and experimentally characterized by the group of Prof. Jishan Wu from the National University of Singapore.

6.1 Organic Macrocycles with Radical Character

Here I present the study of three families of macrocycles (Figure 6.2). [n]cyclopara-biphenylmethines ([n]CPBM) with n=3-8 are formed by n p-diphenyl
units connected through a methine carbon, giving rise to n radical sites in each vertex which are protected by an anthracene derivative. As described in the previous
chapter, p-phenyl bridges bear flexibility to the cycle through the relative tension
between benzene rings, modulating the overall radical character through the equi-

librium between aromatic and quinoidal-like forms. Similar structures are shown by [n]PTM macrocycles, where n = 3 - 6 units of PTM radical are linked together by an ethynylene moiety that provides molecular rigidity avoiding steric hindrance between polychlorated phenyls, and increases the distance between radical centers compared with [n]CPBM. Finally, [8]MC and [10]MC are coronnene-like cycles of fused alternated 5- and 6-member carbon rings, resulting in an arrangement of n fluorenyl radical-like units. Structurally, all these macrocycles are able to hold a large number of radicals, i.e, to be polyradicals. Concretely, they are potential triradicals up to decaradicals according with n. Experimentally, every potential radical site is protected with a bulky group (see Appendix), for computational reasons these groups were reduced (in [n]CPBM) or replaced by hydrogen atoms (in [n]MC).

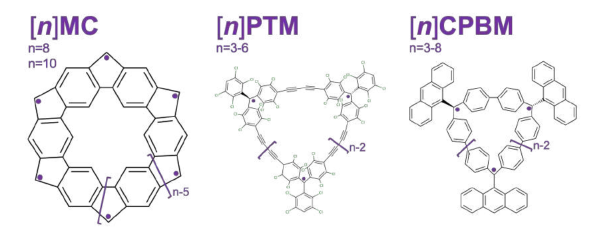


Figure 6.2: Structure of macrocyclic families studied here [n]MC with n = 8 and 10, [n]PTM with n = 3 - 6, and [n]CPBM with n = 3 - 8. Purple dots indicate the radical centers.

Geometrical minima were determined by means of DFT theory. [n]MC are very rigid structures with well defined molecular symmetry. The larger, [10]MC, is a flat macrocycle with D_{2h} symmetry (in good agreement with the X-ray structure) and spin-triplet ground state, while [8]MC holds a singlet ground state, and the ring tension suppress molecular planarity favouring a bowl-shape structure with C_{2v} molecular symmetry (Figure 6.3 a). [n]PTM with n=3 and 5 have a doublet ground state, while n=4 ground state is a singlet and with n=6 the ground state multiplicity is the triplet. The distance between radical centers in [n]PTM slightly increases with n, (~ 14.40 , 15.09, 15.26, and 15.30 Å, for n=3-6, respectively). Ethynylene bridges are aligned in a (pseudo)planar macrocycle in n=3-5, but in the largest member of the family, [6]PTM, the tub-shape conformation of the PTM radical leads to a boat-shaped conformation, similar to the one observed in cyclohexane (Figure 6.3 b). [n]CPBM hold low-spin multiplicity ground states, i.e., if n is odd they are doublets, and singlets when n is even. The relative orien-

tation of the phenyl rings obtained as is an interplay between repulsion inside the bridge but also with the contiguous bridge. As a result, [3]CPBM has a highly tense bowl-shaped structure with nonequivalent distances between radical centers (9.20, 9.06 and 8.63 Å). The tetramer exhibits an almost flat rectangular conformation, which nicely reproduces the crystalline distorted- C_2 geometry with distances between neighboring methyl sites of 9.56 and 9.91 Å computed as 9.73 and 9.93 Å, respectively. As the size of the system increases, the structure of the macrocycle is distorted to a tub-like conformation for n = 6 and 8, and the twisted 8-figure for n = 7 (Figure 6.3 c). Increase of the radical-radical distance suggests the weakness of the coupling between radicals in [n]PTM and [n]CPBM macrocycles with the increasing of n.

The polyradicaloid character of these macrocycles is summarized in Table 6.1. Descriptors, namely, $\Delta E_{gs,es}$ (energy gap from ground state to first excited state of higher multiplicity), y_n and N_u , were computed with RAS-SF methodology, the active space and high-spin multiplicity are system dependent and explicitly shown in the Appendix. Note that radical index y_i is obtained as the LUNO+i occupation, when n is odd y_0 (LUNO), y_1 (LUNO+1), y_2 (LUNO+2), y_3 (LUNO+3) and y_4 (LUNO+4) are the tri, penta, hepta, nona and undecaradicaloid characters, respectively. While, for n-even y_0 , y_2 , y_3 , y_4 and y_5 are the di, tetra, hexa, octa and decaradicaloid index.

[n]	State	Energy	N_u	y_0	y_1	y_2	y_3	y_4
[n]CPBM								
3	Doublet	0.000	1.70	0.35	-	-	-	-
	Quartet	4.186	3.00		-	-	-	-
4	Singlet	0.000	2.95	0.75	0.73	-	-	-
	Triplet	3.396	3.47			-	-	-
	Quintet	6.956	4.00			-	-	-
5	Doublet	0.000	3.94	0.74	0.73	-	-	-
	Quartet	3.395	4.51			-	-	-
	Sextet	6.971	5.00			-	-	-
6	Singlet	0.000	4.61	0.82	0.75	0.73	-	-
	Triplet	1.632	4.98				-	-
	Septet	8.200	6.00				-	-
7	Doublet	0.000	5.42	0.75	0.74	0.72	-	-
	Quartet	3.252	5.93				-	
	Octet	10.390	7.00				-	-
8	Singlet	0.000	5.93	0.76	0.75	0.73	0.73	-
					Conti	nued o	n next	page

Table 6.1 – continued from previous page

[n] State Energy N_u y_0 y_1 y_2 y_3 y_4 Triplet 3.074 6.44 Nonet 13.248 8.00 [n]PTM 3 Doublet 0.0000 3.00 1.00 Quartet 0.0000 3.00 $-$ 4 Singlet 0.0000 3.99 1.00 1.00 Triplet 0.0003 3.99 $-$ Quintet 0.0007 4.00 5 Doublet 0.0000 4.98 1.00 1.00 Quartet 0.0008 4.99
Nonet 13.248 8.00
[n]PTM 3
3 Doublet 0.0000 3.00 1.00
Quartet 0.0000 3.00 -
4 Singlet 0.0000 3.99 1.00 1.00 Triplet 0.0003 3.99 Quintet 0.0007 4.00
Triplet 0.0003 3.99 Quintet 0.0007 4.00
Quintet 0.0007 4.00 5 Doublet 0.0000 4.98 1.00 1.00
5 Doublet 0.0000 4.98 1.00 1.00
Quartet 0.0008 4.99
~
Sextet 0.0016 5.00
6 Singlet 0.0001 5.95 1.00 0.99 0.99 -
Triplet $0.0000 5.95$
Septet 0.0261 6.00
[n]MC
8 Singlet 0.0 4.85 0.73 0.58 0.58 0.50 -
Triplet 2.7
Nonet 20.6 8.00 -
10 Singlet 0.4 7.08
Triplet 0.0 7.04 0.97 0.94 0.57 0.52 0.51
11-et 22.8 10.00

Relative energies of ground state and excited states with lowest and maximum multiplicity (according with n) of [n]CPBM, [n]-PTM and [n]MC. Energy is in kcal/mol, N_u represents the number of unpaired electrons, and y_i is the ith radical index.

In the ground state these macrocycles are characterized by large N_u and y_i values, as well as the small energy gaps between ground state and lowest-lying excited state of different multiplicity. [8]MC is a singlet octaradicaloid ($y_3 = 0.50$) with 4.85 unpaired electrons. In the triplet ground state [10]MC have large y_i values up to $y_4 = 0.51$ ($y_1 = 0.97$, $y_2 = 0.94$, and $y_3 = 0.57$), meaning it possesses moderate decaradicaloid character. In fact, [10]MC was the first stable (finite) organic decaradicaloid system reported at the moment of its synthesis and characterization (2018).

In [n]**CPBM** the polyradical character is moderate, with 1.70, 2.95, 3.94, 4.61, 5.42 and 5.93 unpaired electrons from n = 3 - 8, respectively, and states of the

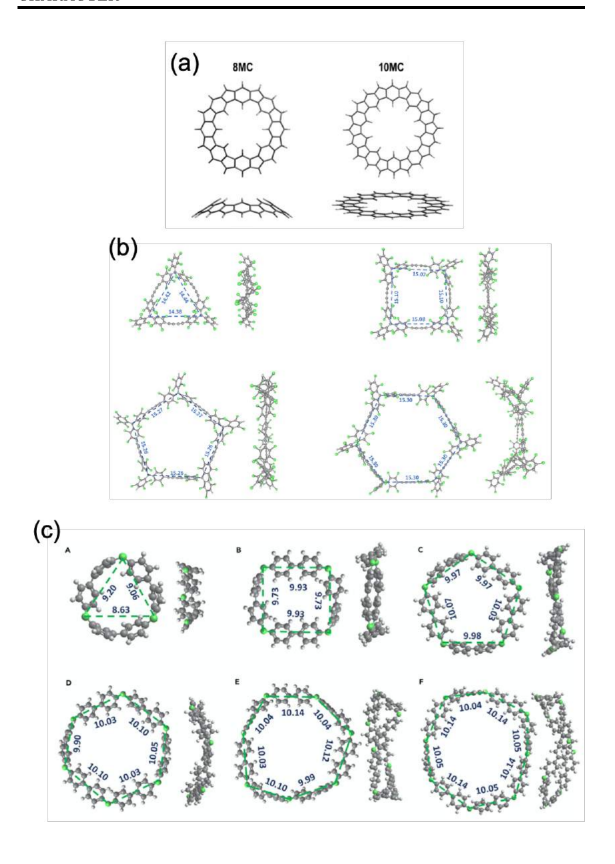


Figure 6.3: Optimized structures and relevant geometrical parameters of [n]MC (a), [n]PTM (b), and [n]CPBM (c).

higher multiplicities computed at \approx 10 kcal/mol with respect to the ground state.

Experimental determinations by SQUID give energy gaps between ground state and states with higher multiplicity of < 4 kcal/mol.

There are $\sim n$ unpaired electrons in $[n]\mathbf{PTM}$, and a quasi-degeneration between states with different multiplicities resulting in an ambiguous ground state spin. To further explore and analyze this situation, specific spin configurations at the ground state geometry were computed by imposing one electron to be localized over each one of the n \mathbf{PTM} radical centers as presented in Table 6.2. Notice that in [3] \mathbf{PTM} the energy of configuration β , α , β is not equal to α , β , α because the sides of the triangle are not equivalent. The imposed spin restrictions split the energy of the states of different multiplicities, but with rather small energy gaps ($\Delta E < 1$ kcal/mol).

[n]PTM	ΔE	Spin configuration
[3]PTM		
	0.000	lpha,eta,lpha
	0.066	eta, lpha, eta
	0.066	α, α, α
[4]PTM		
	0.000	lpha,eta,lpha,eta
	0.077	$\alpha, \alpha, \beta, \beta$
	0.154	$\alpha, \alpha, \alpha, \alpha$
[5]PTM		
	0.000	eta, lpha, eta, lpha, eta
	0.182	$\alpha, \alpha, \alpha, \alpha, \alpha$
	0.581	$\alpha, \alpha, \alpha, \alpha, \beta$
[6]PTM		
	0.000	$\alpha, \beta, \alpha, \beta, \alpha\beta$
	0.172	$\alpha, \beta, \alpha, \alpha, \beta, \alpha$
	0.320	$\alpha, \alpha, \beta, \beta, \alpha, \beta$
	0.940	$\alpha,\alpha,\alpha,\alpha,\alpha,\alpha$

Table 6.2: Energy gap between different spin configurations (kcal/mol) computed with the C-DFT methodology.

Considering the cyclic arrangement of [n]PTM and [n]CPBM the interradical coupling can be approximated by $\Delta E_{S,T} = 2J$ ($\Delta_{D,Q} = 3J$) for even (odd) radical sites.(9) While $J \approx 1.5$ and $J \sim 0$ kcal/mol in [n]CPBM and [n]PTM, respectively, indicating the weak to very weak AFM coupling of these macrocycles. Delocalization of the unpaired electrons in [n]MC prevents the characterization of low-lying states of different spin multiplicities to an effective spin Hamiltonian, and I did not attempt

to define a coupling interaction constant in these systems.

In summary, macrocycles of these families can be regarded as *n*-radicaloid (from tri up to decaradicaloid) with weak to medium AFM coupling between radical centers that favours a low-spin ground state multiplicity. It is remarkable that all of them are stable under ambient conditions.

6.1.1 π -conjugation: [n]annulene analogs

The large number of radical sites in addition to the π -conjugated backbone favors the delocalization of the electronic density over the whole cycle. This is reflected as the formation of alternated aromatic-quinoidal-like structure of the p-phenyl units of [n]CPBM; single-double bond alternation in [n]MC is extended over the both edges of the macrocycles. Differently, the presence of the ethynylene group in the bridge of [n]PTM gives rise to the equilibrium between single-triple bond alternation and the cumulene conjugation. On the other hand, the benzene rings present in these macrocycles tend to form Clar's sextets and recover their local aromaticity, leading to the presence of unpaired electrons and the rise of polyradical character. Equilibrium between quinoidal and local aromatic structures of [8]MC, [4]CPBM and [4]PTM are illustrated in Figure 6.4.

Due to their geometry and conjugation [n]CPBM and [n]PTM could be thought as large-size annulenes with n sides, and the aromatic basic rules apply to them. Namely, if there are $(4n+2)\pi$ electrons according to the Hückel's rule, annulenes are classified as aromatic and tend to delocalize their π electrons over the whole cycle. But, if there are $4n\pi$ electrons in the system, annulenes are considered antiaromatic and they avoid electron delocalization. The interconversion between both valence isomers in annulenes can occur via polyradical transition state (??????), and can be experimentally detected if the rate constant is slow. This process is called "valence tautomerization" and involves the break and posterior formation of double bonds. As a consequence, the valence tautomerization goes through an intermediate with unpaired electrons. Despite of the big efforts to characterize it, the process reminds experimentally unobserved, presumable due to the presence of small reaction barriers.

Particularly, [4]CPBM, where two opposite sides of the macrocycle hold quinoidal-like structure and the other two are aromatic (as radical-radical distance in Figure 6.4), can be compared with the cyclobutadiene (CBD) structure. In the ground state CBD have two clearly localized double bonds connected by two single bonds. Then, cyclobutadiene is the smallest antiaromatic annulene (with 4π electrons). The valence tautomerization of CBD is an ultra fast process with a transition state theoretically predicted as a square diradical.(? ? ? ? ? ? ? ? ? ?) [4]CPBM has 36π electrons, and it can be thought as an antiaromatic annulene. Its size and the large number of bonds that must be broken and formed in the valence tautomerization, in

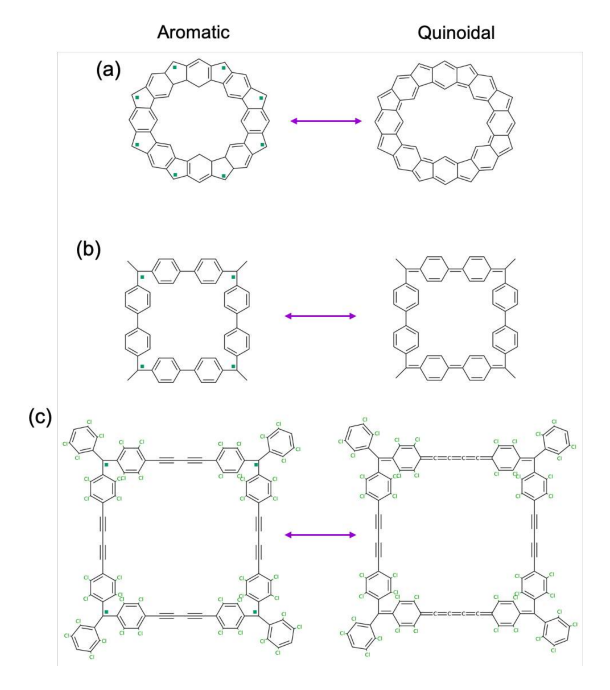


Figure 6.4: Equilibrium between quinoidal and local aromatic structures of [8]MC (a), [4]CPBM (b) and [4]PTM (c).

addition to the loss of local aromaticity in 4 benzene rings, increase significantly the barrier of the process and make it experimentally detectable by variable-temperature (VT)-NMR measurements.

Experimentally, the process was followed by the evolution of the chemical shift of the NMR of the non-equivalent hydrogen atoms of benzene rings in the *p*-diphenyl bridges in the [4]CPBM. Spectra at 310 K (Figure 6.5) shows two signals, suggesting that there are just two kind of hydrogen atoms in the macrocycle backbone. The

first one corresponds to those close to the bulky protector group (7, 7', 9) and 9' in Figure 6.5) and the other one to hydrogen in the centre of each side (8, 8', 10) and (8, 8', 10) are indication of the very fast interconversion between quinoidal and aromatic structures. By cooling the sample the signals are broadened and they coalesce at 263 K. Then, around of 243 K the signals of (8, 8', 10) and (8, 8', 10

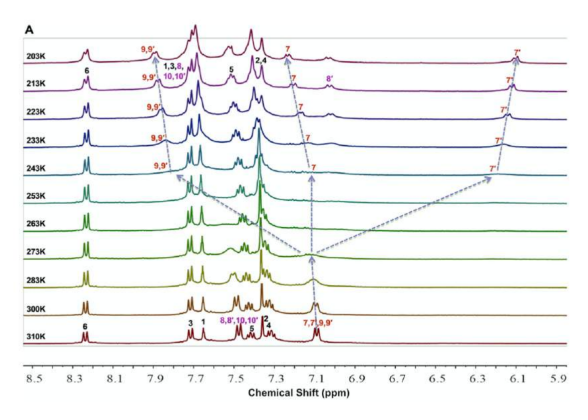


Figure 6.5: VT-NMR spectra of the aromatic region of [4]CPBM. The arrows show the splitting of the protons 7, 7, 9, and 9 at low temperature.

Theoretically, the transition state between valence isomers of [4]CPBM was computed as a global antiaromatic C_4 structure, found at 7.1 kcal/mol higher than the C_2 ground state (Figure 6.6). In the transition state geometry, all the four sides of [4CPBM] are equivalent, meaning that the four benzene rings localized in two opposite sides of the macrocycle in C_2 geometry, lost their aromatic sextet and

contributes with 4 π -electrons each ones to the whole ring. As a consequence, the transition state is globally antiaromatic with $4n\pi$ -electrons.

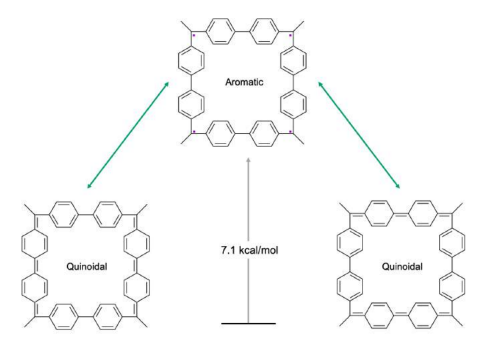


Figure 6.6: Valence tautomerization of [4]CPBM. Two quinoidal structures are connected through an antiaromatic transition state by a energy barrier of 7.1 kcal/mol.

Experimentally, the tautomerization of [n]CPBM with n=6 and 8 is not observed, indicating that the interconversion barrier must be too small enough to be detected. Hence, [4]CPBM represents a rare example of annulene-like system carrying out a detectable valence tautomerization process.

6.1.2 Global aromaticity of [8]MC and [10]MC

Coronoid-like molecules (as [n]MC) are cyclc compositions of fused benzene rings that can be drawn as an alternation of aromatic rings with Clar's sextet or as the union of two peripheral singlet-double bond chains, [n]annulenes, one inside of the other covalently linked but only by single bonds, in other words, whit no conjugation between both annulenes. These structures have been named as annulenne-whitin-an-annulene (AWA) super ring and were proposed for the first time in 1966 with the synthesis of corannulene(?) (Figure 6.7 a). But the AWA idea was discarded by means of theoretical calculations, indicating that inner-outer annulenes are not decoupled, there is at least one double bond linking them.(?) Lately, kekulene(?) (Figure 6.7 b) was proposed theoretically with an AWA structure consisting of an [18]annulene within an [30]annulene, its synthesis in 1978 proved that its structure is dominated by the presence of Clar's rings. Since then, constant efforts have been done, in the design of macrocycles with the AWA structure. For instance, the

alternation of five and six member rings partially prevents the aromatic benzene formation promoting the annulene formation in segments of the macrocycle.(?) However, a truly AWA super ring structure has never been found.

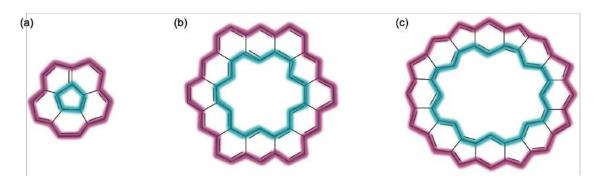


Figure 6.7: Examples of first predicted AWA structures: coronnene (a) kekulene (b). AWA structure of [8]MC (c).

The close inspection of the quinoidal structure (Figure 6.4 c) of [8]MC shows the absence of Clar's sextets and the formation of two, "in" and "out", disconnected annulenes. Actually, the bond lengths of [n]MC correspond to the single-double alternation in the outer and inner rings, and to single bonds connecting conjugated systems in the bridge between both rings (~ 1.46Å). To further characterize the conjugated systems in the singlet (triplet) ground state of [8]MC ([10]MC) I have performed induced current density (ACID) calculations at (U)B3LYP/6-31G(d,p) level of theory. In both macrocycles, ACID plots show two fully decoupled ring current circuits (Figure 6.8 a) localized over the annulenes in the periphery of the macrocycle. These results suggest the formation of two truly AWA super-ring structures. [8]MC and [10]MC are characterized as [24]annulene-within-an-[32]annulene and [30]annulene-within-an-[40]annulene super-ring structures, respectively.

[8]MC, as the combination of [24] and [32] annulenes, has 56 π -electrons and should be classified as an antiaromatic system (4n π with n = 14). Surprisingly, a current flow in the ACID plots show opposite aromaticity than predicted. The diamagnetic current flow in both, inner and outer annulenes in [8]MC is clockwise, corresponding to an aromatic system but since it contains $4n\pi$ electrons, it was expected to exhibit counter-clockwise current (Figure 6.8 b). In summary, it is a [24]annulene-within-an[32]annulene with unusual local and global aromaticity.

On the other hand, the internal ring in [10]MC has $(4n+2)\pi$ electrons, while the external one is a $4n\pi$ electron ring. Then, [10]MC has $(4n+2)\pi$ electrons. Nevertheless, it is worth to remember that the ground state of [10]MC is a triplet, then Baird's rule should be applied to predict the aromaticity of the system. Under Baird's rule the lowest-lying triplet exhibit aromatic (antiaromatic) character if there are $4n\pi((4n+2)\pi)$ electrons, according with it, this macrocycle should be expected

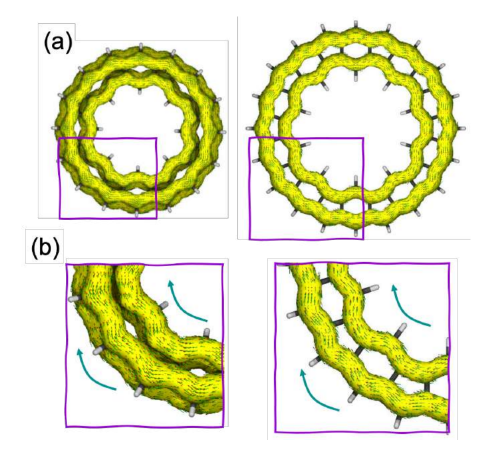


Figure 6.8: ACID current of [8]MC (left) and [10]MC (right) in their ground state conformation, singlet and triplet, respectively (a). Green arrows are clockwise direction an represents the diamagnetic current in each cycle (b).

to be antiaromatic.

Experimentally, the ¹NMR spectra of [8]MC shows two signals that can be associated to hydrogen atoms of the outer and inner perimeter, named as a and b, see Figure 6.9 a and b. Spectra was recorded at few temperatures between 298 and 233 K, showing a group of bands associated to the protons in the bulky protector group, and two peaks, that presumably correspond to hydrogen in inner and outer rings. At very low field and the other one at very high field, last one is small and is better defined at low temperature. Differently from the NMR spectrum of [4]CPBM, the broadening observed by heating is characteristic of open-shell singlet diradicaloids and not by the change of the chemical nature of hydrogen atoms. Our calculations nicely reproduce the ¹H NMR spectrum, showing one signal for the eight b protons at very high field ($\delta = -24.06 - -24.10$ ppm) that correspond to the protons in the inner ring. The other signal, for all the eight hydrogens in the outer ring, was found at very low field ($\delta = 14.80 - 14.78$). Therefore, [8]MC can be regarded as a highly aromatic system. On the other hand, [10]MC is ¹H NMR silent due to its triplet ground state.

In order to rationalize the unusual global aromaticity of both macrocycles, based

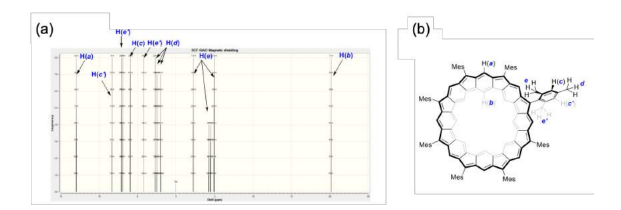


Figure 6.9: Calculated (restricted) 1H NMR spectrum (B3LYP/6-31G(d,p)-GIAO) based on the optimized geometry (a) and hydrogen labels of [8]MC (b). The bowl pointing from the inner hub to the outer rim through the paper. Protons e and c point to the concave side, while protons e' and c' point to the convex side.

on their AWA properties we can consider the whole macrocycle as the sum of two parts (inner and outer rings). Hence the "adiabatic" state (the one describing [n]MC) can be understood as the combination of "diabatic" states with well defined charge and spin on each annulene. Within this approximation, the aromaticity of [8]MC can be explained by two alternative electronic structures: (1) one annulene gives two electrons to the other one, then both are $(4n+2)\pi$ electrons systems and therefore aromatic singlets (Hückel's rule); or each annulene holds a triplet (diradical) ground state and are aromatic according with the Baird's rule. To that end, constrained DFT (C-DFT) calculations were performed with restrictions imposed to inner and outer annulenes, resulting configurations are S/S, T/T, 1+/1-, 1-/1+, 2+/2- and 2-/2+, all them are written as "in/out" annulenes. The main idea is to provide energetic arguments to discern which conformations can stabilize the total wave function.

The relative energy of each pair of diabatic states (Figure 6.10) indicates that 2+/2- and 2-/2+ states are much higher in energy than S/S, T/T, 1+/1- and 1-/1+. Then, the participation of dianionic states in the total wave function is expected to be insignificant. This excludes the option of having two $(4n+2)\pi$ annulenes with singlet ground state exhibiting Hückel aromaticity. Hence, second option have to be considered to explain the global aromaticity of [8]MC, that is, both annulenes are open-shell (computed as triplet) ground state, *i.e.*, diradicals, with aromatic properties according to the Baird's rule. The presence of two open-shell annulenes is in very good agreement with the high polyradical character of this macrocycle and the number of unpaired electrons ($N_u = 4.85$, Table 6.1). Furthermore, this explains the flow currents observed in ACID plots.

The ground state triplet of [10]MC can be described as the combination of

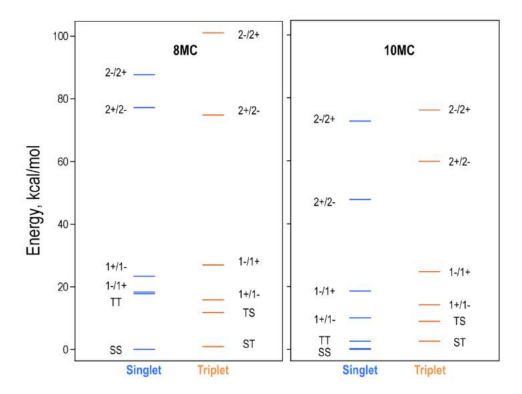


Figure 6.10: State energy diagram for the diabatic states of [8]MC and [10]MC molecules computed with C-DFT. All energies are given with respect to the lowest singlet or triplet state. State labels correspond to the restriction on inner/outer annulene rings, respectively. S/S = singlet/singlet, T/T = triplet/triplet, 1-/1+ = anion/cation, 1+/1- = cation/anion; 2-/2+ = dianion/dication, 2+/2- = dication/dianion.

singlet and triplet annulenes with 30 and 40 π electrons. The lowest energy contribution is the S/T state, and ACID plots reveal their clockwise and counter clockwise current flow, like this both aromaticity rules are satisfied simultaneously, inner ring is an aromatic singlet (Hückel) and the outer ring is an aromatic triplet (Baird's rule). These results confirms its super-aromatic AWA structure.

The ACID plots of the lowest-lying excited states of [n]MC shows antiaromatic (n = 8) and non-aromatic (n = 10) features. Again, their wave function can be rationalized by the contributions of the diabatic states of each ring to the adiabatic state of the macrocycle. The first triplet excited state of [8]MC is mainly the S/T combination of sates and it must be understood as an antiaromatic [24]annulene with singlet state (Hückel rule) within an aromatic diradicaloid with $4n\pi$ electrons (Baird's rule). This combination of states is in good agreement with the counter-clockwise paramagnetic (inner) and clockwise diamagnetic (outer) current flow observed in the ACID plots (Figure X?). The lowest-lying excited state of [10]MC is a singlet with no current flow, therefore is a non-aromatic system, this

might be seen as the result of the aromatic-antiaromatic cancellation. S/S and T/T are close in energy and to differentiate between them is not straightforward, nevertheless, both configurations combines one $(4n+2)\pi$ and one $4n\pi$ annulenes, *i.e.*, one aromatic and one antiaromatic rings under both, Hückel and Baird, rules.

Finally, the ionic configurations (1-/1+ and 1+/1-) are low in energy, and can be considered as a minor contribution to the polyradical character of these macrocycles, but differently from the singlet and triplet states, there is not a well-established rule to determine the aromaticity on doublet states. Mandado et al. suggested an expansion to the Baird's rule to be applied in order to evaluate the aromaticity as the sum of separate contributions from α and β π -electrons (see the details in the Appendix).(?) To this end, we deconstructed the macrocycles as the inner and outer annulenes ([X]MC_in or _out), Table 6.3 summarizes the predictions from the "extended Baird's rule".

Ring model	Spin multiplicity	Charge	α -elec	β -elec	Global			
[8]MC_in	\mathbf{S}	0	12 (AA)	12 (AA)	AA			
[8]MC_in	${ m T}$	0	13 (A)	11 (A)	A			
[8]MC_in	D	-1	13 (A)	12 (AA)	NA			
[8]MC_in	D	+1	12 (AA)	11 (A)	NA			
[8]MC_in	S	-2	13 (A)	13 (A)	A			
[8]MC_in	T	+2	11 (A)	11 (A)	A			
[8]MC_out	\mathbf{S}	0	16 (AA)	16 (AA)	AA			
[8] MC _out	T	0	17 (A)	15 (A)	A			
[8] MC _out	D	-1	17 (A)	16 (AA)	NA			
[8] MC _out	D	+1	16 (AA)	15 (A)	NA			
[8] MC _out	\mathbf{S}	-2	17 (A)	17 (A)	A			
[8]MC_out	Τ	+2	15 (A)	15 (A)	A			
[10]MC_in	S	0	15 (A)	15 (A)	A			
$[10] \mathrm{MC}_{\mathrm{in}}$	${ m T}$	0	16 (AA)	14 (AA)	AA			
[10]MC_in	D	-1	16 (AA)	15 (A)	NA			
$[10] \mathrm{MC} $ _in	D	+1	15 (A)	14 (AA)	NA			
$[10] \mathrm{MC}$ _in	S	-2	16 (AA)	16 (AA)	AA			
$[10] \mathrm{MC_in}$	Т	+2	14 (AA)	14 (AA)	AA			
[10]MC_out	S	0	20 (AA)	20 (AA)	AA			
$[10] \mathrm{MC}_{\mathrm{out}}$	T	0	21 (A)	19 (A)	A			
$[10] \mathrm{MC}_{\mathrm{out}}$	D	-1	21 (A)	20 (AA)	NA			
[10]MC _out	D	+1	20 (NA)	19 (A)	NA			
	Continued on next page							

			1	1 6	
Ring model	Spin multiplicity	Charge	α -elec	β -elec	Global
[10]MC_out	S	-2	21 (A)	21 (A)	A
[10]MC_out	${ m T}$	+2	19 (A)	19 (A)	A

Table 6.3 – continued from previous page

Table 6.3: Contributions to the global aromaticity from α and β π -electrons for each diabatic state of inner (_in) and outer (_out) rings of **[8]MC** and **[10]MC** molecules. S = singlet (closed-shell), D = doublet, T = triplet, A = aromatic, AA = antiaromatic, NA = no aromatic (cancellation between α and β

Anionic annulenes have no aromatic character, then they do not contribute to the global aromaticity of the macrocycles. These results are also in good agreement with the contributions given by the energetic ordering of C-DFT configurations, showing that the singlet of [8]MC can be aromatic only from the combination of triple(inner)-triplet(outer) rings, while the sharing of 2 electrons also leads to global aromaticity in a singlet, energetically their contribution can be neglected. Similarly, the singlet(inner)-triplet(outer) is the only one that gives an aromatic triplet.

It is necessary to stress the qualitative character of these results. They should be used only to distinguish between two extreme situations. The entire complexity of these macrocycles is dictated by their multiconfigurational character and cannot be recovered by C-DFT calculation.

6.2 Fluorenyl dendrons

Etymologically, the term dendrimer comes from the Greek words dendron (tree) and mero (part), meaning "the part of a three". Actually, their structure resembles to snowflakes or treetops.(10) Structurally, they are polymers growing up in branched arrangements surrounding a core. The number of units in the polymer determines the generation of dendrimers and dendrons. Thus, when the core has one layer is the first generation, if it has two layers is a dendrimer of the second generation, and so on up to generation n (Figure 6.11).(? 10; 11) The physical properties triggered by the structure, i.e., viscosity, flexibility, solubility, and density distribution can be modulated by the nature of the end group, leading to applications in a huge number of fields, from medicine as drug carrier agents, tumor sensors or contrast agents for bioimaging, etc.(11; 12) to photoelectronic material science as light-harvesting systems or emitting polymers LEDs.(11? ? ? ? ; 10?)

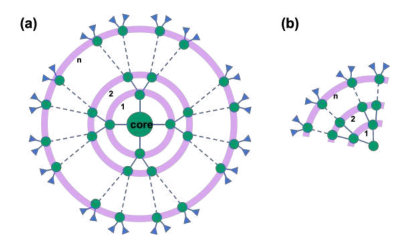


Figure 6.11: General structure of a dendrimer (a) and dendron (b). Central green circle is the core, and the small ones represent branching units. Blue triangles are the end group. Each generation is indicated by purple circles and labeled as first (1), second (2), or n-th (n).

The group of Prof. Rajca has worked through decades extensively in the design and synthesis of denditric radicals (also referred as spin clusters), which exhibit a massive number of radical sites with very strong ferromagnetic interaction between them, exhibiting impressive ground state spin multiplicities (up to 500), but with low stabilities.(13) No evidence of stable radical or polyradical dendrons and dendrimers was available in the literature before 2018 (when the investigation here presented was published).(?)

The expansion of the polyradical macroestructures in branches (following the dendritic arrangement) using the fluorenyl radical (FR) as building block is explored here. The first generation of fluorenyl radical monodendron, FR-G1, is built by connecting two fluorenyl moieties through 9-position to the central FR in 3,6-positions. Expanding FR-G1 with one more layer using the same pattern, the dendron radical of second generation FR-G2 is obtained. The 9-position of central fluorenyl moiety is kinetically blocked by a bulky 9-(3,5-ditert-butylphenyl)anthryl and 3,6-positions of the outer units are blocked by 4-tert-butylphenyl groups, as a result they are stable. Experimentally, half-life of FR-G1 and FR-G2 has been determined upon exposure to air and light ambient conditions monitored by UV/VIS/NIR spectroscopy. FR-G1 has a half-life of 6 days, while the lifetime of the larger FR-G2 is slightly shorter, lasting a bit more than 4 days.(14)

The connectivity of **FR** units in **FR-G1** and **FR-G2** triggers the formation of quinoidal and aromatic forms (Figure 6.12). In one extreme, all six-member carbon rings preserve their aromatic sextets. On the other side, the formation of *para*-quinodimethane-like bonds between fluorenyl units in one or two of the branches of **FR-G1** and **FR-G2**, respectively, leads to the quinoidal-like structure of these fluorenyl dendrimers. Additionally, an intermediate situation can be drawn

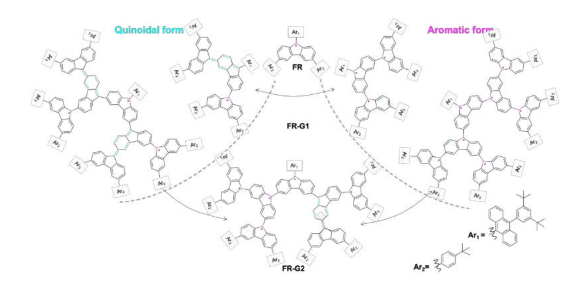


Figure 6.12: Structure of fluorenyl radical (**FR**) surrounded by the first generation fluorenyl radical mono-dendron (**FR-G1**) and the second generation fluorenyl radical mono-dendron (**FR-G2**). Quinoidal bonds are indicated in blue and radical sites in pink.

for FR-G2. Aromatic and quinoidal forms stabilize the open- and closed-shell configurations, respectively. In the aromatic structure, the number of "free" radicals and the multiplicity is expected to be maximized, for instance, FR-G1 has 3 radical centers, and FR-G2 can stabilize up to 7 unpaired electrons. The ground state of FR-G1 is predicted to be a doublet (D_0) , while FR-G2 have a quartet (Q_0) ground state. In these minima, the bond length of the linker between fluorenyl units does not correspond to a double bond, FR-G1 = 1.46, 1.467 Å and FR-G2 = 1.463, 1.467, 1.468, 1.465, 1.467, 1.467 Å. This suggest that the aromaticity of the benzene rings is preserved in both dendrons (Figure 6.12 right) and they are open-shell systems.

The radical characterization of minimum energy structures is summarized in Table 6.4. The doublet ground state of FR-G1 with close to 2 effective unpaired electrons is computed to lie 8.19 kcal/mol below the first excited quartet. The quartet ground state of FR-G2 has more than 5 effective unpaired electrons. The lowest-lying doublet state is computed just 0.78 kcal/mol higher in energy than Q_0 . Computational description of both systems is in good agreement with the doublet-quartet energy gaps of FR-G1 ($\Delta E_{D-Q} = -3.9$ kcal mol⁻¹) and FR-G2 ($\Delta E_{D-Q} = 0.2$ kcal mol⁻¹) obtained through electron spin resonance (ESR) experiments.

The interaction between radicals in these systems can be modeled by means of the effective spin Hamiltonian

$$H = -2J \sum_{i < j} S_i S_j, \tag{6.1}$$

FR-Gn	Multiplicity	Energy	N_u	n _{SONO-3}	n_{SONO-2}	n _{SONO-1}	n _{SONO}	n _{SONO+1}	n _{SONO+2}	n _{SONO+3}
FR-G1	Doublet	0.00	1.82	-	-	1.63	1.00	0.37	-	-
	Quartet	8.19	3.08	-	-	1.00	1.00	1.00	-	-
FR-G2	Quartet	0.00	5.23	1.50	1.42	1.00	1.00	1.00	0.58	0.50
	Doublet	0.78	4.71	1.64	1.42	1.12	1.00	0.88	0.58	0.36

Table 6.4: Characterization of lowest doublet and quartet states of **FR-G1** and **FR-G2**. Energy is in kcal/mol, N_u represents the number of unpaired electrons, $n_{SONO\pm i}$ is the electronic occupation of the semioccupied natural orbital i.

FR-G1 follows a linear arrangement of three radical sites (Figure 6.13 a) in which $H = -2J(S_1S_2 + S_2S_3)$ and $-2J = \Delta E_{D-Q}$, then $J \approx -4.10$ kcal/mol. In FR-G2 the model employed is star-like (Figure 6.13 b), in which one central radical site interacts with the peripheral ones. It is important to notice that this model assumes the antiferromagnetic interaction of three radicals in the molecular center, *i.e.*, site 3 in Figure 6.13 (b) is equivalent to the interaction on (a). The effective Hamiltonian is write then as $H = -2J(S_3S_2 + S_3S_1 + S_3S_4 + S_3S_5)$ with $J = \Delta E_{Q-D} = -0.78$ kcal/mol. Through these models, the coupling between unpaired electrons can be classified to be moderate antiferromagnetic.

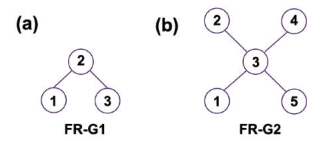


Figure 6.13: Radical-radical interaction models for FR-G1 (a) and FR-G2 (b).

Orbital diagram of the dominant configuration of the ground state of both molecules is depicted in Figure 6.14. (Note: Graphical representation and energy diagrams correspond to molecular orbitals obtained from the ROFH high-spin configuration, while electronic orbital occupancy was obtained at RAS-SF level, as reported in Table 6.4).

Semi-occupied molecular orbitals are delocalized throughout at least two neighbouring fluorenyl moieties. Accordingly, spin density of the doublet (quartet) ground state of **FR-G1** (**FR-G2**) is spread homogeneously over the whole fluorenyl core, illustrating the two-dimensional π -conjugation through the branches. Most of the spin density is localized at the 9-position carbons (Figure 6.15), *i.e.* the radical sites in the cyclopentadienyl radicals.

Electronic occupation of the lowest unoccupied natural orbital (Table 6.4) allows the characterization of the polyradical character of FR-G1 and FR-G2. While FR-

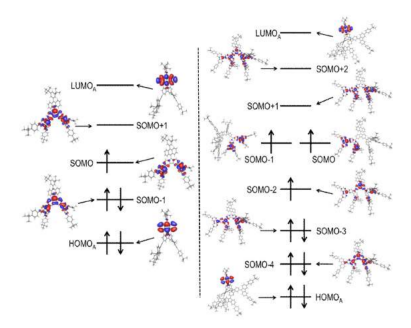


Figure 6.14: Ground state molecular orbital diagram of **FR-G1** (left) and **FR-G2** (right).

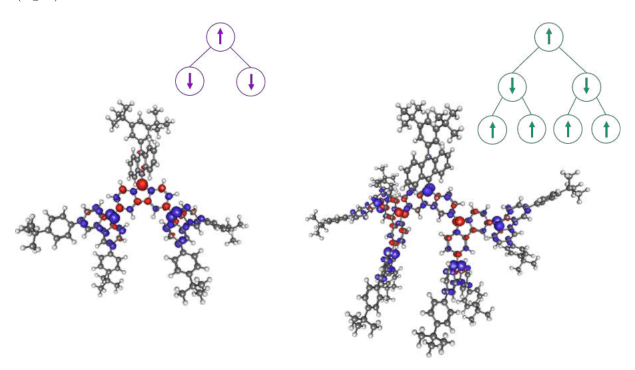


Figure 6.15: Spin density distribution of **FR-G1** (left) and **FR-G2** (right) computed at their respective ground states (doublet and quartet) at the UB3LYP/6-31G* level of theory.

G1 shows moderate triradical character, as is indicated by $y_0 = 0.37$, in the **FR-G2** can be regarded as an heptaradicaloid, with $y_0 = 1.00$, $y_1 = 0.58$, and $y_2 = 0.50$.

All data exposed so far reinforce the moderated AFM exchange interaction be-

tween fluorenyl units in both polyradicaloid dendrons. **FR-G1** and **FR-G2** thus represent the first class of two-dimentionally AFM coupled dendritic polyradicaloids of first and second generation.

Moderate polyradical character provides the feasibility to get states of different multiplicities in a rather narrow window of energy. Main configuration of D_0 (doublet ground state) of **FR-G1** is represented by orbital diagram in Figure 6.14. D_1 (excitation energy 5.26 kcal/mol and $N_u = 2.08$) and Q_1 states correspond to single electron excitation from D_0 within the SOMO-1, SOMO and SOMO+1 space, as $n_{SONO\pm i}$ in Table 6.4 indicates.

Excited states in Table 6.4 can be understood as the result of radical-radical interactions with energies approximated as:

$$E(D_0) = J, (6.2)$$

$$E(D_1) = 0, (6.3)$$

$$E(Q_1) = -J. (6.4)$$

Likewise, Q_0 of **FR-G2** mainly corresponds to the single electron occupation of SOMO-1, SOMO and SOMO+1, as displayed in Figure 6.14 as well as D_1 . Similarly, D_2 to D_4 and Q_2 to Q_3 are obtained as the single excitation from D_1 or Q_0 state, respectively, involving orbitals from SOMO-2 to SOMO+2. D_5 and D_6 are excitations from D_2 , D_3 and D_4 configurations involving the entire set of SOMOs. Due to the moderate AFM coupling, all low-lying states exhibit a high N_u values. The octuplet state is found at higher energies with transitions involving orbitals localized on the anthracene moieties.

Another significant result to highlight is the moderate AFM coupling demonstrated by all the macrostructures, despite their connectivity. AFM allows them to be stable open shell systems with a large number of accessible excited states.

6.3 Appendix

Ground state of all possible multiplicities were explored with unrestricted or restricted DFT, according with the system. Then, table X shows **FR-G1** and **FR-G2** Geometric optimizations were carried out within the DFT framework with restricted and unrestricted approximation of B3LYP functional. In order to determine the ground state geometry, the adiabatic energies of each possible spin multiplicity were computed at DFT level, this means when n = 3 lowest doublet and quartet state, for n = 4 the lowest singlet, triplet and quintet states, etc. [10]MC.

Polyradical character was analysed by the number of unpaired electrons (N_u) and electronic occupation of natural orbitals. Use of C-DFT as tool to confirm the relative stability between specific spin configurations and

State	Energy	N_u	n _{SONO-3}	n_{SONO-2}	n_{SONO-1}	n _{SONO}	n_{SONO+1}	n _{SONO+2}	n_{SONO+3}
Q_0	0.00	5.23	1.50	1.42	1.00	1.00	1.00	0.58	0.50
D_1	0.78	4.71	1.64	1.42	1.12	1.00	0.88	0.58	0.36
D_2	2.47	4.32	1.62	1.54	1.22	1.00	0.78	0.46	0.38
D_3	2.65	4.38	1.60	1.52	1.22	1.00	0.78	0.48	0.40
D_4	2.77	4.90	1.47	1.37	1.24	1.00	0.75	0.63	0.53
Q_1	6.04	5.50	1.43	1.24	1.10	1.00	0.89	0.76	0.57
Q_2	6.09	5.68	1.37	1.22	1.10	1.00	0.90	0.78	0.63
Q_3	6.11	5.59	1.41	1.26	1.06	1.00	0.93	0.74	0.59
D_5	6.32	5.08	1.61	1.37	1.01	0.99	0.99	0.63	0.39
D_6	6.53	5.56	1.43	1.22	1.10	1.00	0.90	0.78	0.56
O_1	58.18	7.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00

Table 6.5: Calculated relative vertical energies (kcal/mol) of **FR-G2** for different spin states, and the number of unpaired electrons (N_u) and electronic occupation of SONOs at the quartet ground state (n_{SONO-3} , n_{SONO-2} , n_{SONO-1} , n_{SONO+1} , n_{SONO+2} , and n_{SONO+3}).

Table 6.6: Vertical energy gaps (kcal/mol) between ground and low-lying states of different multiplicities computed at the $B3LYP/6-31G^*$ level.

PTM-n	state	Multiplicity	ΔE	S^2
PTM-3				
	D_0	Doublet	0.000	1.83
	Q_1	Quartet	0.066	3.82
PTM-4				
	S_0	Singlet	0.000	2.11
	T_1	Triplet	0.076	3.10
	Q_1	Quintet	0.154	6.09
PTM-5				
	D_0	Doublet	0.000	2.88
	Q_1	Quartet	0.090	4.88
	S_1	Sextet	0.182	8.87
PTM-6				
	S_0	Singlet	0.000	3.16
	T_1	Triplet	0.071	4.15
	Q_1	Quintet	0.506	7.14
	S_1	Septet	0.940	12.13

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Chapter 7

Singlet fission

Singlet fission (SF) is a photophysical process in which one singlet exciton shares its energy with a molecule in the ground state, yielding the formation of two triplet excitons. SF term was conceived to explain the generation of triplet states in anthracene crystals in 1965. (1) Later on, the same mechanism was detected in crystalline tetracene. (2) At that moment, several theoretical efforts were done to develop a model to explain the kinetics of the process, (3; 4?) while the experiments were focused on go further in the understanding of the mechanism and exploration of new materials. (5) Despite, the interesting advances in the field during the decade of 1990, the interest for SF gain a lot of attention until the early 2000s, when Hanna and Nozik suggested SF as a possible strategy to overcome the efficiency barrier in solar-cells. (6; 7) Since then, extensive theoretical and experimental work has been done in order to reveal the deepest details ruling the process. (8; 9; 10)

Involved states, mechanisms, and some conditions have been explored and stablished. Organic materials able to undergo this process efficiently have been investigated, providing valuable information to the design of the new ones with improved capabilities that allows SF research to reach its main goal: inclusion of SF materials in the third generation of solar cells. Experimentally, solar cells incorporating SF materials have reached above 100 % of external quantum yield efficiency and up to 200% of internal quantum efficiency.(11; 12)

Through this chapter, I will explain first the basic concepts to understand the singlet fission process followed by the explanation of the well-accepted five-state model. Finally, I will show how the expansion of this model to consider the role of two more states provides some light in the mechanistic aspect, but also can help to improve the design of singlet fission chromophores.

7.1 Singlet fission in a nutshell

SF is a down conversion process in which the excitation energy of one singlet exciton is shared to produce two "independent" triplet excitons.(9; 10) The singlet fission

process can be described in two steps as follows:

$$S^* \rightleftharpoons {}^{1}TT \rightleftharpoons T_1 + T_1. \tag{7.1}$$

The first step starts with the singlet exciton formed (no matter the source) while ends with the formation of the multiexcitonic state (¹TT). The second step is related to the decoupling of the triplet-triplet pair.

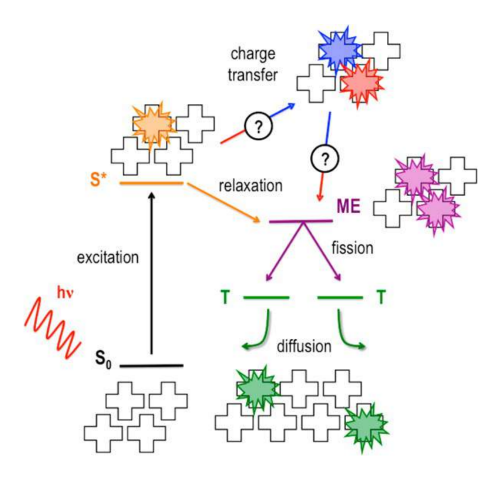


Figure 7.1: Overview of the singlet fission process. Ground state (a), excitation (1), optical exciton (b), relaxation (2), triplet-pair state (c), charge separation (2'), charge transfer states (c'), fission (3), independent triplets (d), diffusion (4). Image adapted from (13).

The whole process can be described as the succession of photophysical events, as illustrated in Figure 7.1. The activation is usually, but not only, achieved by the absorption of one photon leading to the singlet excitation; then the multiexcitonic state is formed thorought a internal conversion process since the triplet-triplet state is coupled as an overall singlet. Then, by the fission of this state, two triplets localized over different monomers are generated and, in excited systems, migration through neighbour chromophores can occur. The SF process requires some energetic

conditions to be fulfilled:

$$E(S_1) \ge 2E(T_1). \tag{7.2}$$

The thermodynamic condition establishes that the energy of the singlet exciton should be at least twice the energy of the triplet (Equation 7.2). This ensures the exothermicity of the whole process. But it is not enough to fulfil the Equation 7.2, to ensure that the process is efficient, $E(S_1) - 2E(T_1) \sim 0$, otherwise if the reaction is exothermic the fission process is kinetically inefficient. On the other hand, if the energy of S_1 is slightly lower than $2T_1$, SF can be carried out as an endothermic process, as observed in tetracene crystals.(1)

$$E(T_2) > 2E(T_1),$$
 (7.3)

$$E(Q_1) > 2E(T_1).$$
 (7.4)

Equation 7.3 and 7.4 are related to the triplet-triplet recombination, which after singlet fission is desirable to avoid. Concretely, the energy of the second triplet (T_2) must be higher than twice the energy of T_1 to discard the triplet-triplet recombination to the triplet manifold. The population of Q_1 is not often observed due to the high energy of this state, and can be related to the tetraradical character of the SF chromophore, which is detrimental to the process.

In addition to the state energies of the monomer, the spatial arrangement of the chromophores is very important. The relative orientation between π -conjugated systems can lead to the formation of aggregates that modify the energy of S_1 , giving rise to more accessible deactivation paths than SF. It has been observed that co-valently linked dimers are beneficial to modulate the electronic interaction between monomers by imposing restrictions to their relative orientation without affecting their photophysical properties. (14; 15; 9)

7.1.1 Singlet fission chromophores

Organic molecules with very small to moderate diradical character have been proved to fulfil the energetic requirements of SF, such is the case of pentacene (according with our calculations in Chapter 4, it holds a moderate diradicaloid character).

Actually, one of the fundamental topics in the field, is the design of new chromophores to increment the efficiency of singlet fission and allow the expansion of its applications.

The minimum model for SF is a dimer, to have a system that can localize both triplets separately. Nevertheless, the properties of the monomer, *i.e.*, relative energy of excited states, absorption coefficients, structure, radical character, and chemical photostability, have to be considered in order to have a chromophore appropriate for

SF. Historically, polyacenes have been extensively explored as SF materials, since they fulfil nicely the energetic and structural requirements, concretely, tetracene and pentacene. In longer polyacenes the stability is not good as exposed in the previous Chapter.

There are three main classes of organic chromophores that can be found in the literature, classified according to the ordering of excited singlet states (Figure 7.2).

7.1.1.1 Class I

In this class of chromophores, the lowest excited singlet state corresponds to the optical transition, which is characterized as the HOMO-to-LUMO excitation. Higher singlet states are considerable energetically above it, avoiding mixing or additional decaiment channels. Compounds in this class hold a closed-shell configuration. Tetracene, anthracene and perylene are chromophores of this class that have been investigated to carry out singlet fission.

7.1.1.2 Class II

The optically allowed excitation is to the S_2 state in chromophores of class II. Configuration of S_1 corresponds to the linear combination to the HOMO-1-to-LUMO and HOMO-to-LUMO+1 excitations. S_1 is energetically found slightly below S_2 , and can be populated by an internal conversion process. SF chromophores of this class are naphthalene, pyrene and phenanthrene.

7.1.1.3 Class III

In chromophores of class III, as in those of class II, the second excited singlet is the optically allowed, while the energy gap respect to the S_1 state is quite small. S_2 involves HOMO-to-LUMO transition and S_1 is characterized as the double excitation HOMO²-to-LUMO². This doubly excited state is very common in π -conjugated systems like polyenes, where 1B_u and 1A_g excited states are almost degenerated.

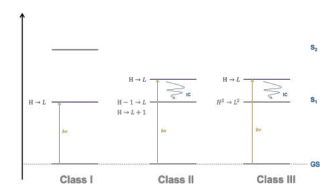


Figure 7.2: Excited state diagram of singlet fission chromophores of class I (a), II (b) and III (c). H = HOMO, L= LUMO.

The relative energy between T_1 and S_1 must be modulated to fulfil the Equation 7.2. Energy splitting between both states can be approximated to twice the HOMO-LUMO exchange integral ($\Delta E_{ST} \approx 2K_{hl}$), which depends on the repulsion of two electrons located in different orbitals and its magnitude is related to the spatial overlap of HOMO and LUMO. There are two main manners to tune ΔE_{ST} and open the gap between S_1 and T_1 : i) destabilization of S_1 (increment of K_{hl}) in class I, since in closed-shell molecules the $\Delta E_{S_1T_1}$ is smaller compared with $\Delta E_{S_0T_1}$. This can be achieved by means of inclusion of heteroatom or increasing the size of the conjugated system. ii) stabilization of the T_1 state, *i.e.*, decreasing $\Delta E_{S_0}T_1$, by inducing low to moderate diradical character in chromophores of class I and III. (9) Systems with strong diradical character are detrimental for the SF efficiency, since SF becomes too exothermic, resulting in a large energy loss in the fission process.

7.2 Five-state model

Normally, within the dimer model, the first step of the SF process is described within a five-state model, including the singlet excited state $(S_1S_0 \text{ or } S_0S_1)$, and the multiexcitonic state (^1TT) , as indicated in Equation 7.1. Additionally, charge transfer states are included as potential intermediates or assisting states of the process.

7.2.1 Electronic states

7.2.1.1 Singlet excited states

The S_1 state is assumed to be the lowest excited singlet state of the SF chromophore, usually described by the HOMO-to-LUMO transition. Within the SF context, it is expected than if the optical state is not the lowest one, S_1 will be populated before the fission through internal conversion. Still, exist a small possibility to higher singlets (S_n) undergo SF.

7.2.1.2 Charge transfer states

First, it is important to clarify the difference between charge transfer (CT) and charge resonance (CR) states. Both kind of states are related to the exchange of electrons to form the cation/anion (CA or AC) pair. CT is referred to the formal migration of one electron from one monomer to the other, while CR indicates the presence of charged configurations in the wave function, but does not imply the formation of charged chromophores. In other words, it is possible to describe the CR state as the linear combination of CT states. The big impact of these states to SF process has been extensively studied and the most common SF mechanism is the one mediated by charge transfer states.(15)

7.2.1.3 The triplet-pair state

The core of the singlet fission reaction is the formation of the multiexcitonic tripletpair state (1 TT). According to the coupling of two angular moments, the triplet-pair leads to the formation of 9 microstates, 1 TT (1), 3 TT (3), and 5 TT (5). The most relevant in SF is the one corresponding to the triplet-pair coupled as an overall singlet. In this way, singlet fission is a process allowed by spin symmetry rules, since no change of multiplicity between $S_{1}S_{0}$ and 1 TT states occurs. Normally, 1 TT is more stable than 3 TT and 5 TT, and the extra stabilization can be described as the needed energy to dissociate two triplets, specifically:

$$E_b \approx {}^5TT - {}^1TT. \tag{7.5}$$

⁵TT can be understood as 4 unpaired electrons (two on each triplet), while ¹TT has a larger tendency to mix with other configurations (as local excitations or charge transfer states).

This triplet-pair state was first computationally characterized through wave function based methods. (16?) Later on, it was experimentally detected by Chan and collaborators (17) as a SF intermediate. Since then, the nature and properties of the coupled double triplet state have been largely investigated both experimentally and theoretically by several authors. (18; 15; 19; 20)

The minimal configuration space necessary to describe the S_1S_0 , S_0S_1 , AC, CT, and 1TT states must contain the two frontier orbitals (HOMO and LUMO) of the two monomers. Low-lying excited singlet states of molecular dimers can be build in terms of local excitations to S_1 and ionic configurations corresponding to the cationanion (CA) and anion-cation (AC) charge transfer (CT) states, and the triplet-pair state (1TT). In the HOMO/LUMO frozen orbital approximation for equivalent homodimers,(?) these diabatic states are expressed as:

$$|S_1S_0\rangle = \frac{1}{\sqrt{2}} \left(|h_A\bar{l}_Ah_B\bar{h}_B\rangle + |l_A\bar{h}_Ah_B\bar{h}_B\rangle \right),$$
 (7.6)

$$|S_0S_1\rangle = \frac{1}{\sqrt{2}} \left(|h_A\bar{h}_Ah_B\bar{l}_B\rangle + |h_A\bar{h}_Al_B\bar{h}_B\rangle \right), \tag{7.7}$$

$$|CA\rangle = \frac{1}{\sqrt{2}} \left(|h_A \bar{l}_B h_B \bar{h}_B\rangle + |l_B \bar{h}_A h_B \bar{h}_B\rangle \right),$$
 (7.8)

$$|AC\rangle = \frac{1}{\sqrt{2}} \left(|h_A \bar{h}_A l_A \bar{h}_B \rangle + |h_A \bar{h}_A h_B \bar{l}_A \rangle \right), \tag{7.9}$$

$$|^{1}\text{TT}\rangle = \frac{1}{\sqrt{3}}[|h_{A}l_{A}\bar{h}_{B}\bar{l}_{B}\rangle + |\bar{h}_{A}\bar{l}_{A}h_{B}l_{B}\rangle - \frac{1}{2}\left(|h_{A}\bar{l}_{A}h_{B}\bar{l}_{B}\rangle + |l_{A}\bar{h}_{A}l_{B}\bar{h}_{B}\rangle - |h_{A}\bar{l}_{A}l_{B}\bar{h}_{B}\rangle - |l_{A}\bar{h}_{A}h_{B}\bar{l}_{B}\rangle\right)], \quad (7.10)$$

where h_X (l_X) is to the HOMO (LUMO) of molecule X, the upper bar indicates spin- β orbital. The exciton Hamiltonian in the basis of orthonormal diabatic states is expressed as:

$$H_{ex} = \sum_{I < J} |I\rangle H_{IJ}\langle J|, \tag{7.11}$$

where $H_{IJ} \equiv \langle I|\hat{H}|J\rangle$ are defined as the electronic energies (I=J) and couplings $(I \neq J)$. The electronic coupling between the five electronic states is:

$$\langle S_1 S_0 | \hat{H} |^1 TT \rangle = \sqrt{\frac{3}{2}} \left[(l_A h_B | l_A l_B) - (l_B h_A | h_A h_B) \right],$$
 (7.12)

$$\langle S_1 S_0 | \hat{H} | S_0 S_1 \rangle = 2(l_A h_A | l_B h_B) - (l_A l_B | h_A h_B),$$
 (7.13)

$$\langle S_1 S_0 | \hat{H} | CA \rangle = (l_A | \hat{F} | l_B) + 2(h_A l_B | h_A l_A) - (h_A h_A | l_A l_B),$$
 (7.14)

$$\langle S_1 S_0 | \hat{H} | AC \rangle = -(h_A | \hat{F} | h_B) + 2(h_A l_B | h_B l_B) - (h_A h_B | l_B l_B),$$
 (7.15)

$$\langle {}^{1}\text{TT}|\hat{H}|\text{AC}\rangle = \sqrt{\frac{3}{2}} \left[(l_{A}|\hat{F}|h_{B}) + (l_{A}h_{B}|l_{B}l_{B}) - (l_{A}h_{B}|h_{A}h_{A}) \right], \quad (7.16)$$

$$\langle \operatorname{CA}|\hat{H}|\operatorname{AC}\rangle = 2(l_B h_A | l_A h_B) - (l_A l_B | h_A h_B), \tag{7.17}$$

(7.18)

where orthonormal orbitals have been considered $(\langle i|j\rangle = \delta_{ij})$, $(i|\hat{F}|j)$ is the Fock matrix element between orbitals i and j, and (ij|kl) is a two-electron interaction integral written as:

$$(ij|kl) = \int \phi_i^*(\mathbf{r}_1)\phi_j(\mathbf{r}_1) \frac{1}{r_{12}} \phi_k^*(\mathbf{r}_2)\phi_l(\mathbf{r}_2) d\mathbf{r}_1 d\mathbf{r}_2.$$
 (7.19)

Notice that for symmetric homodimers $\langle IJ|\hat{H}|KL\rangle = \langle JI|\hat{H}|LK\rangle$, with $|IJ\rangle$ corresponding to any of the diabatic states of the dimer built as the coupling of monomeric states $|I\rangle$ and $|J\rangle$ in equations 7.6-7.10.

In the following, I use the co-planar ethylene dimer as a model system. Molecular geometry of each ethylene molecule has been frozen at the MP2/cc-pVTZ ground state D_{2h} optimized structure with CC and CH bond lengths of 1.3320 and 1.0804 Å, respectively, and HCC angle of 121.3°. The conformational space considered corresponds to the two ethylene molecules stacked at 3.5 Å with different relative in-plane displacements along the long and short molecular axes (Figure 7.3).

Computed electronic couplings for the co-planar conformational space of the ethylene dimer (Figure 7.4) are in very good agreement to similar calculations performed by Smith and Michl.(21) The coupling between S_1S_0 (and S_0S_1) with ¹TT state, which is optimal at the alternate disposition along the bond axis (x coordinate), is 1-2 orders of magnitude smaller than those of S_1S_0 or ¹TT to charge separated configurations (540-740 meV absolute maxima), since the latter contain

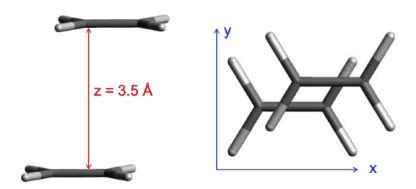


Figure 7.3: Side (left) and top (right) views of the co-planar ethylene dimer employed in this work. Vertical intermolecular distance (z = 3.5 Å) indicated in red.

single electron Fock matrix elements (equations 7.14-7.16), while the (direct) interaction is obtained as the difference between two bielectronic integrals (equation 7.12). Such strong coupling of the initial and final states to charge separation states, is the driving force for CT-mediated singlet fission mechanism invoked in numerous studies.(? 22?; 23)

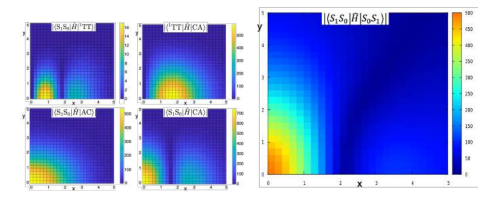


Figure 7.4: Absolute electronic couplings (in meV) between S_1S_0 , ¹TT, CA, and AC diabats for the stacked ethylene dimer computed at a constant vertical separation (z = 3.5 Å) and different longitudinal (x in Å) and horizontal (y in Å) displacements as shown in Figure 7.3.

7.3 Seven-state model

Beyond the extensively studied states, S_1S_0 , 1TT , T_1 and CT, in organic conjugated molecules other electronic states can be close in energy and be available to be populated and play an active or secondary role in the SF process. Such is the case of chromophores in class III, in which the doubly excited (dark) state is almost degenerated to the bright state. The dark state has been characterized to be mainly the double excitation from HOMO-to-LUMO and becomes the lowest excited state in polyenes larger than but adiene. (24; 25). Herein, this state is named

as doubly excited state ("D state"). Within the framework of SF it is important to stress the difference between the D (HOMO² \rightarrow LUMO²) and the ¹TT states, even when both are not optically active (dark states), and D state has been described as a coupled triplet-pair,(26; 27; 28) the electronic coupling between triplets in ¹TT is significantly weaker compared with the triplet-triplet electronic coupling in the D state. Moreover, while ¹TT exist as a delocalized state holded by at least two choromopores, D state is a monomeric doubly excited state. Its participation in the SF mechanism was suggested to occur in pentacene,(16) although this possibility was later disregarded by means of experimental measurements(29; 30) and accurate electronic structure calculations(22) indicating that the optical state lies below the dark state.

Here an extension of the SF model (previous section) is presented by the inclusion of the D state. The possible role in the ¹TT formation: i) direct when is the lowest excited state; or 2) as a mediator when is a higher state, is explored by means of the analysis of the electronic couplings and the time evolution of excited states.

7.3.1 Electronic couplings with the D state

Within the simple model of SF states described in previous section (HOMO and LUMO of each monomer), the DS_0 and S_0D states, monomeric HOMO to LUMO double excitation, are expressed as:

$$|DS_0\rangle = |l_A\bar{l}_Ah_B\bar{h}_B\rangle,$$
 (7.20)

$$|S_0D\rangle = |h_A\bar{h}_Al_B\bar{l}_B\rangle.$$
 (7.21)

(7.22)

The energy of the D state under this approximation is computed as $E(DS_0) \approx E(D) + E(S_0)$, and $E(^1TT) \approx 2E(T_1)$, the relative energy of DS_0 with respect to 1TT and S_1S_0 is then:

$$E(DS_0) - E(S_1S_0) \approx \varepsilon_l - \varepsilon_h - 3(ll|hh) + (ll|ll),$$
 (7.23)

$$E(DS_0) - E(^{1}TT) \approx (ll|ll) - (hh|hh) + 2[(ll|hh) + (lh|lh)].$$
 (7.24)

The variables used here follow the definition and nomenclature as in Equation 7.19. Within the HOMO-LUMO model, the electronic couplings between SF states (equations 7.6-7.10) and D state can be calculated as:

$$\langle DS_0|\hat{H}|^1TT\rangle = -\sqrt{3}(h_A h_B |l_A l_B), \qquad (7.25)$$

$$\langle DS_0 | \hat{H} | S_1 S_0 \rangle = \sqrt{2} \left[(l_A |\hat{F}| h_A) + (h_A l_A | l_A l_A) - (h_A l_A | h_A h_A) \right], \quad (7.26)$$

$$\langle DS_0|\hat{H}|CA\rangle = \sqrt{2}(l_A h_A | l_A l_B), \qquad (7.27)$$

$$\langle DS_0|\hat{H}|AC\rangle = -\sqrt{2}(l_A h_A | h_A h_B), \qquad (7.28)$$

$$\langle \mathrm{DS}_0 | \hat{H} | \mathrm{S}_0 \mathrm{D} \rangle = \langle \mathrm{S}_0 \mathrm{D} | \hat{H} | \mathrm{DS}_0 \rangle = 0.$$
 (7.29)

Equations 7.12-7.17 and 7.25-7.29 complete the electronic Hamiltonian of the seven-state model. Using the same grid of the displaced co-planar ethylene dimer (Figure 7.3), the coupling between DS_0 and S_1S_0 , 1TT , CA and AC diabatic states is represented in Figure 7.5.

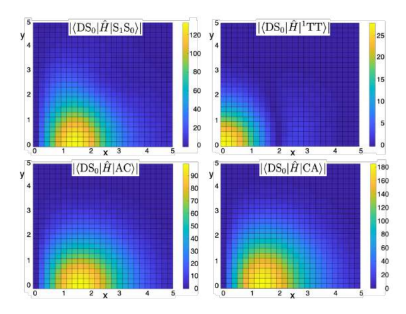


Figure 7.5: Absolute electronic couplings (in meV) between DS₀ and S₁S₀, ¹TT, CA, and AC diabats for the stacked ethylene dimer computed at a constant vertical separation (z = 3.5 Å) and different longitudinal (x in Å) and horizontal (y in Å) displacements as shown in Figure 7.3

The doubly excited localized state shows sizeable couplings with S_1S_0 , 1TT , CA, and AC states. The DS_0/S_1S_0 couplings present the strongest interaction at the longitudinal displacement (y = 0.0 Å), with eclipsed alternate carbons, where the electronic interaction is 133 meV. The couplings of DS_0 to AC and CA show very similar conformational profiles, with the strength of the interaction to CA being larger than for S_1S_0 and about twice the coupling to AC. Interestingly, the couplings to the triplet-pair state are maximum at the eclipsed structure, since it simultaneously maximizes the overlaps between localized HOMOs and LUMOs (equation 7.25). The computed interaction at the D_{2h} arrangement ($\sim 30 \text{ meV}$) is considerably stronger than the optimal coupling between S_1S_0 and 1TT (Figure

7.4). This result can be rationalized by comparing equations 7.12 and 7.25, where the former corresponds to the difference between two bielectronic integrals, resulting in a rather small overall interaction.

7.3.2Singlet fission couplings

Even if the D state lies energetically above the optical transition, it can be directly or indirectly involved in the SF process, likewise charge separated configurations are able to catalyze the formation of the ¹TT state through the CT-mediated mechanism.

Normally, the reaction rate for the formation of the triplet-pair can be approximated by the Fermi's Golden Rule:(31)

$$k_{SF} = \frac{2\pi}{\hbar} |V_{SF}|^2 \delta(E_I - E_F), \tag{7.30}$$

where I and F subscripts refer to initial and final states, respectively, and $\delta(E_I - E_F)$ ensures energy conservation. The electronic coupling in SF (V_{SF}) can be perturbatively expanded to second order to take into account the participation of one mediator state, which corresponds to the CT (CA and AC) states (five-state model). To model the case in which both states, CT and D (or S_0S_1), can act as mediator, the V_{SF} can be expanded to third order:

$$V_{SF} = \sum_{n=1}^{\infty} V_{IF}^{(n)}, \quad (7.31)$$

$$V_{IF}^{(1)} \equiv V_{IF} = \langle I|\hat{H}|F\rangle, \quad (7.32)$$

$$V_{IF}^{(1)} \equiv V_{IF} = \langle I|\hat{H}|F\rangle, \quad (7.32)$$

$$V_{IF}^{(2)} = \sum_{K \neq I,F} \frac{V_{IK}V_{KF}}{(E_I + E_F)/2 - E_K}, \quad (7.33)$$

$$V_{IF}^{(3)} = \sum_{K,L \neq I,F} \frac{V_{IK}V_{KL}V_{LF}}{(E_I - E_K)(E_F - E_L)} - V_{IF} \sum_{K \neq I,F} \frac{V_{IK}V_{KF}}{(E_I - E_K)(E_F - E_K)}, \quad (7.34)$$

where $E_K = \langle K | \hat{H} | K \rangle$. Such expression has been employed in SF to discern between different mechanisms for the formation of the triplet-pair state depending on the energy distribution of electronic states and the coupling between them. The direct transition from initial state (n = 1) depends only on the direct coupling between 1 TT and $S_{1}S_{0}$, in our model system $\langle S_{1}S_{0}|\hat{H}|^{1}$ TT \rangle is maximazied when the ethylene dimer is slipped in the x-axis. On the other hand, the mediated mechanism through virtual states (K states can be CT and D) is described when n=2. In fact, the mediated mechanism has shown to be dominant in many systems, in which the CT states (CA and AC) act as virtual mediators due to the strong couplings with S₁ and ¹TT states. In Figure 7.4 it can be noticed that the couplings with CT states are considerably bigger than the direct one. Additionally, in the seven-state model the L and K can be CT (AC and CA) and D. In such way, Equation 7.34 represents the mechanism mediated simultaneously by both states.

If D is the lowest molecular excited state, the first step of SF can occur directly from it, in such case the activation of D occurs as consequence of the fast exciton decay after photoabsorption by the optical state (S*) populating the D state and then the formation of ¹TT state as:

$$S^* \longrightarrow D \stackrel{S_1, CT}{\rightleftharpoons} {}^1TT.$$
 (7.35)

Generation of ¹TT from D can take place thought different microscopic mechanisms, that can involve, or not, other electronic states if they are close in energy. Further, as has been observed with the CT states the S*-CT-¹TT couplings determine their participation in SF process. The possibility and the relative importance of different decay paths was explored by computing the total couplings (equation ??-7.34) for the conformational space of the co-planar ethylene dimer considering different relative energetic ordering of states.

First, if D and 1 TT states are much lower than the rest of singlet excitations in the system, second (and third) order terms in equation 7.31 can be disregarded, and the only available channel is the direct transfer from D to the triplet-pair state. In that case, the rate of 1 TT formation is expected to be controlled by the direct coupling, which is favored in the eclipsed conformation (Figure 7.5). It is worth noticing that the eclipsed form also corresponds to the most stable arrangement of excimeric states,(?) since it produces the strongest exciton coupling ($\langle S_1S_0|\hat{H}|S_0S1\rangle$ in Figure 7.4). Therefore, if the S_1 state is not much higher in energy than the D and 1 TT states the excimer formation could compete with SF from D state. The feasibility between excimer and triplet-pair state formation will be largely controlled by the relative energies of S_1 and D states of the chromophore.

Alternatively, if there are other excitations lying not too high in energy, such as the local exciton (S_1) or CT states, they could mediate the SF mechanism from D state (Figure 7.6).

Despite that DS_0/S_1S_0 couplings are rather strong (Figure 7.5), mediation by S_1 states is effectively hindered by the weak $S_1S_0/^1TT$ interaction, and SF from D with low-lying S_1 states is still dominated by the direct mechanism (Figure 7.6a). On the other hand, the presence of low-energy CT states has an important impact on SF. If CA and AC configurations are energetically close to D (Figure 7.6b), the CT-mediated mechanism is expected to be the main channel to populate 1TT , with optimal couplings for slip-stacked conformations along the x-axis with a maximum absolute value of ~ 320 meV. If both CT and S_1 states are energetically close to D, the third order couplings simultaneously involving CT and S_1 (CT, S_1 -mediated mechanism) contribute to an important increase of the total SF coupling of slip-stacked conformations of ~ 110 meV with respect to the (second order) CT-mediated

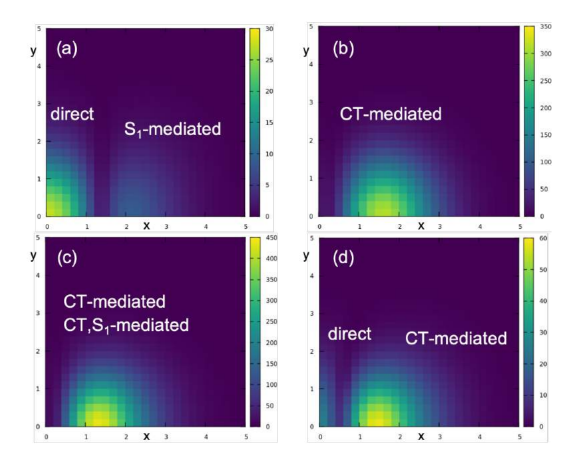


Figure 7.6: Absolute total SF couplings (in meV) between DS₀ and ¹TT with $E(DS_0) - E(^1TT) = 0.25$ eV, and S_1S_0/S_0S_1 and CT states at (a) 0.4 and 20 eV, (b) 20 and 0.4 eV, (c) 0.4 and 0.4 eV, and (d) 0.4 and 3.4 eV above the D state, for the stacked ethylene dimer computed at a constant vertical separation (z = 3.5 Å) and different longitudinal (x in Å) and horizontal (y in Å) displacements as shown in Figure 7.3.

couplings (Figure 7.7).

Probably, the case with small D/S_1 gaps, and with CT states a few eVs above them might represent a realistic energy distribution of molecular dimers, aggregates or crystal structures of organic chromophores with the D state below S_1 . Interestingly, in that situation we might have both SF mechanisms available, with direct transition from D favored for eclipsed dimers and CT-mediated formation of ¹TT in displaced arrangements (Figure 7.8).

The fact that the D state cannot be connected to the ground state through the dipole moment operator (dark state) might be an advantage for the SF reaction. Contrary to singlet fission from the HOMO \rightarrow LUMO optical singlet (S₁), radiative decay to the ground state (fluorescence) cannot compete with the formation of the triplet-pair, potentially resulting in increased fission efficiencies.

In many organic chromophores the doubly excited state lies above the S_1 excitation as in class II chromophores. Hence, the SF photophysical reaction will occur

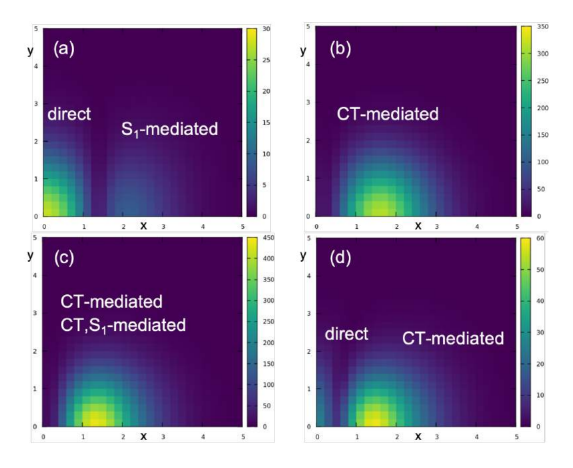


Figure 7.7: Absolute total, first, second and third order SF couplings (in meV) between DS₀ and 1 TT with $E(DS_{0}) - E(^{1}$ TT) = 0.25 eV, and S₁S₀/S₀S₁ and CT states at 0.4 and 0.4 eV above the D state, for the stacked ethylene dimer computed at a constant vertical separation (z = 3.5 Å) and different longitudinal (x in Å) and horizontal (y in Å) displacements as shown in Figure 7.3.

from the S_1 exciton, while the D state, if energetically not too high, might play an indirect role as a virtual state catalyzing the formation of the triplet-pair state, similar to the CT-mediated SF mechanism:

$$S_1 \stackrel{D,CT}{\rightleftharpoons} {}^1TT$$
 (7.36)

Direct S_1 to 1TT coupling (Figure 7.4) is of the order of tens of meV or weaker, with a maximum at the alternate conformation (with x displacement of half of the carbon-carbon bond distance, ~ 0.8 Å). Our calculations confirm that doubly excited states can mediate the transition to 1TT through second order contributions (equation 7.33). Interestingly, both direct and D-mediated mechanisms are maximum and in phase at the alternate geometry (with second order D-mediated couplings being stronger than $S_1/{}^1TT$ direct interactions), in which they cooperatively assist the formation of the triplet-pair state with a rather strong overall SF coupling (~ 60 meV in Figure 7.9).

As shown in numerous occasions, (32; 23?; 22; 33) CT-mediated SF can be rather

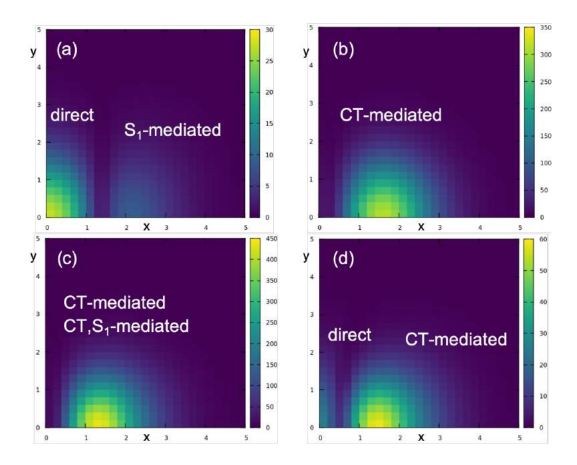


Figure 7.8: Absolute total, first, second and third order SF couplings (in meV) between DS₀ and 1 TT with $E(DS_{0}) - E(^{1}$ TT) = 0.25 eV, and S₁S₀/S₀S₁ and CT states at 0.4 and 3.4 eV above the D state, for the stacked ethylene dimer computed at a constant vertical separation (z = 3.5 Å) and different longitudinal (x in Å) and horizontal (y in Å) displacements as shown in Figure 7.3.-split d) 1-2-3

efficient, with strong SF couplings achieved through superexchange, i.e., second order contributions to the $S_1/^1TT$ coupling. When CT states are close to S_1 , $|V_{SF}|$ is as high as 706 meV (Figure 7.10 at 1.0 Å). Due to the strength of S_1/CT and $CT/^1TT$ couplings, second order CT terms largely determine the overall SF couplings when both D and CT states are energetically low (Figure ??c), with virtually no difference with the $|V_{SF}|$ profile obtained with high energy D state in Figure 7.10.

If D state is energetically close to S_1 , but CT is higher in energy (by a few eVs), the CT-mediated channel is still responsible of the stronger contribution to SF couplings, with direct and CT,D-mediated paths also contributing, although to a lesser degree (Figure ??d). Concretely, the direct interaction exhibits the opposite sign, effectively decreasing the overall $|V_{SF}|$ values, in particular for x < 1 Å. The third order CT,D-mediation is also out-of-phase with respect to superexchange for $0 \le x \le 0.8$ Å, but in-phase for x > 0.8 Å structures, resulting in a maximum coupling shifted to higher longitudinal displacements with respect to the CT-mediated mechanism alone.

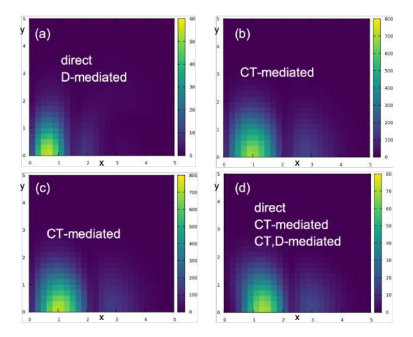


Figure 7.9: Absolute total SF couplings (in meV) between S_1S_0 and 1TT with $E(S_1S_0) - E({}^1TT) = 0.25$ eV, and DS_0/S_0D and CT states at (a) 0.4 and 20 eV, (b) 20 and 0.4 eV, (c) 0.4 and 0.4 eV, and (d) 0.4 and 3.4 eV above the S_1 state, for the stacked ethylene dimer computed at a constant vertical separation (z = 3.5 Å) and different longitudinal (x in Å) and horizontal (y in Å) displacements as shown in Figure 7.3.

7.3.3 Singlet fission dynamics with D state

The potential impact of the D state in the SF dynamics, either as being the initial state for the transition to ¹TT or as a mediator state was evaluated by computing the time evolution of the density matrix of excitonic states through the second order time convolutionless quantum master equation (TCL QME) expressed as:

$$\frac{d}{dt}\rho_{ex}(t) = -i\left[H_{ex}, \rho_{ex}(t)\right] + \sum_{\omega, K} \gamma_K(\omega, t) \left(A_K(\omega)\rho_{ex}(t)A_K^{\dagger}(\omega) - \frac{1}{2}\left\{A_K^{\dagger}(\omega)A_K(\omega), \rho_{ex}(t)\right\}\right),$$
(7.37)

where $\rho_{ex}(t)$ is the density matrix at time t, H_{ex} is the excitonic Hamiltonian in equation 7.11, $A_I(\omega)$ is the I-diabat transition matrix defined as:

$$A_{I}(\omega) = \sum_{K',L'} |K'\rangle\langle K'|I\rangle\langle I|L'\rangle\langle L'|\delta(E_{L'} - E_{K'} - \hbar\omega), \tag{7.38}$$

where prime indices K' and L' run over adiabatic states with energies $E_{K'}$ and $E_{L'}$, and $\gamma_K(\omega, t)$ is the relaxation rate that in the Markov approximation becomes time

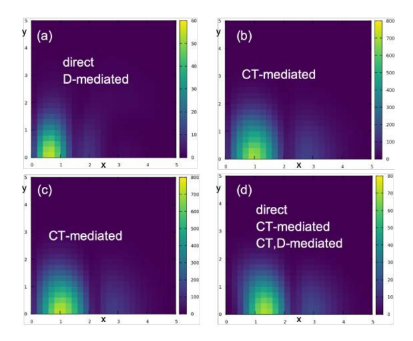


Figure 7.10: Absolute total SF couplings (in meV) between S_1S_0 and 1TT with $E(S_1S_0) - E({}^1TT) = 0.25$ eV, and DS_0/S_0D and CT states at (a) 0.4 and 20 eV, (b) 20 and 0.4 eV, (c) 0.4 and 0.4 eV, and (d) 0.4 and 3.4 eV above the S_1 state, for the stacked ethylene dimer computed at a constant vertical separation (z = 3.5 Å) and different longitudinal (x in Å) and horizontal (y in Å) displacements as shown in Figure 7.3.

independent and can be written as:

$$\gamma_{I}(\omega) = \begin{cases} 2\pi J_{I}(\omega)(1 + n(\omega, T)); \ \omega > 0, \\ \frac{4\lambda_{I}k_{B}T}{\Omega_{I}}; \ \omega = 0, \\ 2\pi J_{I}(-\omega)n(-\omega, T); \ \omega < 0, \end{cases}$$
(7.39)

with the spectral density $J_I(\omega)$ approximated by an Ohmic function with a Lorentz-Drude cutoff:(? ?)

$$J_I(\omega) = \frac{1}{\pi} \frac{2\omega \lambda_I \Omega_I}{\omega^2 + \Omega_I^2},\tag{7.40}$$

where k_B is the Boltzman constant, and $n(\omega, T)$ the Bose-Einstein distribution at temperature T. In the following, I use $\lambda_I = 50$ meV and $\Omega_I = 180$ meV for all diabats I, corresponding to the stretching modes in acene molecules.(?).

In order to evaluate the time evolution, H_{ex} from specific structural conformations of the ethylene dimer previously computed were employed to evaluate Equation 7.37. To simplify, I use S_1 , D and CT state labelling of the dimer to simultaneously refer to S_1S_0/S_0S_1 , DS_0/S_0D and AC/CA pair of states, respectively.

SF dynamics from D

For the stacked eclipsed disposition of the two chromophores, direct coupling between dark states and ${}^{1}\text{TT}$ is maximum, while couplings to S_{1} and CT states become

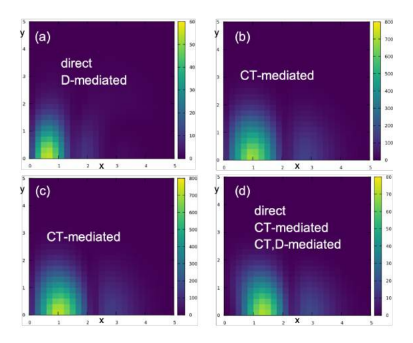


Figure 7.11: Absolute total SF couplings (in meV) between S_1S_0 and 1TT with $E(S_1S_0) - E({}^1TT) = 0.25$ eV, and DS_0/S_0D and CT states at (a) 0.4 and 20 eV, (b) 20 and 0.4 eV, (c) 0.4 and 0.4 eV, and (d) 0.4 and 3.4 eV above the S_1 state, for the stacked ethylene dimer computed at a constant vertical separation (z = 3.5 Å) and different longitudinal (x in Å) and horizontal (y in Å) displacements as shown in Figure 7.3.

zero (Figure 7.5). Thus, at this arrangement of the molecular dimer, generation of the triplet-pair state can only follow a direct mechanism, as shown in Figure 7.12, in which the population of the D state is directly transferred to 1 TT. The first order DS₁/ 1 TT coupling at the D_{2h} symmetry is in the order of 30 meV, resulting in a rather fast formation of 1 TT, which is completed with less than 2 ps.(34)

In reality, the symmetry restrictions on the interstate couplings imposed at the eclipsed disposition can be alleviated due to static or dynamic disorder. Small distortions lowering the D_{2h} symmetry can allow the interaction of D states with CT and S_1 configurations. To simulate such disorder effects, we perform the time evolution for a small longitudinal displacement between the two molecules (x = 0.2 Å in Figure 7.13a). Initially, the system evolves towards the triplet-pair state, as in the ideal D_{2h} symmetry, but very rapidly the D (and 1 TT) state follow internal conversion to the S_1 state. At the nearly eclipsed (and eclipsed) conformation the exciton and superexchange interactions are rather strong (in the order of 500 meV or larger), stabilizing the formation of an excimer-like adiabatic state that delocalizes over the two molecules, and exhibits strong S_1 character and sizeable symmetric contributions of CT configurations ($\sim 10\%$). This state acts as a trap state blocking the formation of 1 TT.

The decay to an excimeric trap state can be circumvented in chromophores with

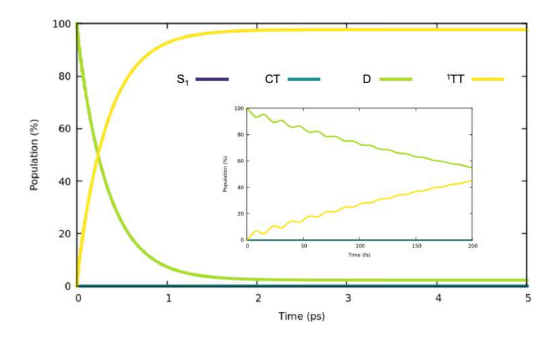


Figure 7.12: Time evolution of state population (in %) for two co-planar eclipsed ethylene molecules (z = 3.5 Å) from DS₀ state, with $E(DS_0) - E(^1TT) = 0.25 \text{ eV}$, and S₁S₀/S₀S₁ and CT states at 0.4 and 3.4 eV. D, S₁ and CT indicate the total population of DS₀/S₀D, S₁S₀/S₀S₁ and CA/AC, respectively. Early time evolution shown in the inset.

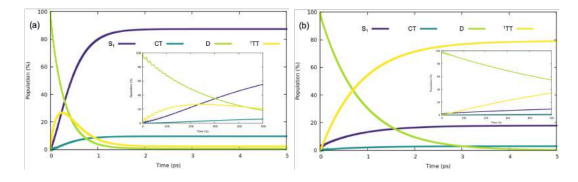


Figure 7.13: Time evolution of state population (in %) for two co-planar ethylene molecules (x = 0.2, y = 0.0, z = 3.5 Å) from DS₀ state, with $E(DS_0) - E(^1TT)$ equals to (a) 0.25 eV and (b) 0.55 eV, and S_1S_0/S_0S_1 and CT states at 0.4 and 3.4 eV. D, S_1 and CT indicate the total population of DS_0/S_0D , S_1S_0/S_0S_1 and CA/AC, respectively. Early time evolution shown in the inset.

a larger relative energy gap between S_1 and 1TT . In Figure 7.13b we simulate the exciton dynamics corresponding to the same situation as in Figure 7.13a, but with a larger exothermicity for the SF process from the dark state, that is from 0.25 eV to 0.55 eV as evaluated with the state energies of the monomer $(E(D) - 2E(T_1))$. Now, excimer stabilization due to exciton (S_1S_0/S_0S_1) and superexchange (S_1/CT) couplings is not large enough to overcome the $S_1/^1TT$ diabatic energy gap, and the

triplet-pair is the lowest adiabatic state. It is worth noticing that in this case 1 TT mixes with S_{1} ($\sim 18\%$) and CT ($\sim 3\%$) configurations.

As illustrated in Figure 7.5, slip-stack configurations do not favor the direct coupling, and as a consequence the direct mechanism is also unfavorable. On the other hand, the dark state presents quite strong interactions with excitonic (S_1) and CT states, with S_1/CT couplings being also strong. Overall, SF couplings predict CT and S_1 mediated processes (Figure 7.6). As a result, the excited state time evolution from the dark state in the x=1.4 Å dimer exhibits a rather different population profile than in the eclipsed case (Figure 7.14). At early times, both excitonic and CT states are involved in the decay of the dark state. Initially, specially in the first 50 fs, the D state shows intense population oscillations coherent with the fluctuations in the total excitonic population. The S_1 occupation reaches $\sim 20\%$, and rapidly decays to $\sim 3\%$. The fast depopulation of S_1 coincides with the modest raise of CT population. Moreover, the steep formation of 1TT occurs from the very beginning, suggesting an intricate participation of D, S_1 and CT states in the SF mechanism. Notably, the $DS_0 \rightarrow ^1TT$ photophysical reaction is nearly completed with 0.5 ps, faster than in the direct case.

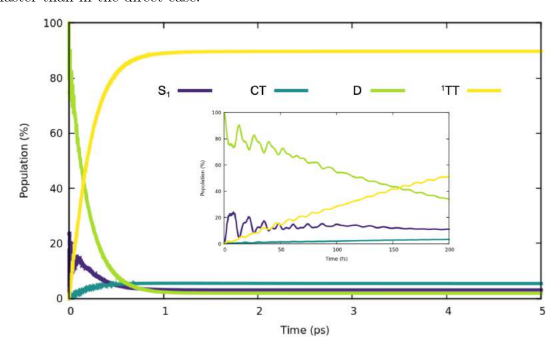


Figure 7.14: Time evolution of state population (in %) for two co-planar ethylene molecules (x = 1.4, y = 0.0, z = 3.5 Å) from DS₀ state, with $E(DS_0) - E(^1TT) = 0.25$ eV, and S_1S_0/S_0S_1 and CT states at 0.4 and 3.4 eV. D, S_1 and CT indicate the total population of DS_0/S_0D , S_1S_0/S_0S_1 and CA/AC, respectively. Early time evolution shown in the inset.

To confirm the simultaneous participation of S_1 and CT states, we simulate the SF dynamics without AC/CA (Figure 7.15a) and without S_1S_0/S_0S_1 (Figure 7.15b) configurations. In both cases the dynamics of the process changes. The SF reaction

becomes slower without S_1 states and specially in the absence of CT states. Without the presence of AC and CA configurations at relatively low energies, the initial DS_0 state strongly interacts with S_1S_0 , producing a mixed D ($\sim 80\%$) and S_1 ($\sim 20\%$) population in the first few fs. From there, the formation of the triplet-pair state is achieved through the simultaneous direct transition from D and S_1 states, with a faster reaction rate from the dark state. The changes observed without excitonic states are less extreme and mainly related with the evolution at early times. The formation of 1TT is nearly completed at 2 ps, which is twice the time required when S_1 states are energetically close (Figure 7.14), demonstrating the simultaneous mediation of CT and excitonic states.

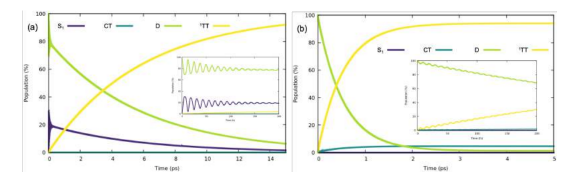


Figure 7.15: Time evolution of state population (in %) for two co-planar ethylene molecules (x = 1.4, y = 0.0, z = 3.5 Å) from DS₀ state, with $E(DS_0) - E(^1TT) = 0.25$ eV, and (a) S₁S₀/S₀S₁ states at 0.4 eV (no CT states) and (b) CT states at 3.4 eV (no S1 states). D, S₁ and CT indicate the total population of DS₀/S₀D, S₁S₀/S₀S₁ and CA/AC, respectively. Early time evolution shown in the inset.

D-mediator SF dynamics

In this section we explore the potential mediator role of the dark state in the SF dynamics with S_1S_0 as the initial state, and by considering two different coupling hot spots in the configurational space of two co-planar ethylene molecules.

The first case corresponds to the x-axis displaced conformation. In Figure 7.9 is illustrated the quite strong direct and D-mediated (second order) SF couplings, which are expected to dominate the excited state dynamics if CT states lie energetically high. This situation is represented with the simulation of exciton dynamics for the x = 0.8 Å conformation, suggesting a direct $S_1 \rightarrow {}^1\text{TT}$ mechanism (Figure 7.16). At this molecular arrangement the coupling between S_1S_0 and DS_0 (106 meV) is about an order of magnitude stronger than the $S_1S_0/{}^1\text{TT}$ interaction (17 meV). As a consequence, the S_1S_0 state initially interacts with the dark state, which reaches a population as high as 40%, but vanishes after less than 200 fs, while S_1 is totally recovered. During the first 200 fs the triplet-pair remains depopulated. At later times the S_1 and ${}^1\text{TT}$ populations simultaneously decay and grow following a first order kinetics. Thus, in this case, the D state is not actually mediating the forma-

tion of the triplet-pair state, but assisting exciton delocalization, that is from S_1S_0 to $S_1S_0+S_0S_1$, prior to the (direct) S_1 to ¹TT transition.

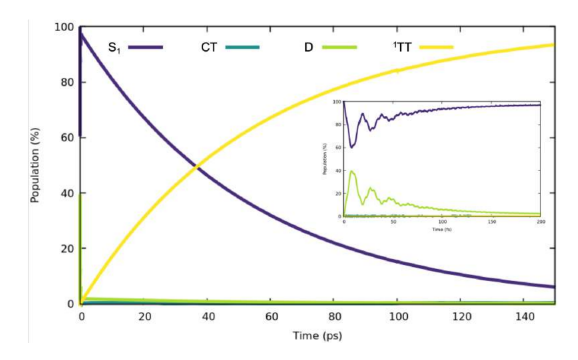


Figure 7.16: Time evolution of state population (in %) for two co-planar ethylene molecules (x = 0.8, y = 0.0, z = 3.5 Å) from S_1S_0 state, with $E(S_1S_0) - E(^1TT) = 0.55$ eV, and DS_0/S_0D and CT states at 0.4 and 20 eV. D, S_1 and CT indicate the total population of DS_0/S_0D , S_1S_0/S_0S_1 and CA/AC, respectively. Early time evolution shown in the inset.

Finally, I present the simulation of the SF dynamics for the ethylene dimer structural conformer with a long axis displacement of 1.4 Å, and D and CT states at 0.4 and 3.4 eV with respect to S_1 (Figure 7.17), for which SF couplings predict the possibility of direct, CT-mediated and CT,D-mediated contributions to the total coupling (Figure 7.11). The formation of the triplet-pair state is very fast, i.e., completed with 140 fs, although the final ¹TT state is strongly mixed with S_1 (20%) and CT (6%) configurations. D and CT states are largely involved in the early exciton dynamics, with strong oscillations between $S_1/D/{}^1$ TT. The initial decay of S_1 is concomitant to the raise of D, immediately followed by the increase in the 1 TT population. Although the occupation of the CT state remains always quite low due to its higher energy, it seems to also effect the time evolution of 1 TT.

Internal conversion to the triplet-pair becomes nearly an order of magnitude slower if the CT states are removed from the system (Figure 7.18a). The D state strongly interacts with S₁ and presents sizeable occupation at early times (0-60 fs), coinciding with a raise of ¹TT. The population of the dark state considerably diminishes approximately at 60-80 fs, just when the increase of ¹TT decelerates. This profile indicates a short-time D-mediated SF mechanism followed by the direct

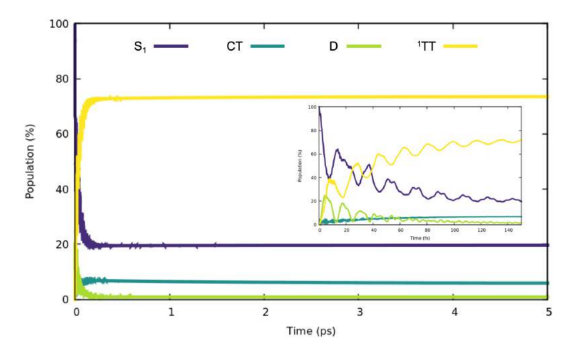


Figure 7.17: Time evolution of state population (in %) for two co-planar ethylene molecules (x = 1.4, y = 0.0, z = 3.5 Å) from S_1S_0 state, with $E(S_1S_0) - E(^1TT) = 0.25$ eV, and DS_0/S_0D and CT states at 0.4 and 3.4 eV. D, S_1 and CT indicate the total population of DS_0/S_0D , S_1S_0/S_0S_1 and CA/AC, respectively. Early time evolution shown in the inset.

 $S_1 \rightarrow {}^1TT$ transition. Alternatively, if the dark state is very high in energy (Figure 7.18b), the formation of 1TT is dominated by the CT-mediation, as it becomes evident when compared to the slower dynamics obtained when only the direct mechanism is taken into account (Figure 7.18c). Overall, despite that in this case the sole presence of the direct mechanism results in a rather fast 1TT generation, the different mediation mechanism through D and CT states are more efficient and are expected to dominate SF in slip-stacked conformations.

7.4 Appendix

7.4.1 Computational details

Electronic couplings between diabatic states have been computed using orthonormalized orbitals localized on each of the two ethylene molecules. Fragment orbitals have been obtained as linear combinations of canonical (ground state) Hartree-Fock orbitals of the dimer as described in Chapter 3.(35; 36) The 6-31G basis set has been employed in all electronic structure calculations. Notice that the D state in ethylene lies at quite high energy with respect to S_1 and twice the T_1 energy. On the other hand, here we use the ethylene dimer as a model to (only) compute electronic couplings, while we explore different relative energies between D, S_1 , 1TT and

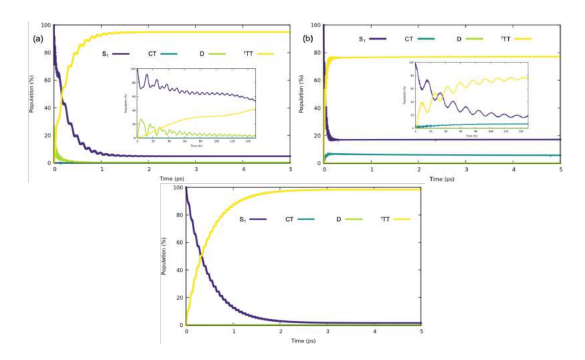


Figure 7.18: Time evolution of state population (in %) for two co-planar ethylene molecules (x = 1.4, y = 0.0, z = 3.5 Å) from S_1S_0 state, with $E(S_1S_0) - E(^1TT) = 0.25$ eV, and (a) DS_0/S_0D at 0.4 eV, and (b) CT states at 3.4 eV. D, S_1 and CT indicate the total population of DS_0/S_0D , S_1S_0/S_0S_1 and CA/AC, respectively. Early time evolution shown in the inset.

CT states, aiming to possible excited state distributions in realistic singlet fission sensitizers.(1; 2) These calculations have been performed with a development version of the Q-Chem program.(37)

The TCL QME equation has been solved by means of the fourth-order Runge-Kutta approach at T=300 K. Excited to ground state radiative decays have not been included in the exciton dynamic simulations. Singlet fission exciton dynamics calculations have been done with an in-house program developed by the group.

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Chapter 8

Spiroconjugated dimers

Spiro compounds are defined by the IUPAC as the compounds having one atom (usually a quaternary carbon) as the only common member of two rings. (?????) If both halves are constituted by π -electron systems, they are named as spiroconjugated compounds. (1) Then, spiro-conjugation is the term used to describe the indirect interaction between two systems of π -orbitals in perpendicular planes and linked together through a non-conjugated atom, *i.e.*, an sp^3 carbon. (2).

Spiro-conjugated systems are very attractive due to their molecular properties triggered by relative orientation between both halves. First, the orthogonality and rigidity provided by the spiro-linkage avoids sterical hindrance between π -systems, and increase the stability and solubility with respect to the non-spiro precursors. Spectroscopically, the special molecular geometry minimizes electronic interactions, and the relative orientation of the monomeric transition dipole moments exclude the possibility of excimer formation and energy splitting of excited states as a consequence of molecular aggregation. These properties make them materials with potential applications in optoelectronic devices. They have been extensively investigated and used as field-effect transistors, components of solar cells, photodiodes and phototransistors, light-emitting devices, etc.(3; 4) Among all these processes SF has not been considered until very recently. By means of transient absorption spectroscopy, Kumarasamy et al. obtained fast SF rates and slow triplet pair recombination in a dimer of spiro-pentacene.(5)

In this chapter, spiroconjugated systems are studied as promising materials fulfilling the requirements to carry out two photophysical process to generate triplet excitons. In the first section, an exhaustive theoretical study of the suitability of proposed spiroconjugated dimers to undergo the SF process is presented. The potential to be good SF sensitizers of a family of pro-spiro chromophores is investigated. Then, the modelling of the SF intricacies in spiroconjugated dimers in terms of the nature of electronic states, interstate couplings, and SF mechanisms, as well, the effect of vibrational distortions were studied to obtain a rather complete picture of the process.

An alternative photophysical process to generate exciton triplets is the intersystem crossing (ISC). In collaboration with the group of Prof. Jinquan Chen from the East China Normal University, we explored the assistance of charge resonance to an efficient singlet to triplet conversion featured by a different family of spiroconjugated systems. In the last part of this chapter, depth characterization of the radical character in spiro compounds is provided.

8.1 Spiro-conjugated dimers in singlet fission

8.1.1 Chromophores pro-spiro

In the design of spiro-conjugated compounds suitable for SF, it is necessary to consider monomers with the proper energetics (section 7.1.1). The full characterization of SF chromophores expose their electronic features in order to determine if they are or not able to fulfill the SF requirements, i.e., $E(S_1) \ge 2E(T_1)$, $E(T_2) > 2E(T_1)$, etc. In this case, we consider two conjugated scaffolds 2,7-disubstituted-fluorene **Fn** and 2,7-disubstituted-5,10-dihydro-dibenzopentalene **DBPn**, containing one and two sp^3 (pro-spiro) carbon atoms, respectively, combined with two substituents: diemetilene (1) and cyanometilene group (2), see Figure 8.1.

Figure 8.1: Quinoidal (left) and diradical (right) electromers of the studied chromophores: **F1**: 2,7-dimethylene-fluorene, **F2**: 2,7-bis(dicyanomethylene)-fluorene, **DBP1**: 2,7-dimethylene-5,10-dihydro-dibenzopentalene and **DBP2**: 2,7-bis(dicyanomethylene)-5,10-dihydro-dibenzopentalene. Starred position indicate pro-spiro atoms.

Both structures present two planar resonance forms, namely quinoidal (closed-shell) and aromatic (diradical), exhibiting tendency to have diradical character. In the ground state both, **Fn** and **DBPn**, are planar with C_{2v} and C_{2h} molecular

symmetry, respectively.

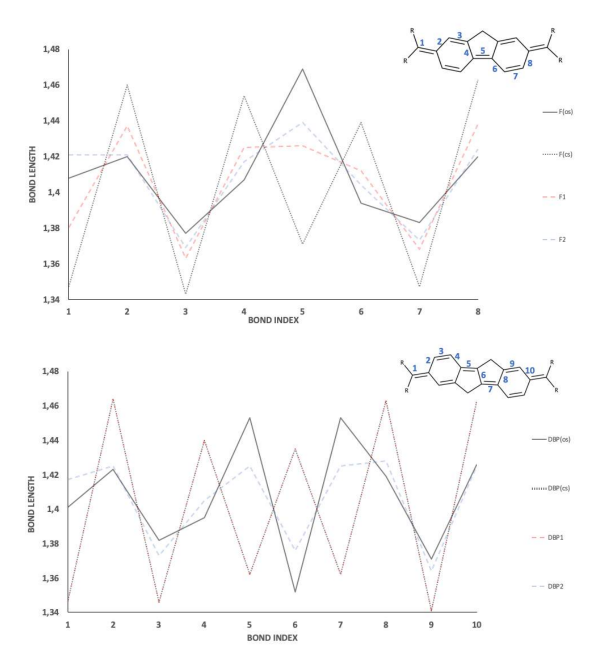


Figure 8.2: Bond length alternation of ground state optimized geometries of \mathbf{Fn} (a) and \mathbf{DBPn} (b) compared to the purely closed-shell (CS) and diradical (OS) structures. CS corresponds to the restricted singlet while OS to the unrestricted triplet for $\mathbf{R} = \mathbf{H}$.

The effect of the electron withdrawing power of the dicyano groups promotes the recovery of local aromaticity of the six-member rings in the backbone, leading to the apparition of diradicaloid character. The bond length alternation (Figure 8.2) shows that **F1** and **F2** have a similar bond length distribution that resembles to aromatic structure represented by the open-shell structure (diradical structure)

	F 1	F2	DBP1	DBP2
ΔE_{ST}^{ad}	10.0	5.5	19.3	5.4
y_0	0.39	0.44	0.17	0.46
N_u	0.92	0.99	0.51	1.05

Table 8.1: Adiabatic singlet-triplet energy gaps ($\Delta^{ad}E_{ST}$) computed at CAM-B3LYP level (in kcal/mol) and ground state diradical index (y_0) and number of unpaired electrons (N_u) obtained with RAS-SF of **Fn** and **DBPn** molecules.

in Figure 8.2, pointing that **F1** holds some diradical character. Differently, the minimum of **DBP1** is clearly quinoidal-like and corresponds to the closed-shell geometry since its radical form is destabilized by the ring strain around the vinylene group shared by two pentagons, as discussed in the study of carbon-bridged phenylenevinylene oligomers. (6). Computed singlet-triplet energy gaps brings an idea of the diradicaloid character of this monomers. Further, as signalised before, the relative energy between first excited singlet and triplet states determines the suitability of a compound to experience SF. Table 8.1 illustrates the intermediate diradical character of **F2** and **DBP2** ($y_0 = 0.44$ and $y_0 = 0.46$, respectively), which agree with the small ΔE_{ST}^{ad} . The spatial representation of the FOD density in the ground state (Figure 8.3 b) captures the diradical stabilization effect by the CN groups through delocalization of unpaired electrons. Radical index of $\mathbf{F1}$ is close to that of its analogous cyano-derivative as indicated by geometrical parameters, while the singlet-triplet energy gap is around twice the computed one for F2. On the other hand, **DBP1** shows rather mild diradical character as indicated by the larger ΔE_{ST}^{ad} and its diradical index is quite small $(y_0 = 0.17)$, in conjunction with a quinoidal-like structure.

HOMO and LUMO of the monomers are π -orbitals delocalized over the entire backbone but the sp^3 carbon atom (Figure 8.3 a). Pro-spiro atom plays an structural role helping to maintain the rigidity and planarity of the π -systems in both halves.

Computed low-lying excited states are summarized in Table 8.2. The main contributions to the electronic structure of the T_1 and the two lowest excited singlets involve the HOMO and LUMO frontier orbitals (Figure 8.3). While S_1 is mainly the HOMO² \rightarrow LUMO² transition, the most important configuration in S_2 corresponds to the HOMO-to-LUMO excitation. HOMO-1 \rightarrow LUMO and/or HOMO \rightarrow LUMO+1 configurations contributes to both excited singlets. In the T_2 state electronic excitation from HOMO-1 and HOMO to LUMO and LUMO+1 are involved. Vertical energies to the lowest triplet preserve the trend obtained for the adiabatic energies at the DFT level and dictated by the diradicaloid character of the molecules. In chromophores with the dicyano group, T_1 is considerably lower than in **F1** and **DBP1** (Table 8.1). Calculated excitations predict the presence of a dipole forbidden

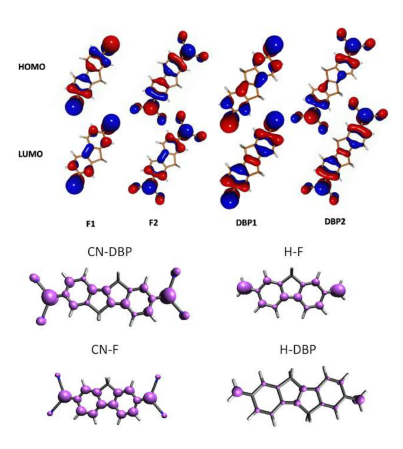


Figure 8.3: Frontier molecular orbitals (top) and FOD of studied monomers.

transition ($f \sim 0$) to a singlet state located just below the optical excitation, which holds an important double excitation character. This state ordering is in agreement with the 2^1A_g state lying below 1^1B_u in the closely related diphenylpolyenes. The optical state is S_2 , 1B_2 and 1B_u for **Fn** and **DBPn**, respectively, which corresponds mainly to the HOMO-to-LUMO single electron promotion and lies about 3 eV above the ground state for the dimethylene substitute monomers, while this dipole allowed transition is notably redshifted upon CN substitution, in agreement with the excitation energies predicted by TDDFT(7) (2.03 eV) and with the reported absorption maximum in solution (1.98 eV).(8)

molecule	state	ΔE	f	contributions (%)
F 1	${}^{3}B_{2}$	0.78	-	$H \to L (91)$
	$2 \times {}^3B_u$	1.57		
	$^{1}A_{1}$	2.75	0.008	$\mathrm{H}^2 \to \mathrm{L}^2 \ (46)$
				$\text{H-1} \to \text{L} (20)$
	${}^{1}B_{2}$	3.11	2.096	$H \to L (48)$
	9			$H-1 \rightarrow L/H \rightarrow L (25)$
	3A_1	3.44	-	` /
-				$H-1 \to L+1 \ (16)$
$\mathbf{F2}$	${}^{3}B_{2}$	0.41	-	$H \to L (94)$
	$2 \times {}^3B_u$	0.82		
	${}^{1}A_{1}$	2.17	0.005	$\mathrm{H}^2 \to \mathrm{L}^2 \ (41)$
				$H-1 \rightarrow L (29)$
	$^{1}B_{2}$	2.51	2.124	$H \to L (50)$
	3 4	0.00		$H-1 \rightarrow L/H \rightarrow L (28)$
	${}^{\circ}A_1$	2.82	-	()
				$H-1 \to L+1 \ (12)$
DBP1			-	$H \to L (87)$
	$2 \times {}^{3}B_{u}$			
	$^{1}A_{g}$	2.97	0.000	$H^2 \rightarrow L^2 (42)$
	1 D	2.05	0.706	$H \rightarrow L+1 (15)$
	$^{1}B_{u}$	5.05	2.720	$H \rightarrow L (52)$
	3 <u>/</u>	2 0/1	-	$H-1 \rightarrow L/H \rightarrow L (19)$ $H \rightarrow L+1 (35)$
	1 1 g	2.01		$H-1 \rightarrow L (34)$
DBP2	$^{3}B_{u}$	0.35		. ,
DDP2	$2 \times {}^3B_u$		-	$H \to L (93)$
			0.000	$\mathrm{H}^2 \to \mathrm{L}^2 \ (38)$
	2 1 9	1.00	0.000	$H-1 \rightarrow L (28)$
	$^{1}B_{n}$	2.28	2.393	$H \rightarrow L (42)$
	u			$H-1 \rightarrow L/H \rightarrow L (35)$
	3A_g	2.37	-	$H-1 \rightarrow L(60)$
				$\text{H1} \rightarrow \text{L+-1} (18)$

Table 8.2: Vertical excitation energies (ΔE in eV), oscillator strengths (f), and electronic transition contributions (in %) to the low-lying excited singlet and triplet states of **Fn** and **DBPn** computed at the RAS(4,4)-srVWN level. For the sake of comparison we include twice the energy of the lowest triplet. H = HOMO, L = LUMO.

Data in Table 8.2 and diradical character (Table 8.1) gives all the important quantities to establish relationships between excited states and preliminary conclusion about the feasibility of SF in these compounds. Since $2E(T_1) < E(S_1)$ SF for the four pro-spiro chromophores would be an exothermic process. Additionally, the second triplet state is energetically higher than twice the energy of T_1 , *i.e.*, $2E(T_1) < E(T_2)$, which is beneficial for SF since it hinders the efficient recombination of the fission triplets to the triplet manifold as a competing decay channel. Nevertheless, there is a large difference in **F1**, **F2** and **DBP2** between the $2T_1$ energy and the excitation to the lowest singlets (S_1 and S_2), which predicts a kinetically unfavorable SF photophysical reaction ($E(S_1)$ - $2E(T_1) > 1$ eV). On the other hand, transition energy to the lowest excited singlets in **DBP1** are rather close to twice the S_0 to T_1 gap (0.1-0.2 eV above $2E(T_1)$), satisfying thermodynamic and kinetic conditions for an efficient photophysical reaction.

These results are computed at the Franck-Condon (FC) region, thus the relaxation on the excited state potential energy surface (PES) has not been taken into account. The relative energy between both states S_1 and T_1 should be evaluated at the minima of the potential energy surface since the fission rate is slower than the vibration relaxation rate. Before the generation of the multiexcitonic state (1 TT state), the ${}^{1}B_{u}$ state is expected to follow a vibrational relaxation to the minimum in its PES in an ultrafast process that can lead to unfavorable energy gaps between S_2 and T_1 . Geometry optimization of **DBP1** on the excited ${}^{1}B_{u}$ PES systematically lowers the excitation energies with respect to the gaps at the ground state structure as shown in Table 8.3. The ordering of states is the same as in the FC geometry. With the relaxation of ${}^{1}B_{u}$, the ${}^{3}B_{u}$ state is also stabilized while the energy of ${}^{3}A_{g}$ is not significantly affected. Importantly, an increase on the SF exothermicity is produced upon the relaxation of the state, which could be beneficial in order to inhibit the reverse SF reaction once the triplets are formed.

state	S_0 geom.	$^{1}B_{u}$ geom.
3B_u	1.41	1.02
$^{1}A_{g}$	2.97	2.55
$^{1}B_{u}$	3.05	2.87
3A_g	2.94	2.83
$2 \times {}^3B_u$	2.85	2.03

Table 8.3: RAS(4,4)-srVWN vertical transition energies (in eV) to low-lying excited states of **DBP1** at the CAM-B3LYP ground and ${}^{1}B_{u}$ optimized geometries. For the sake of comparison we include twice the energy of the lowest triplet.

All these results point towards **DBP1** as the most promising candidate for effi-

cient SF. Then, the possibility of SF in the *spiro-DBP1* will be investigated in the following.

8.1.2 Spiro conjugated DBP1 dimer

The spiro system is obtained by fusion of two **DBP1** monomers through the prospiro carbon atom. The ground state characterization shows that the properties $(N_u, y_0, \text{ etc.})$ are obtained as twice the values in the monomer. A an example, Figure 8.4 shows the comparison between **DBP1** and its spiro derivative. Both monomers in the dimer are equivalent and the bond length alternation remains the same as in the **DBP1**. In general, the formation of the spiro compound implies a rather small separation between chromophores and the sp^3 -hibridization of the spiro atom forces the orthogonality between the two π -systems. Additionally, the bond lengths related to the spiro-linkage, as well as the angle C-C_{spiro}-C are marginally distorted with respect to ones in the pro-spiro chromophore (Figure 8.4b).

The spiro-DBP1 dimer contains a 2-fold rotation axis (C_2 symmetry point group) that goes through the spiro atom in the direction resulting from the sum of the normal vectors of the two monomeric planes. Although in the dimer the two π -electron systems are perpendicular to each other, they can effectively interact contingent upon their local symmetry. (2?) More concretely, only those orbitals from the monomer that are simultaneously antisymmetric to the two perpendicular reflexion planes in the dimer have the right symmetry to interact. For the case of π -orbitals it corresponds to those orbitals invariant under a C_2 rotation around the axis in the molecular plane that goes through the spiro atom and is parallel to the short monomeric axis (pseudo C_{2v} in Figure 8.5). Here, this condition has to be considered only approximatively, since the **DBP1** is not invariant under such C_2 rotation. In this sense, the LUMO of **DBP1** is locally (around the spiro atom) C_2 symmetric, while the HOMO is antisymmetric. As a result, the two HOMOs of spiro-DBP1 are degenerate (non-interacting monomeric HOMOs) and the LUMO and LUMO+1 present a small energy splitting ($\Delta_L = 54$ meV). Considering the poor intermonomer orbital overlap, such orbital spliting can be approximated as twice the two-electron exchange integral of the LUMO on the two chromophores (K_{ll})

$$\Delta_L \approx 2K_{ll}.\tag{8.1}$$

From such distribution of the frontier molecular orbitals a moderate redshift of the transition energies to the lowest excited states with respect to the monomer might be predicted. It is also worth noticing the lack of electron density on the spiro atom in all the frontier molecular orbitals, which suggests that the bridging sp^3 carbon is not directly involved in the electronic structure of the low-lying states,

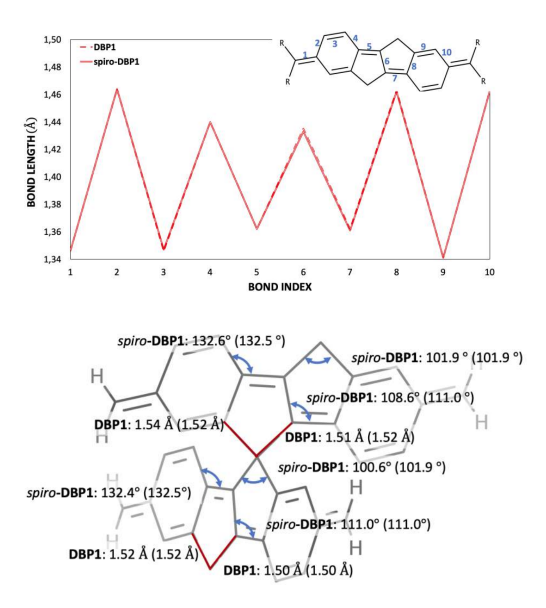


Figure 8.4: Bond length variation of **DBP1** and *spiro-***DBP1** (a), distortion leaded by the spiro-link in *spiro-***DBP1** with the values of **DBP1** shown in parenthesis for comparison (b).

and that its impact in the relative energies and couplings is merely structural.

8.1.2.1 Electronic states of spiro-DBP1

Energy distribution of the lowest-lying electronic excited states, singlets, of the spiro-**DBP1** are shown in Table 8.4. Pairs of quasi-degenerated excited states can be observed, which is in good agreement with the idea of two chromophores weakly interacting. Further, as predicted by the Δ_L , there is a very small redshift of the transition energies with respect to the monomeric excited states. The nature of all these excited states (adiabatic states) can be characterized in terms of coupling of

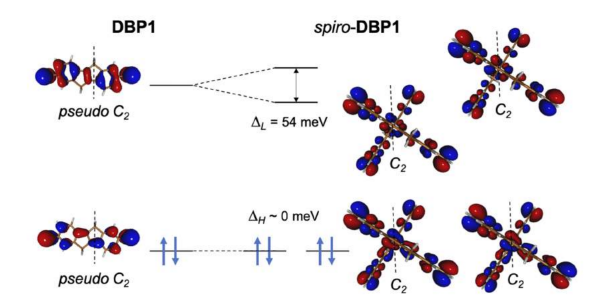


Figure 8.5: Frontier molecular orbital energy diagram of **DBP1** (left) and *spiro-***DBP1** (right). The vertical dashed line on **DBP1** indicates the $pseudo-C_{2v}$ rotation discussed in the text. The vertical dashed line on spiro-DBP1 corresponds to the C_{2v} symmetry axis.

monomeric states (diabats). Then, the wave function of every excited state can be expressed in terms of local and collective excitons. In Table 8.4 the composition of the adiabatic states (herein labeled in boldface) reflects the main electronic character. Note that the SS contribution includes all possible combinations of S_iS_j with i, j = 0, 1, 2, ... configurations.

State	Symm	ΔE	f	ω^{SS}	ω^{CT}	ω^{TT}	N_u
$^{1}\mathrm{TT}$	A	2.83	0.000	0.09	2.94	96.97	4.29
\mathbf{S}_1	B	2.94	0.013	98.35	1.64	0.01	3.63
$\mathbf{S}_{1}^{'}$	A	2.95	0.002	89.60	9.75	0.65	3.63
\mathbf{CR}_1	B	3.02	0.089	3.50	96.07	0.43	2.79
\mathbf{CR}_2	A	3.04	0.021	10.65	86.80	2.55	2.74
\mathbf{S}_2	B	3.05	2.604	96.70	3.29	0.01	2.83
\mathbf{S}_{2}^{\prime}	A	3.09	2.643	98.54	1.44	0.02	2.89

Table 8.4: Excitation energies (ΔE in eV), oscillator strengths (f), and composition contributions with singlet-singlet (ω^{SS}), charge transfer (ω^{CT}), and multiexcitonic (ω^{TT}) character in % of spiro-**DBP1** computed at the RAS-srVWN level.

The lowest-lying excited state of the spiro-DBP1 has an excitation energy (2.93)

State	Excitation energy (eV)	Oscillator strength	N_u
T_2	1.41	0.003	2.25
T_3	2.93	0.000	3.42
T_4	3.39	0.000	2.49
T_5	3.71	1.858	2.35

Table 8.5: DBP1 $T_1 \to T_n$ transitions computed at RAS-SF/6-31G**.

eV), very close to twice the lowest triplet in the monomer (2.92 eV). This transition has a multiexcitonic character, that can be assigned to a pair of triplets localized in each monomer and coupled as an overall singlet, as can be reflected in the large number of unpaired electrons ($N_u = 4.29$).

The ${}^{1}\mathbf{TT}$ state is composed by 96.97 % of triplet-triplet character, but also charge transfer contribute to it, suggesting that there is a weak electronic interaction between both triplets. Indeed, following the approximation to the triplet-triplet energy binding, E_b , as the energy difference between the triplet-pair quintet and singlet states:

$$E_b = E(^5\text{TT}) - E(^1\text{TT}) = -42 \text{ meV},$$
 (8.2)

results in a very small triplet-triplet interaction. At the RAS(8,8)-2SF/6-311+G(d,p) level, ${}^{1}\mathbf{TT}$ is higher in energy than ${}^{5}\mathbf{TT}$ (E_{b} is negative). Triplet-pair in this situation is named as unbounded biexciton and means that triplet separation will be exothermic (9).

Since the electronically coupled triplet-pair state typically exhibits characteristic transitions in the excited state spectra, (10; 11) in order to further investigate the nature of the SF precursor state we characterize electronic transitions from ${}^{1}\mathbf{T}\mathbf{T}$ compared to triplet-triplet promotions from T_{1} state in **DBP1** (Table 8.5). Computed high-energy states and their transition dipole moments from the ${}^{1}\mathbf{T}\mathbf{T}$ state, indicate that excited state optically allowed transitions correspond to excitations to ${}^{1}\mathbf{T}_{1}\mathbf{T}_{i}$ associated to $\mathrm{T}_{1} \to \mathrm{T}_{i}$ transitions in the **DBP1** monomer (Table 8.6), and hence predict the lack of additional transient absorptions than those corresponding to triplet-triplet transitions. All these results point towards the formation of a spin coherent triplet-pair state with very small electronic coupling between the localized triplets. Such state has been labeled by Pensack and coworkers as ${}^{1}(\mathbf{T} \cdots \mathbf{T}).(12)$

About 0.1 eV above ${}^{1}\mathbf{TT}$ lie a pair of near-degenerated singlet dark states with A and B symmetries resulting from the coupling of two local S_1 (${}^{1}A_g$) excitations on each moiety (\mathbf{S}_1 and $\mathbf{S}_{1'}$). Both states exhibit LE/CT mixings, specially the C_2 -symmetric transition ($\mathbf{S}_{1'}$) for which charge separation configurations represent almost 10% of the transition. Similarly, the two linear combinations of the monomeric

State	Excitation energy (eV)	Transition dipole moment	N_u
$^{-1}$ $\mathrm{T_{1}T_{2}}$	1.42	0.003	4.40
1 $\mathbf{T}_2\mathbf{T}_1$	1.44	0.001	4.25
$^1~\mathbf{T}_2\mathbf{T}_2$	2.83	0.000	4.47
1 $\mathbf{T}_1\mathbf{T}_3$	2.94	0.000	5.43
1 $\mathbf{T}_3\mathbf{T}_1$	2.95	0.000	5.41
1 $\mathbf{T}_1\mathbf{T}_4$	3.40	0.000	5.04
$^{1}\mathbf{X}$	3.65	0.052	3.31
$^{1}\mathbf{T}_{1}\mathbf{T}_{5}$	3.70	1.852	4.53
$^{1}\mathbf{T}_{5}\mathbf{T}_{1}$	3.73	1.231	4.51

Table 8.6: spiro-**DBP1** transitions from ${}^{1}\mathbf{T}_{1}\mathbf{T}_{1}$ to higher excited states computed at RAS-SF/6-31G** using an active space of 8 electrons into 8- π orbitals. ${}^{1}\mathbf{X}$ is a computed state that does not correspond with the TT character.

bright state (S_2) on each DBP1 moiety are very close in energy. Excitations to S_2 and $S_{2'}$ exhibit transition dipole moments perpendicular (B symmetry) and parallel (A symmetry) to the molecular rotational axis, in agreement with the dipole-dipole interaction model of Kasha.(13) The small energy gap between these states and ${}^{1}\mathbf{TT}$ suggests that an efficient SF reaction might occur from S_1/S'_1 or S_2/S'_2 .

At the same transition energies as the S_i excitonic states (~ 3 eV), two novel excitations appear that cannot be directly associated with any of the electronic transitions of the monomer. Decomposition of their electronic wave function characterizes them as symmetric combinations of CT configurations between the two DBP1 fragments, that is charge resonant states (14) (CR₁ and CR₂) belonging to the A and B representations of the molecular symmetry. Charge transfer or resonance states have been extensively studied as mediators in the SF process. In the case of not too high CT energies (~ 1 eV above $S_1/^1TT$),(15) the charge separation configurations act as virtual states assisting the formation of the triplet-pair (superexchange), leading to the commonly referred CT-mediated SF, which is the most commonly invoked SF mechanism. (16; 17; 18; 19; 20) In spiro-DBP1 these states appear rather low in energy compared to CT energies computed for other SF materials, such as in covalent dimers of pentacene. (21) The presence of such low-energy CR states can be rationalized by the 1/R dependence of the energy of separated charges and the rather short distance between DBP1 fragments in the dimer, which is approximately 2.5 Å in spiro-DBP1, 3.7-3.8 Å in crystalline tetracene and pentacene and 3.1 Å in efficient SF tetracene covalent dimers(22; 23) (as measured by the shortest inter-monomer C-C distance). More importantly, in this case \mathbf{CR}_1 and \mathbf{CR}_2 are virtually isoenergetic to the lowest singlets, which could potentially facilitate the efficient formation of ¹**TT**. On the other hand, the existence of very low CT states could eventually provide a nonradiative decay channel back to the ground state, possibly through the formation of trap states,(24; 25; 26; 23) and hence being detrimental to SF.

8.1.2.2 Singlet fission mechanism

Results presented until this point suggest than, indeed, the spiro-dimers are promising candidates to carry out SF. In one hand, the selection of the chromophore with the optimum excited state distribution is necessary for an efficient process. On the other hand, the weak interaction between monomers in spiro-compounds does not importantly modify the electronic structure of the monomer, and additionally, the short distance between them makes the lower charge transfer/resonance states available to play a (significant) role in the SF mechanism.

The first step in the SF reaction is completed with the formation of the multiexcitonic state (Equation 7.1). Here, the presence of the large number of excited states in a small range of energy allows all them to participate in the formation of ${}^{1}\mathbf{TT}$. The relative rates of formation from different electronic states $(r_{1,i})$ are evaluated by means of the kinetic model based on the transition state theory modification of the Fermi golden rule developed by Feng et al. (27)

$$r_{1,i}[\mathbf{S}_i/\mathbf{C}\mathbf{R}_i \to {}^{1}\mathbf{T}\mathbf{T}] \approx |V|^2 e^{-\alpha\Delta G_i/kT},$$
 (8.3)

where i is the first or second single excitonic state or charge resonance state, V is the nonadiabatic coupling between initial and final states, α is the free energy approach coefficient accounting for the magnitude of the activation energy (as suggested previously,(?) $\alpha = 0.5$ is used), ΔG_i is the free energy difference between initial and final states, k is the Boltzmann constant, and T is the temperature. Previously, this equation has been simplified by approximating the nonadiabatic coupling V (Equation 8.3) to the norm of the symmetrized one-particle transition density matrix (γ)(?) (Equation 8.4) and the ΔG_i by the electronic energy difference between states (ΔE_i).

$$|V| \propto \frac{||\gamma_i||}{\Delta E_i}.\tag{8.4}$$

The transition between excitonic and CR states will be much faster than the formation of 1 TT. Therefore, we assume that after absorption the excited states population follows a Boltzmann distribution among \mathbf{S}_{i} , \mathbf{CR}_{i} , and \mathbf{S}_{i} . Then, the overall rate of the first step in SF is approximated as the sum of the contributions from the set of rapidly populated states

$$r_1 \approx \sum_i p_i r_{1,i},\tag{8.5}$$

where p_i is the Boltzmann's factor of state i. In order to evaluate the importance of each transition to the final product (${}^{1}\mathbf{TT}$) the factor f_i is defined as:

$$f_i \equiv p_i r_{1,i}. \tag{8.6}$$

Table 8.7 resumes the values for these parameters. As pointed out by the energy distribution of excited states, ΔE_i goes from 0.1 to \sim 0.3 eV. At a given temperature (300 K here), the population of the lowest excited states is considerably larger than in higher states, as indicated by the Boltzmann's factors, suggesting the fast internal conversion to CR_i and later to S_1/S_1 states after photo-excitation to the bright states. The strongest coupling $(||\gamma_i||^2)$ is obtained between \mathbf{CR}_i and the triplettriplet pair. Actually, these states contribute with more than the 50% to r_1 despite they hold less than 4% of population. Nonadiabatic couplings of S_2 and S_2' with ¹TT are at least one order of magnitude smaller with respect to the CR/TT ones. This is in good agreement with previous studies reporting that the CR/TT coupling in SF materials is significantly stronger than S_i/TT interaction. (?) The two lowest excitonic states have more than 90% of the electronic population. Nevertheless, $||\gamma_{S_1,^1TT}||^2$ and f_i of S_1 are rather small. On the other hand, S_1 ' contributes to the formation of ¹TT with close to 40%. This can be rationalized as a result of the contribution of the CT configurations ($\sim 10\%$) to its wave function. Bright states play a marginal role in the SF mechanism, pointing that the direct transition from S_2 to TT is not favorable in spiro-**DBP1**.

state	Sym.	ΔE_i	p_i	$ \gamma_i ^2$	f_i
\mathbf{S}_1	В	0.11	0.590	0.004	0.072
$\mathbf{S}_{1'}$	A	0.12	0.363	0.034	0.383
\mathbf{CR}_1	В	0.19	0.023	0.296	0.335
\mathbf{CR}_2	A	0.21	0.014	0.263	0.205
\mathbf{S}_2	В	0.22	0.008	0.026	0.003
$\mathbf{S}_{2'}$	A	0.26	0.002	0.016	0.003

Table 8.7: Electronic energy gap ΔE_i (in eV), Boltzmann's factor p_i (T = 300 K), squared norm of γ_i and contribution to r_1 (f_i) for the transition from excitonic and \mathbf{CR}_i excitations to the triplet-pair state in spiro- $\mathbf{DBP1}$.

These results highlight the crucial role of CT contributions in spiroconjugated dimers. SF in $spiro-\mathbf{DBP1}$ is predicted to mainly occur through the population of low-lying excited states with strong CR character (\mathbf{CR}_i) and through the lowest excitonic states with important mixing with CT configuration.

Interestingly, \mathbf{S}_1 and $\mathbf{S}_{1'}$ are dark states in which the main transition is HOMO² \rightarrow LUMO², meaning they are mostly the doubly ("D") excited state described in

previous chapter. Energetically, the situation of the *spiro*-**DBP1** could be described by the couplings in Figure 7.7, but the relative orientation of the monomers in the dimer is different in each case, *i.e.*, (slip-)stacked in ethylene and orthogonal in *spiro*-**DBP1**.

The complete SF photophysical reaction requires the splitting of ${}^{1}\mathbf{TT}$ into two independent triplets. The relative rate of this reaction can be evaluated by means of the binding energy (E_b) as:

$$r_2[^1\mathbf{TT} \to \mathbf{T}_1 + \mathbf{T}_1] \approx e^{-\alpha E_b/kT}$$
 (8.7)

It is expected that the weak electronic interaction between the two triplets in ${}^{1}\mathbf{TT}$, given by the negative sign of E_{b} (-42 meV), results in $r_{1} << r_{2}$. But rates are computed taking account different approximations and comparison between them is not possible. Still it is safe to conclude that the formation of the ${}^{1}\mathbf{TT}$ will be the rate limiting step for the SF fission efficiency in spiro-**DBP1**, and probably in general for spiroconjugated compounds with triplet and singlet energies suitable for singlet fission.

8.1.3 Vibronic couplings

Even with the rigidity of the dimeric scaffold provided by the spiro-atom, some vibrational modes can distort the ground state and favour a deactivation path different from SF or modify the mechanism in which it occurs. Actually, it has been shown that vibrational distortions can change the interaction between two moieties in spiroconjugated systems. (28) So far, the Franck-Condon geometry has been explored, suggesting the suitability of the spiro-DPB1 to perform SF. This static picture does not consider the structural variation that can occur due to finite temperature effects or via the energy relaxation of the photoactivated system on the excited state PES. The first class are related to low-frequency modes and corresponds to geometrical distortions within the kT energy range that can modify the relative orientation and, as a consequence, interaction between the two moieties. On the other hand, the structural rearrangement of excited molecules from the ground state geometry to the excited state minima might involve much larger energy changes, triggering the movement of high-frequency vibrations. This is the typical case in conjugated systems, for which $\pi \to \pi^*$ electron promotions couple to CC stretching modes of the order of $1400-1600 \text{ cm}^{-1}$.

In the following, the effect of low- and high-frequency vibrations in the SF capabilities of *spiro-DBP1* will be explored.

8.1.3.1 Low-energy vibrational modes

In order to investigate the impact of the thermal activation of the excited vibrational levels of low-frequency molecular modes, we explore the distortion along the five lowest-energy vibrational modes of spiro-**DBP1**. We do not aim to exhaustively analyze all vibrations; rather, we consider the five molecular modes $[Q_i(\Gamma), i = 1, 5, \Gamma = A, B]$ with harmonic frequencies $< 50 \text{ cm}^{-1}$ (Figure 8.6), which mostly corresponds to symmetric (A) and antisymmetric (B) bending deformations between the two monomers modifying their relative disposition (dihedral angle).

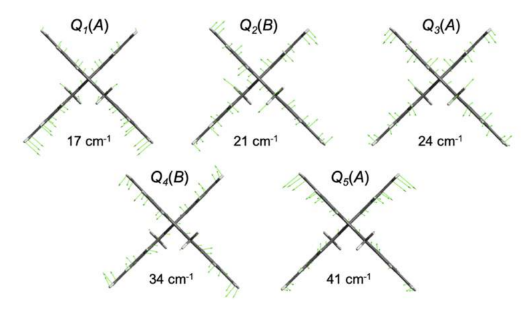


Figure 8.6: Lowest frequency modes of *spiro-DBP1* computed at the CAM-B3LYP level.

To infer the potential role of these modes, we evaluate excited state energies and interstate couplings ($||\gamma||$) at the distorted geometries of the dimer following each $Q_i(\Gamma)$ deformation, implying an energy change of the order of kT ($T=300\mathrm{K}$). Overall, the relative energies of low-lying states of spiro-**DBP1** do not change very much with respect to the ground state distribution. The low-lying states remain within a rather small energy range (0.1-0.3 eV) for the five distortions with the ${}^{1}\mathbf{TT}$ state systematically being the lowest excited singlet (Figure 8.7a). Additionally, the geometrical changes induce small modifications of the interstate couplings ($||\gamma||^{2}$ in Figure 8.7b). The only exception is the distortion along the $Q_3(A)$ mode, which exhibits reduced relative energies with respect to ${}^{1}\mathbf{TT}$ and sizable modifications of the computed couplings.

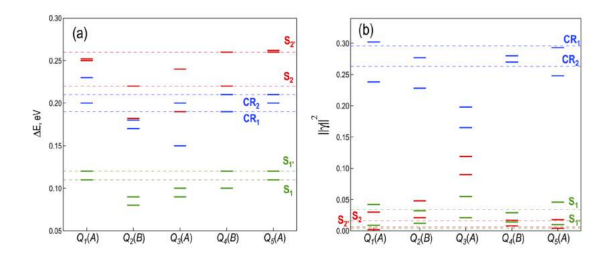


Figure 8.7: Excited state energy diagram (in eV) with respect to ${}^{1}\mathbf{TT}$ (a) and $||\gamma||^{2}$ values to ${}^{1}\mathbf{TT}$ (b) at the $Q_{i}(\Gamma)$ -distorted structures of spiro-**DBP1** computed at the RAS(8,8)-srVWN level. Slashed lines indicate the values at the Franck-Condon geometry.

These changes can be rationalized by means of the electronic composition of states in terms of diabatic contributions. The distorted geometry induced by the symmetric $Q_3(A)$ bending mode exhibits a quite different composition of the low-lying states with increased degrees of CT configurations into the triplet-pair and excitonic states (Figure 8.8), which explains the increased $||\gamma||^2$ values between ${}^1\mathbf{TT}$ and excitonic states.

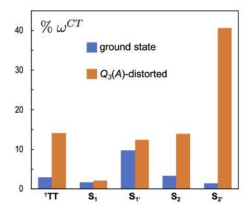


Figure 8.8: CT contributions ω^{CT} (in %) of low-lying states of *spiro-DBP1* at the ground state and $Q_3(A)$ -distorted structures computed at the RAS-*srVWN* level.

Changes in the interstate couplings might affect the SF mechanism. Indeed, the slight energy lowering of excitonic states with respect to ${}^{1}\mathbf{TT}$ and the increase of electronic couplings notably modify the weight of the different SF paths (Table 8.8), with the transition from $\mathbf{S}_{1'}$ becoming the main path toward the triplet-pair state

state	Sym.	ΔE_i	p_{i}	$ \gamma_i ^2$	f_i
\mathbf{S}_1	B	0.09	0.512	0.021	0.211
$\mathbf{S}_{1'}$	A	0.10	0.410	0.055	0.442
\mathbf{CR}_1	B	0.15	0.054	0.198	0.245
\mathbf{CR}_2	A	0.20	0.009	0.165	0.064
\mathbf{S}_2	B	0.24	0.002	0.090	0.028
$\mathbf{S}_{2'}$	A	0.19	0.013	0.119	0.011

Table 8.8: Electronic energy gaps with respect to ${}^{1}\mathbf{TT}$ (ΔE_{i} , in eV), Boltzmann's factors p_{i} (T = 300 K), squared norms of γ_{i} , contributions to r_{1} (f_{i}) for the transition from excitonic and \mathbf{CR}_{i} excitations to the triplet-pair state for the $Q_{3}(A)$ -distorted spiro-**DBP1** computed at the RAS(8,8)-srVWN level.

and with a notably larger overall rate of formation, i.e., $r_1(Q_3)/r_1(FC) = 1.54$ ($\sim 50\%$ increase).

The increase of the TT/CT mixing also modifies the triple-triplet interaction in ${}^{1}\mathbf{TT}$, stabilizing the spin-singlet coupling of the two triplets with respect to ${}^{5}\mathbf{TT}$. At the Q3(A)-distorted geometry, the two spin multiplicities become virtually degenerate ($\Delta E_b \sim 0$ meV), reducing the ground state exoergicity for the triplet-triplet decoupling.

Although, these results do not predict major modifications of the photophysics of *spiro-DBP1* with temperature, the change on the state population can help to improve the SF process and its efficiency.

8.1.3.2 High-energy vibrational modes

Now we consider the possible impact that high-energy structural distortions might have on the SF reaction in spiro-**DBP1**. Here, I focus my attention to molecular vibrational modes involving stretching of carbon-carbon bonds (Figure 8.9). At 1359 and 1361 cm⁻¹, we identify two modes corresponding to the symmetric and antisymmetric stretching of C-C single bonds in the quinoidal form of **DBP1** moieties $[Q_{s1}(\Gamma), \Gamma = A, B]$, while symmetric/antisymmetric C=C stretchings $[Q_{s2}(\Gamma), \Gamma = A, B]$ appear at higher frequencies (1585 cm⁻¹).

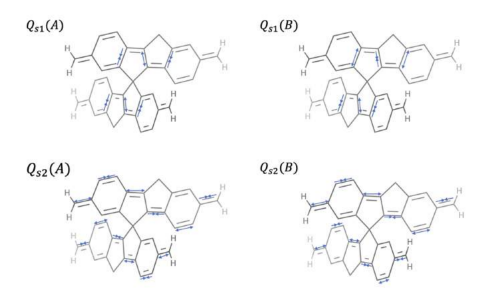


Figure 8.9: Symmetric (A) and antisymmetric (B) CC stretching vibrational modes of *spiro-***DBP1**.

These vibrational modes allow spiro-DBP1 to relax on the PES of the low-lying excited states. The symmetric $Q_{s1}(A)$ vibration toward the simultaneous shortening of the bonds shared by the two pentagons in each DBP1 monomer (negative displacements in Figure 8.10 a) stabilizes the singlet excited states, while in the opposite direction of $Q_{s1}(A)$ all low-lying singlets are strongly destabilized with respect to their Franck-Condon energies. These energy trends can be rationalized as the result of increasing and decreasing the π bonding and antibonding interactions in the frontier MOs, which can differently stabilize/destabilize the HOMO and LUMO of the monomers (Figure 8.3). Importantly, due to the simultaneous electron promotion to both monomeric LUMOs, energy stabilization upon $Q_{s1}(A)$ is more pronounced in ¹TT than for excitonic and CR states. This result is in line with the SF exothermicity increase observed in the excited state optimization of DBP1 (Table 8.3). Moreover, the relative stability of ${}^{1}\mathbf{TT}$ comes with a diminution of its CT character, which reduces from 3% at the Franck-Condon geometry to <0.5% for a 0.08 Å decrease in the central CC bond lengths. The $Q_{s1}(B)$ stretching breaks the C_2 symmetry of the dimer, generating inequivalent DBP1 units. Both directions of the vibrational distortion stabilize (destabilize) the HOMO (LUMO) of one of the monomers, while destabilize (stabilize) the HOMO (LUMO) of the other DBP1 fragment. As a result, ¹TT presents a rather flat and symmetric PES around the ground state geometry. On the other hand, excitonic states tend to localize on one of the two chromophores and CR contributions become unbalanced CT excitations with neat charge separation character (Figure 8.10 b). Moreover, $Q_{s1}(B)$ distortion diminishes the energy gap between the lowest excitonic state and ¹TT, which might facilitate the formation of the triplet-pair.

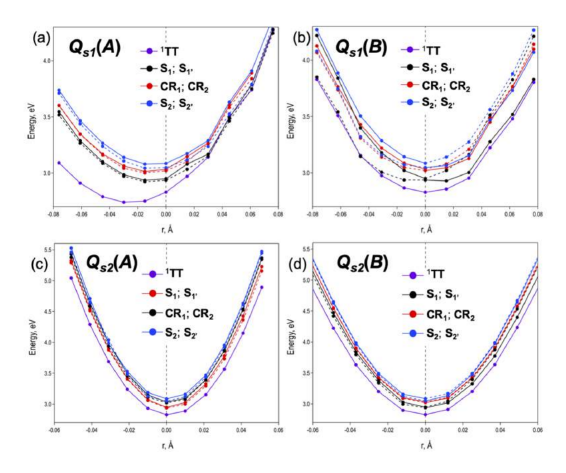


Figure 8.10: Excited state energy profiles (in eV) along the CC stretching bonds in Figure 8.9 computed at the RAS(8,8)-srVWN level.

Double bond stretchings in $Q_{s2}(\Gamma)$ systematically increase the energy of all lowlying singlets. Both symmetric, $Q_{s2}(A)$, and antisymmetric, $Q_{s2}(B)$, distortions largely preserve the relative state energies (Figure 8.10 (c) and (d), respectively); hence, we do not expect them to have a strong impact on the SF capabilities.

In summary, this investigation points out that spiroconjuged dimers are suitable materials to carry out SF. **DBP1** is a promising chromophore with the proper excited state distribution to make SF process efficient. The spiro linkage favors the presence of low-lying charge resonance states, which play a major role in the formation of the triplet-pair state. Overall, these results suggest that spiroconjugated dimers are potentially good candidates to efficiently generate independent triplet states through SF.

8.2 Spin orbit charge transfer - intersystem crossing in spiroconjugated dimers

Triplet excited states play an important role in excited states dynamics, since they can provide a deactivation path. Differently from excited singlet states, triplets

have long-lifetime and can reach longer diffusion lengths. (29; 30) Excited triplet states have application in organic optoelectronic devices (as OLEDs, photodynamic therapy, organic memory devices, sensors, among other electronic, magnetic and medical applications). (29; 31; 32?) In addition, the relative ΔE_{ST} is associated with the diradical character, which as has been exposed through these pages, is related to potential uses in organic photovoltaics. Electronic transition from S_0 to T_1 is not spin allowed since there is a net change in the state multiplicity, i.e., $\Delta S \neq 0$. Nevertheless, the population of the excited triplet states can be reached from the deactivation of higher singlet excited states through radiativeless process intersystem crossing (ISC).(33) ISC is the transition process between states of different multiplicities, form singlet to triplet, or triplet to singlet as is the case in reverse-ISC. In principle, this transition is forbidden by spin symmetry selection rule, but the process is facilitated by the spin-orbit coupling (SOC) effect which mixes the angular orbital moment L with the spin S of the singlet and triplet states represented by the wave functions, Ψ_S and Ψ_T , as $\langle \Psi_S | \hat{S} \cdot \hat{L} | \Psi_T \rangle$. To enhance the SOC it is important the energy gap between both states to be small. Also, big atomic numbers results in large orbital and spin magnetic moments, which favors the spin reorientation, i.e., from β to α in order to obtain the triplet state. The dependence of the SOC strength with the atomic number is named as "heavy atom effect" (HAE).(34) Based on the HAE effect, modifications to organic scaffolds have been made to include heavy atoms, organic (S, Te, Se, P) or transition metals and study the ISC in organic molecules. (35; 36)

Another mechanism to induce the ISC is through charge resonance process. To this end donor-aceptor moieties of a systems are ideally well separated, and the molecular overlap of both halves is rather poor.(33). The spin-flip is facilitate since the orbital angular momentum tends to change during the electron migration in the CR process. As the SOC is enhanced by the CT states this mechanism is called as spin-orbit charge transfer intersystem crossing (SOCT-ISC, Figure 8.11).

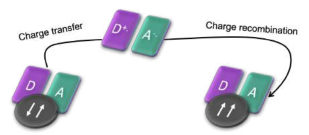


Figure 8.11: ISC process assisted by CT states in a donor-acceptor system. After its formation, the charges in the cationic/anionic monoradical pair are recombined promoting the spin-flip to form a triple state. Figure adapted from reference (33).

Few is known about this mechanism, but some studies have sited the basis to

enhance its efficiency. (37; 38) A large dihedral angle between π -systems, orthogonal molecular orbitals, and of course a low energy CT state are desirable. All these requirements can be fulfilled by the spiro-linkage as exposed in the previous section. Actually, the reverse-ISC process has been studied in spiroconjugated systems recently. (35; 39) Herein, I present the computational characterization of the SOCT-ISC mechanism in a family of spiroconjugated systems designed and experimentally studied by the group of Prof. Jinquan Chen from the East China Normal University. In the following I will discuss the theoretical aspects and results of this study. Complete experimental characterization of the pothophysics of these spiro-conjugated dimers can be found in reference (40)

Four 2,7-substituted benzophenone moieties are used as building blocks for the spirocompounds, ($\mathbf{m0}$ - 3 in Figure 8.12 a). All monomers are characterized having a singlet ground state with closed-shell structure. Computation of singlet and triplet excited states was performed with different methods. While T_1 in $\mathbf{m0}$ is a $\pi\pi^*$ transition, it has $n\pi^*$ character in $\mathbf{m1}$ and $\mathbf{m2}$, while in $\mathbf{m3}$ it has an intramonomer charge transfer (iCT) character ($\pi\pi^*$), and its $n\pi^*$ transition corresponds to T_2 . On the other hand, the first excited singlet is a dipole forbidden transition with $n\pi^*$ character, and the bright state (f > 0) with $\pi\pi^*$ character is higher in energy. At the CAM-B3LYP level, for all the monomers the excitation energies are $T_1 \sim 3.3$ eV, $S_1 \sim 4.0$, and bright state (S_2) of $\mathbf{m0}$, $\mathbf{m2}$ - 3 is around 4.5 eV, and 5.3 (S_4) for 1. The relative energies between S_n and T_1 are quite small, which discard $\mathbf{m0}$ - 3 as potential singlet fission chromophores.

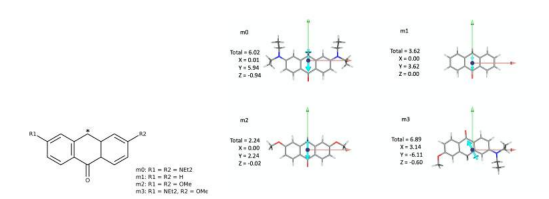


Figure 8.12: Structure of 2,7-substituted benzophenone (a) and permanent dipole moment (b) of **m0** - **3**.

The intensity of the permanent dipole moment for the ground state is modulated by the acceptor-donor character of the susbtituents, and is oriented in the short molecular axis pointing to the dione group with the exception of **m3**, where it is slightly diagonal due to the molecular asymmetry 8.12 b).

8.2.1 Spiro[bisanthracene]dione analogues

The three spiroconjugated dimers, **SF1** - **3**, respectively, were obtained by the combination of **m0** with **m1** - **3** (Figure 8.13). As expected from the spiro-linkage, monomers are oriented perpendicularly to each other.

Figure 8.13: Structure of 2,7-substituted benzophenone. Bottom part is **m0**, combined with **m1-3** monomers give rise to **SF1-SF3**.

Frontier molecular orbitals of **SF1** - **3** can be correlated to molecular orbitals of the monomers, particularly, HOMO and LUMO orbitals of the dimer are localized on a monomer, while the HOMO is over the **m0** moiety, the LUMO is over **m1-3** (Figure 8.14 a). The total dipole moment is given by the vector summation of the dipole moment of both monomers, pointing in the direction to the dione group of **m0** (Figure 8.14 b).

8.2.2 Photophysics of Spiro[bisanthracene]dione

Experimentally, the variation of the solvent polarity, from the non-polar cyclohexane to the polar acetonitrile going through the 2-methyltetrahydrofuran, shifts proportionally the main absorption band of the **SFn** to lower energies (Figure 2 of (40)). The absorption of spirocompounds were simulated in gas phase, (Figure 8.15), reproducing nicely the experimental observations. Calculations shown a large number of triplet excitations (5 at least) lying energetically below the excited singlets. Both, low-lying triplets and singlets, correspond to $\pi\pi^*$ and $n\pi^*$ transitions, but in **SF3** also iCT appears, directly related to the iCT state in **m3**. T₁ is characterized as a $\pi\pi^*$ transition (since iCT in **m3** is a $\pi\pi^*$ state) and have an excitation energy of \sim 3.2 eV. The bright singlet is computed around 4.2 eV having also a $\pi\pi^*$ character. By comparison with the excitation energy of the monomers; it is easy to notice that absorption bands of the dimers mostly match with the maximum absorption localized on each monomer. The most intense band corresponds to the optical transition

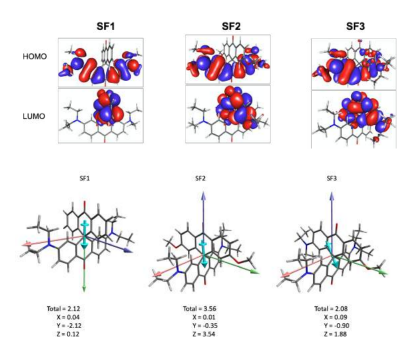


Figure 8.14: HOMO and LUMO orbitals (a) and permanent dipole moment in Debyes (b) of **SF1-3**.

of the monomer $\mathbf{m0}$ (4.48 eV), while the local excitations in $\mathbf{m1}$ - 3 are higher in energy and weaker on intensity. As in the *spiro*-**DBP1** homodimer, the properties of the monomer are conserved and the small redshift from $\mathbf{m0}$ to \mathbf{SFn} found can be rationalizated as the weak interaction between monomers in the spiro-compound. Optically active and dark states can be analysed in terms of the contribution of the monomeric excitations. Interestingly, all states computed in the energy range of the bright state (S*, *i.e.*, S₄ of $\mathbf{SF1}$ and S₃ of $\mathbf{SF2}$ and $\mathbf{SF3}$) have small contributions, and less than 1 eV above the S* there is a state with $\sim 90\%$ charge transfer character from $\mathbf{m0}$ to $\mathbf{m1}$ -3 (transitions from $\mathbf{m1}$ -3 to $\mathbf{m0}$ are energetically higher).

State	ΔE	f	Character	Transition	Composition (%)
SF1					
T_1	3.28	-	LE/CT	$\mathrm{m0}(\pi\pi^*)/\mathrm{m0} ightarrow \mathrm{m1}$	76.9/7.9
T_2	3.30	-	LE/CT	$\mathrm{m0}(\pi\pi^*)/\mathrm{m0} ightarrow \mathrm{m1}$	74.7/17.3
T_3	3.38	-	LE/CT	$\mathrm{m1}(\mathrm{n}\pi^*)/\mathrm{m0} o \mathrm{m1}$	89.1/5.6
T_4	3.57	-	LE/CT	$\mathrm{m0}(\mathrm{n}\pi^*)/\mathrm{m1} \to \mathrm{m0}$	76.6/10.7
T_5	3.66	-	LE/CT	$m0(\pi\pi^*)/m1 \rightarrow m0$	65.6/11.7
S_1	3.86	0.000	LE/CT	$\mathrm{m1}(\mathrm{n}\pi^*)/\mathrm{m0} o \mathrm{m1}$	85.0/6.4
				Con	tinued on next page

Table 8.9 – continued from previous page

State	ΔE	f	Character	Transition Transition	Composition (%)
$\overline{S_2}$	4.03	0.000	LE/CT	$m0(n\pi^*)/m1 \rightarrow m0$	79.0/10.5
S_3	4.14	0.001	CT	$\mathrm{m0} ightarrow \mathrm{m1}$	97.5
S_4	4.24	0.864	$_{ m LE}$	$m0 \ (\pi\pi^*)$	91.0
S_5	4.46	0.000	CT	$m0 \rightarrow m1$	97.9
SF2					
T_1	3.29	-	LE/CT	$\mathrm{m0}(\pi\pi^*)/\mathrm{m0} ightarrow \mathrm{m2}$	82.3/4.8
T_2	3.31	-	LE/CT	$\mathrm{m0}(\pi\pi^*)/\mathrm{m0} o \mathrm{m2}$	82.1/7.6
T_3	3.51	-	LE/CT	$\mathrm{m2}(\pi\pi^*)/\mathrm{m0} o \mathrm{m2}$	69.5/5.1
T_4	3.52	-	LE/CT	$\mathrm{m2}(\mathrm{n}\pi^*)/\mathrm{m0} o \mathrm{m2}$	87.6/7.4
T_5	3.56	-	LE/CT	$\mathrm{m2}(\pi\pi^*)/\mathrm{m0} o \mathrm{m0}$	70.2/5.9
S_1	3.99	0.000	LE/CT	$\mathrm{m2}(\mathrm{n}\pi^*)/\mathrm{m0} o \mathrm{m2}$	88.2/8.2
S_2	4.03	0.000	LE/CT	$\mathrm{m0}(\pi\pi^*)/\mathrm{m0} ightarrow \mathrm{m2}$	82.1/7.2
S_3	4.26	0.848	$_{ m LE}$	$\mathrm{m}0(\pi\pi^*)$	91.0
S_4	4.42	0.020	CT	$m0 \rightarrow m2$	94.5
S_5	4.70	0.316	LE/CT/LE	$\mathrm{m2}(\pi\pi^*)/\mathrm{m0} \rightarrow \mathrm{m2/m0}(\pi\pi^*)$	46.1/19.4/12.8
SF3					
T_1	3.21	-	iCT/CT	$\mathrm{m}3(\pi\pi^*)/\mathrm{m}0 ightarrow \mathrm{m}3$	68.9/5.0
T_2	3.28	-	LE/CT	$\mathrm{m0}(\pi\pi^*)/\mathrm{m0} ightarrow \mathrm{m3}$	74.9/4.5
T_3	3.31	-	LE/CT	$\mathrm{m0}(\pi\pi^*)/\mathrm{m0} ightarrow \mathrm{m3}$	75.1/14.2
T_4	3.56	-	LE/CT	$\mathrm{m3}(\mathrm{n}\pi^*)/\mathrm{m0}(\mathrm{n}) \to \mathrm{m3}(\pi^*)$	66.8/20.0
T_5	3.58	-	LE/CT	$\mathrm{m0}(\mathrm{n}\pi^*)/\mathrm{m3}(\mathrm{n}) \to \mathrm{m0}(\pi^*)$	70.4/15.0
S_1	4.02	0.000	LE/CT	$\mathrm{m3}(\mathrm{n}\pi^*)/\mathrm{m0}(\mathrm{n}) \to \mathrm{m3}(\pi^*)$	77.2/14.3
S_2	4.04	0.000	LE/CT	$\mathrm{m0}(\mathrm{n}\pi^*)/\mathrm{m3} \to \mathrm{m0}$	72.3/14.8
S_3	4.24	0.845	${ m LE}$	$\mathrm{m0}(\pi\pi^*)$	88.7
S_4	4.33	0.677	iCT/CT	$\mathrm{m3}(\pi\pi^*)/\mathrm{m0} \rightarrow \mathrm{m3}$	70.7/14.6
S_5	4.62	0.030	CT/iCT	$m0 \rightarrow m3/m3(iCT)$	76.4/16.1

Vertical excitation energies (in eV), oscillator strengths (f), excited state character, transitions and orbital composition to the low-lying excited states of **SF1-3** computed at the ground state geometry. Character of the transition has been indicated in % for the main orbital contributions.

Big changes are observed in the emission spectra derived from the change of the polarity of the solvent. A main band is identified as the radiative decaiment from the S* state around the same wavelength in all **SFn** (~ 400 nm). A new band appears at lower energy (> 500 nm) which, broadens with the polarity of the solvent. This new

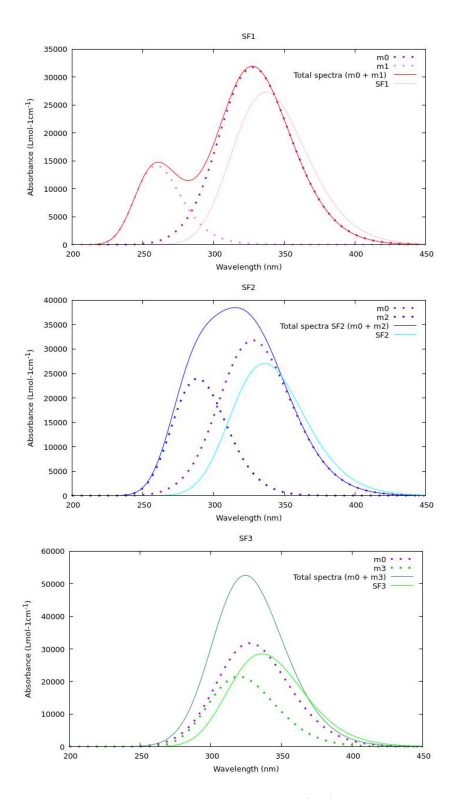


Figure 8.15: Simulated UV-VIS spectra of SF1-3 (a-c) with the contributions of excitations of each monomer. Total spectra represents the sum of the bands of each monomers while **SFn** is the spectra of the spiro dimer.

broad emission feature is associated to the stabilization of interchromophore charge transfer. At the minimum of the PES of each state the contribution of each monomer is manifested. The most intense band corresponds to the energy de-excitation of the

 $\pi\pi^*$ state of $\mathbf{m0}$, while in the CT minimum this state is strongly stabilized being the lowest excited state. Fluorescence from the CT state is then allowed by Kasha's rule. For instance, at the CT minimum the geometry of $\mathbf{SF1}$ is distorted and the CT character is maximized up to 96 % of $\mathbf{m0} \to \mathbf{m1}$. Notice that, due to the lack of molecular symmetry, CT states are not degenerated ($\mathbf{m0} \to \mathbf{m1}$ -3 $\neq \mathbf{m1}$ -3 $\to \mathbf{m0}$). The role of $\mathbf{m0}$ as a donor stabilizes the charge separation. The stabilization of CT states is modulated by the substituents, in $\mathbf{SF2}$ and $\mathbf{SF3}$ MeO and $\mathbf{Net_2}$ are electron donating groups, which reduce the charge transfer between both halves compared with $\mathbf{SF1}$, as reflected by the high mixing with $\pi\pi^*$ configurations in the minimum of CT PES (Table 8.10).

State	ΔE	f	character	%
SF1	3.23	0.004	$m0 \rightarrow m1$	96.1
$\mathbf{SF2}$	3.44	0.000	$\mathrm{m2}(\mathrm{n}\pi^*)/\mathrm{m0} o \mathrm{m2}$	70.9/22.6
SF3	3.47	0.000	$\mathrm{m3}(\mathrm{n}\pi^*)/\mathrm{m0} \rightarrow \mathrm{m3}$	83.7/9.6

Table 8.10: Vertical de-excitation energy (in eV), oscillator strength (f), and state character from the energy minimum of the lowest CT state of **SF1-3** computed at the CT state geometry.

8.2.3 Spin-orbit coupling

8.2.3.1 A brief experimental background

The fast generation of long-life triplets (9.22, 3.84 and 4.04 μ s in SF1-3, respectively) was experimentally detected by femtosecond transient absorption. In cyclohexane a long-lived excited-state absorption (ESA) signal is observed for SF1-3, which decays about 200 ns in SF1 similar results are obtained in SF2 and SF3. This signal is associated to the formation of a $\pi\pi^*$ triplet. With the increase of polarity of the solvent (up to DMSO) the ESA band associated to the triplet gradually disappears until it is only present in SF3 (about 4 ns after excitation). A ground-state bleach signal is observed before the signal of triplet formation, indicating the presence of another state that acts as an intermediate in the triplet generation. According with the computational characterization presented in Table 8.2.2, the excited states that are energetically accessible from S* have important contributions of CT configurations. Experimental observations and computational predictions suggest an excited state dynamics as Figure 8.16 illustrates for SF3. After the population of the $\pi\pi^*$ bright state, the CT state is formed and generates the triplet state by charge recombination.

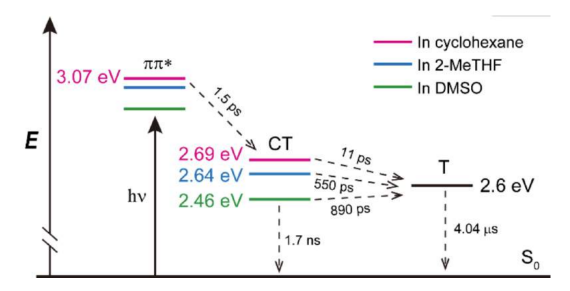


Figure 8.16: Simplified Jablonski diagram of the photophysics processes in non-polar to polar solvents of **SF3**. Figure adapted from reference (40)

In summary, a fast conversion from singlet to triplet process is experimentally detected and depends in the stabilization of CT states provided by the solvent polarity. The ISC rate constant (k_{ISC}) can be approximated by the Marcus form of the Fermi's golden rule:

$$k_{ISC} = \frac{2\pi}{h} < \Psi_{CT}^{0} |H_{SO}| \Psi_{T}^{0} >^{2} (4\pi\lambda RT)^{-\frac{1}{2}} \times e^{\frac{(\Delta E + \lambda)^{2}}{4\lambda RT}},$$
 (8.8)

where H_{SO} is the spin-orbit Hamiltonian, λ is the reorganization energy, and ΔE is the splitting between CT and triplet state.

To describe the generation of triplets by means of CT states, the spin-orbit coupling between them was computed at the geometrical minimum of the CT state. Table 8.11 shows that at this structure there are several triplets close in energy with $\langle CT|H_{SO}|T_n \rangle \neq 0$. Large H_{SO} couplings are found with triplets that have large mix of configuration with CT states

The family of spiro-compounds analyzed here are characterized by their tunable SOCT-ISC dependent of the solvent polarity.

In non-polar environments, the CT state assist the SOCT-ISC, providing rate constants that can be tuned by 2 orders of magnitude according with the electron withdrawing or donating character of the substituent group. In polar solvents the CT state is stabilized and can acts as a trap state favoring other deactivation pathways different from the triplet formation.

These results highlight the main role of the low energy CT states assisting and promoting the fast $S \to T$ conversion.

State	ΔE	SOC	Character	%	
SF1					
T_1	-0.16	1.00	$\mathrm{m1}(\mathrm{n}\pi^*)/\mathrm{m0} o \mathrm{m1}$	72.3/9.3	
T_2	-0.12	1.62	$\mathrm{m0}(\pi\pi^*)/\mathrm{m0} ightarrow \mathrm{m1}$	70.6/10.2	
T_3	-0.06	0.77	$\mathrm{m}0 ightarrow \mathrm{m}1/\mathrm{m}1(\mathrm{n}\pi^*)$	73.3/15.0	
T_4	0.06	0.18	$\mathrm{m0}(\pi\pi^*)/\mathrm{m0} o \mathrm{m1}$	68.5/5.1	
T_5	0.09	0.40	$\mathrm{m1}(\pi\pi^*)/\mathrm{m0} ightarrow \mathrm{m1/m1}(\mathrm{n}\pi^*)$	70.4/4.6	
SF2					
T_1	-0.47	0.89	$\mathrm{m2}(\mathrm{n}\pi^*)/\mathrm{m2} o \mathrm{m0}$	69.7/21.7	
T_2	-0.43	58.06	$\mathrm{m2} o \mathrm{m0}$	78.7	
T_3	-0.15	0.12	$\mathrm{m0}(\pi\pi^*)/\mathrm{m0} o \mathrm{m2}$	75.8/4.7	
T_4	-0.13	0.69	$ m m0(\pi\pi^*)/m0 ightarrow m2$	74.0/7.7	
T_{5}	-0.08	1.96	$\mathrm{m2}(\pi\pi^*)/\mathrm{m2} o \mathrm{m0}$	76.0/5.4	
T_6	0.13	0.02	$\mathrm{m0}(\mathrm{n}\pi^*)/\mathrm{m}2 o \mathrm{m}0$	87.5/6.4	
SF3					
T_1	-0.61	49.70	$m3(iCT)/m0 \rightarrow m3$	75.3/5.9	
T_2	-0.47	2.13	$\mathrm{m}3(\mathrm{n}\pi^*)/\mathrm{m}0 ightarrow\mathrm{m}3$	81.5/9.6	
T_3	-0.20	27.25	m3(iCT)	76.6	
T_4	-0.20	0.69	$\mathrm{m0}(\pi\pi^*)/\mathrm{m3} o \mathrm{m0}$	74.3/4.6	
T_5	-0.17	1.61	$\mathrm{m0}(\pi\pi^*)/\mathrm{m3}(\mathrm{iCT})/\mathrm{m3}(\mathrm{n}\pi^*)/\mathrm{m3} \rightarrow \mathrm{m0}$	68.8/7.4	
T ₆	0.10	0.01	$m0(n\pi^*)/m3 \rightarrow m0$	74.0/15.3	

Table 8.11: Relative energies (in eV) and spin-orbit couplings (in cm⁻¹) between the lowest CT singlet and low-lying triplet states of **SF1** - **3** computed at the CT state geometry. Character of the transition has been indicated in % for the main orbital contributions.

8.3 Spiroconjugation of a diradicaloid-pair

spiro-**DBP1** has suitable electronic features to carry out efficient SF, as highlighted in the first section of this chapter. This means, a good excited state distribution in the monomers is predicted, specifically the $E(S^*) - 2E(T_1)$ value is positive and small. On the other hand, **Fn** and **DBP2** chromophores can be regarded as diradicaloids (see Table 8.1), meaning the energy of T_1 is considerably low with respect

to S* excitation energy, and the singlet fission process is not favored. Interestingly, the tetra-phenyl substituted **DBP2** has been synthesized, and theoretically and experimentally analysed,(8) reporting a $\Delta E_{ST} = 5.68$ kcal/mol at B3LYP/6-31G(d,p), that is in good agreement with our calculations (8.07 kcal/mol). In the same study, an extended version of the π -conjugated **DBP2**, two fused units, (Figure 8.17 c) was characterized to hold a strong diradical character ($\Delta E_{ST} = 2.12$ kcal/mol, determined by SQUID measurements, and 1.83 kcal/mol at B3LYP/6-31G(d,p) level). Recently, experimental group of Professor Eiichi Nakamura from The University of Tokyo achieved the synthesis of the *spiro*-**DBP2**(41), while the group of the University of Malaga leaded by Prof. Juan Casado characterized its spectroscopic features. Together with them, we described and rationalized the electronic structure of this compound.

Through this chapter it has been suggested that the main role of the spiro linkage is structural, allowing both halves to be together in a rigid conformation with weak interaction derived from the orthogonality of the π -backbones, which also results in a low-lying pair of charge transfer states. This means, that the final properties of the spiro system are largely dictated by the electronic structure of the "independent" monomers. Hence, is it reasonable to envisage the spiro-DBP2 as a pair of independent diradicaloids? Or the diradicaloid character of DBP2 can lead to the interaction between both fragments to form a tetraradicaloid?

Here, I'll present the theoretical and computational investigation of the electronic structure of the spiro-**DBP2**. For sake of comparison, the pro-spiro, spiro and an extended monomer were analyzed at the same level of theory (see the Appendix). In order to differentiate with the **DBP2** and spiro-**DBP2**, hydrogen atoms in the pro-spiro position where replaced by phenyl groups, thus hereafter the pro-spiro chromophore will be named as ph**DBP2**, its extended version, based on bis(dibenzo)pentalene (ph**BDBP2**) exph**DBP2**, and the spiro dimer spiro-ph**DBP2** (Figure 8.17).

Figure 8.17: Structure of $ph\mathbf{DBP2}$ (a), the spiro dimer $spiro-ph\mathbf{DBP2}$ (b), and $ph\mathbf{BDBP2}$ (c).

All of them hold a singlet open-shell ground state computed at the DFT level of theory. The closed-shell singlet is higher in energy, with open-shell to closed-shell gaps following the $ph\mathbf{DBP2} < spiro-ph\mathbf{DBP2} < ph\mathbf{BDBP2}$ tendency. Adiabatic

energy gaps, in kcal/mol, are 5.79 and 1.78 for $ph\mathbf{DBP2}$ and $\mathbf{ex}ph\mathbf{DBP2}$, respectively, computed with the B3LYP functional (with ω B97X-D are 3.48 and 0.78, respectively). There is a noticeably drop of ΔE_{ST}^{ad} , i.e., substantially grow of diradical character, from $ph\mathbf{DBP2}$ to $ph\mathbf{BDBP2}$. This can be explained as the lost of interaction between radicals in the molecular ends through the π -conjugated system as an effect of the molecular size increment. Interestingly, the adiabatic quintet in state in $spiro-ph\mathbf{DBP2}$ is computed at 10.9 (6.69) kcal/mol above the ground state open-shell singlet at . This result indirectly indicates the presence of two triplets, one in each $ph\mathbf{DBP2}$ unit of the spiro-compound. Further characterization of the radical nature of these compounds is present in Table 8.12.

Geometry	Compound	state	ΔE	N_u	y_0	y_1
B3LYP	33LYP ph DBP2		-	0.94	0.40	0.06
		T_1	10.10	2.10	1.00	0.04
	$spiro$ - $ph\mathbf{DBP2}$	S_0	-	2.02	0.44	0.43
		T_1	9.11	3.10	0.98	0.46
		T_2	9.16	3.11	0.97	0.46
		Q_1	18.36	4.19	1.00	1.00
	exphDBP2	S_0	-	1.68	0.80	0.04
		T_1	0.58	2.08	1.00	0.03
$\omega \mathrm{B}97\mathrm{X}\text{-}\mathrm{D}$	$ph\mathbf{DBP2}$	S_0	_	1.13	0.51	0.05
	-	T_1	6.66	2.08	1.00	0.03
	spiro- ph DBP2	S_0	-	2.39	0.54	0.53
		T_1	6.02	3.29	0.90	0.64
		T_2	6.09	3.29	0.90	0.64
		Q_1	12.15	4.17	1.00	1.00
	$\mathbf{ex}ph\mathbf{DBP2}$	S_0	-	1.75	0.84	0.03
		T_1	0.21	2.06	1.00	0.02

Table 8.12: Low-lying state relative energies (ΔE) in kcal/mol), number of unpaired electrons (N_u), and diradical (y_0) and tetraradical (y_1) indices of $ph\mathbf{DBP2}$, $spiro-ph\mathbf{DBP2}$ and $\mathbf{exph\mathbf{DBP2}}$ computed at the RAS-SF/6-31G(d,p) level for the ground state geometry optimized at B3LYP and ω B97X-D functionals.

The following discussion will be done using the results obtained with the B3LYP optimized geometry, but it is worth noting that conclusions are qualitatively the same with geometries obtained with ω B97X-D. Large N_u values of S₀ of the three compounds reaffirms their open-shell character. As presented before phDBP2 is a moderate diradicaloid with $y_0 = 0.40$. After construction of the spiro-conjugated system, two vertical triplets are calculated close to the energy of the phDBP2

 T_1 , in $spiro-ph\mathbf{DBP2}$ T_1 and T_2 are the \pm linear combinations of the configurations that localize the triplet in one or another $ph\mathbf{DBP2}$ unit. Additionally, the Q_1 state is $\sim 2T_1$ of $ph\mathbf{DBP2}$. This small energy difference can be associated to the diradical-diradical interaction through the spiro connection (as discussed in Figure 8.5 and Equation 7.19). This result points toward the rise of a system with ~ 4 unpaired electrons, i.e., a tetraradical. By comparison of the ground state of $spiro-ph\mathbf{DBP2}$ and $ph\mathbf{DBP2}$. In latter there are 2.02 unpaired electrons, approx twice than the monomer. Moreover, due to their similar value of y_0 (around 0.4), but $spiro-ph\mathbf{DBP2}$ has a $y_1 = 0.43$), meaning, it should be named as a tetraradicaloid instead of diradicaloid. $exph\mathbf{DBP2}$ S_0 and T_1 states are almost degenerated with an energy gap inferior to 1 kcal/mol, and a diradical index $y_0 > 0.8$. Clearly, $exph\mathbf{DBP2}$ holds the strongest diradical character, but $spiro-ph\mathbf{DBP2}$ is the only tetraradicaloid of the series.

The radical character of spiro-ph**DBP2** was determined through experiments of molar magnetic susceptibility $(\chi_M T - T)$. Three main signals are characterized as one singlet, triplet, and quintet that can be assigned to the S₀ (ground state), T_{1,2}, and Q₁ in Table 8.12, respectively. A small amount of an impurity with diradical character is also detected ($\Delta E_{ST} = 0.09 \text{ kcal/mol}$). The impurity is predicted to be produced by the strong interaction between two spiro-compounds when they are close to each other. By steric hindrance the π - π stacking between two monomeric backbones is not possible, but radicals localized in the molecular ends can interact covalently to form a σ -bond between them, (Figure 8.18 a).

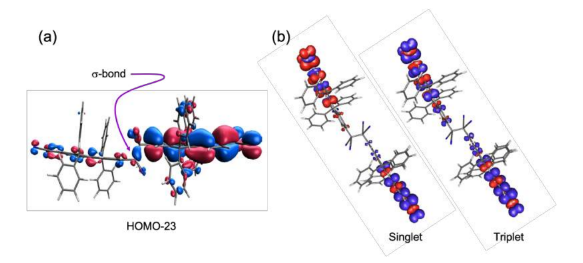


Figure 8.18: σ -bonding orbital (HOMO-23) (a) and spin density representation of the lower singlet and triplet (b) of the σ -dimer at the ω B97X-D/6-31G(d).

This situation was evaluate in a σ -bonded dimer of $ph\mathbf{DBP2}$. The computed electronic energy binding between two monomers is ~ 25.3 kcal/mol. Two unpaired electrons are localized one in each extreme of the σ -dimer (Figure shows the spin

density of lowest singlet and triplet states 8.18 b). Due to the large distance between both radicals their interaction fades away and the singlet-triplet energy gap tends to zero ($\Delta E_{ST} < 0.03 \text{ kcal/mol}$), in good agreement with the aforementioned singlet-triplet energy gap experimentally determined (0.09 kcal/mol).

8.4 Appendix

8.4.1 Computational details

Molecular structures have been optimized within the density functional theory (DFT) in conjunction with the CAM-B3LYP energy functional.

The ground state diradical character (y_0) has been quantified as the occupation number of the lowest unoccupied natural orbital (LUNO).(42) The number of unpaired electrons of electronic states has been evaluated according to the linear Head-Gordon index (N_u) .(43)

All calculations were performed with the Q-Chem program. (44) Molecular orbitals (MOs), RAS-srVWN natural orbitals (NOs) and FODs have been plotted with a homemade version of Avogadro (Version 1.2). (45)

8.4.1.1 Spiroconjugated dimers in singlet fission

Optimized geometries were confirmed to be minima on the ground state potential energy surface (PES) by means of frequency analysis within the harmonic approximation. Raw frequencies are 17.68, 22.21, 24.86, 35.70, 42.42 cm⁻¹. Excited states have been computed employing the restricted active space configuration interaction method with the *hole* and *particle* approximation and the spin-flip excitation operator (RAS-SF),(? 46) and with the inclusion of short-range DFT correlation through a range-separation scheme (RAS-srDFT).(?) The lowest restricted open-shell Hartree-Fock (ROHF) high-spin triplet (quintet) state is used as the reference configuration for all RAS-srVWN calculations of the studied monomers (spiro dimers) with 4 (8) electrons into 4 (8) π -orbitals in the RAS2 space. RAS1 and RAS3 subspaces include all virtual and doubly occupied orbitals, respectively.

The spatial distribution of unpaired electrons has been represented by means of the fractional occupation density (FOD). Characterization of electronic excited states of spiroconjugated compounds in terms of local excitations (LE), charge transfer (CT) or resonance (CR), and triplet-triplet configurations (TT), i.e. diabatic states, has been carried out by the decomposition of the adiabatic state wave functions and density matrices. (47; 48) CT diabatic states have been also computed with the constrained-DFT (C-DFT)(?) method with the CAM-B3LYP functional.

8.4.2 **SOCT-IC**

Vertical excitation energies at different molecular geometries (ground, excited and CT states) were calculated with the LR-TDDFT with the Tamm-Dancoff approximation (TDA) and with the same functional. RAS-SF calculations were performed with the high-spin quintet as the reference configuration, with four electrons in four orbitals in the RAS2 space, all occupied (virtual) orbitals in the RAS1 (RAS3) space, and the 6-31G(d,p) basis set.

Geometry relaxation on the excited states potential energy surfaces were performed with B3LYP and CAM-B3LYP functionals for different excited triplet and singlet states. Solvation effects of cyclohexane and DMSO solvents were included through the SMD solvation model. Inter-state spin orbit couplings between low-energy singlets and triplets were computed at the same electronic structure level (CAM-B3LYP/6-311G(d,p)) taking into account one-electron contributions to the Breit-Pauli spin-orbit Hamiltonian.

8.4.3 spiro-DBP2

Molecular geometries of ground state open- and closed-shell configurations and the lowest triplet state have been optimized within the density functional theory (DFT) in combination with the ω -B97XD and B3LYP and in addition to CAM-B3LYP, and the 6-31G(d,p) basis set. Formation of the σ -dimer has been simulated by a pair of Q₁ molecules at the ω B97X-D/6-31G(d) computational level. Restricted active space spin-flip (RAS-SF) calculations were done by including all doubly occupied and virtual orbitals in RAS1 and RAS3 subspaces, respectively. In the active space are included 4 electrons in the 4 frontier π -orbitals, and 4 electrons in the 5 orbitals in the σ -dimer; while RAS2 in the spiro-compound included 8 electrons in 8 orbitales. The high spin triplet (monomers) and quintet (spiro and σ -dimer) Hartree-Fock determinants were used as the reference configurations. The 6-31G(d,p) basis set was used in all RAS-SF calculations.

Vertical excitation energies to the lowest excited states where obtained with the timedependent (linear response) version of DFT (TDDFT), and with the ω – B97X - Dfunctional and 6-31G(d, p) basis set.

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Chapter 9

General conclusions

The results exposed throughout these pages present the full characterization of the radical character of organic systems by means of adequate electronic structure methods. More specifically, the origin of the diradical (polyradical) character was rationalized for each system and put in relation with the most remarkable molecular properties associated with the presence of unpaired electrons.

To highlight the structure-diradical character relationships established in this work, the radical characters of [6]Ac, [6]CC, and 2 can be used as an example (Chapter 4). All of them are composed by six fused benzene rings having zigzagedges. When rings are fused following a one dimensional pattern, the experimentally well-known [6]Ac holds a moderate diradicaloid character. If this linear arrangement is closed in a belt shape, one obtains the [6]CC compound that exhibits a stronger diradicaloid character, resulting in a severe chemical instability, which has prevented its synthesis. Finally, the triangular arrangement of six fused rings in 2 presents a triplet-spin ground state and can be considered a perfect diradical.

Besides these purely topological considerations, a relationship between the radical character of a system and the spatial arrangement of the radical sites within the molecular structure can be drawn. This has been demonstrated in Chapter 6, where the radical-radical interaction (ferromagnetic or antiferromagnetic) is modulated by the characteristics of the bridge linking the individual units that carry an unpaired electron, namely its flexibility, length and nature of π -conjugation. Alternatively, the controlled tuning of radical character in organic molecules can be achieved through chemical modifications of the molecular structure, as addressed in Chapter 4. More specifically, heteroatom substitution, which has been shown to affect the spin multiplicity of the ground state, provides a viable strategy to design new stable (di)radicaloid systems.

The establishment of the aforementioned structure-properties relationships called

for methods incorporating strong correlation effects to achieve the proper description of the radical nature of the systems. Concretely, the characterization of (poly)radical systems of small to large size reported in this thesis mainly rely on the use of the powerful RAS-SF family of methods. Indeed, these approaches provide a multiconfigurational wave function able to deal with several strongly correlated electrons. Moreover, this accurate treatment is achieved at reasonable computational cost as evidenced from the study of the large macrostructures reported in Chapter 6. The theoretical-experimental synergy in which some parts of this thesis has been carried out highlights the crucial role of such quantum chemistry methods in the understanding of radical systems. In this context, I have been able to provide an interpretation of the spectroscopic features of newly synthesized stable diradical dianions oligothiophene derivatives and illustrate at the molecular level their peculiar ability to stabilize polaron-pairs (Chapter 5). In the same chapter, I have rationalized the temperature related interconversion between two stable conformers with different diradical character.

With a more applied aspect in mind, and from the understanding gained from the previous chapters, it is evident that key photophysical processes at stake in organic based devices can beneficiate from the presence of radicals. In this regard, I have focused my interest on the unique spin-allowed singlet fission (SF) process that is currently considered as one of the most promising approaches in the development of the next generation photovoltaic technologies. Among the requirements for this process to occur efficiently, the control of the singlet-triplet energy gap, which can be modulated indirectly via the alteration of the radical character, is crucial. From the theoretical point of view, I have extended the well-established five state model to describe SF by including doubly excited states. The obtained results from the application of the new model show that indeed these states can potentially play an important role in the SF mechanism as the initial or mediator states (Chapter 7). In particular, I have identified the parameters, i.e., relative energy of states and adequate electronic couplings, controlling the involvement of the doubly excited states in the SF process, and I have explored the optimal conditions for the fast and efficient generation of the triplet-pair state.

Ultimately, the last chapter is devoted to the in-depth exploration of spiroconjugated molecules on the basis of the structure-diradical relationships and photophysical processes derived from the previous chapters. These systems hold a very unique chromophoric arrangement (*i.e.*, two weakly interacting perpendicular chromophores close in space) via the spiro atom. Spiro-compounds exhibit low energy charge-transfer (CT) states, which makes them promising candidates for CT-mediated photophysical processes such as SF or intersystem crossing.