Density Functional Calculations

Distortion-Controlled Redshift of Organic Dye Molecules

Ayush K. Narsaria,[a] Jordi Poater,[b, c] Célia Fonseca Guerra,[a, d] Andreas W. Ehlers,[e, f] Trevor A. Hamlin,[a] Koop Lammertsma,*[a, f] and F. Matthias Bickelhaupt*[a, g]

Abstract: It is shown, quantum chemically, how structural distortion of an aromatic dye molecule can be leveraged to rationally tune its optoelectronic properties. By using a quantitative Kohn–Sham molecular orbital (KS-MO) approach, in combination with time-dependent DFT (TD-DFT), the influence of various structural and electronic parameters on the HOMO–LUMO gap of a benzenoid model dye have been investigated. These parameters include 1) out-of-plane bending of the aromatic core, 2) bending of the bridge with respect to the core, 3) the nature of the bridge itself, and 4) π–π stacking. The study reveals the coupling of multiple structural distortions as a function of bridge length and number of bridges in benzene to be chiefly responsible for a decreased HOMO–LUMO gap, and consequently, red-shifting of the absorption wavelength associated with the lowest singlet excitation (λ ≈ 560 nm) in the model cyclophanes systems. These physical insights together with a rational approach for tuning the oscillator strength were leveraged for the proof-of-concept design of an intense near-infrared (NIR) absorbing cyclophane dye at λ = 785 nm. This design may contribute to a new class of distortion-controlled NIR absorbing organic dye molecules.

Introduction

Modulation of frontier orbital energies has elucidated many aspects of materials and synthetic chemistry. Illustrative is the aspect of decreasing the HOMO–LUMO gap (ΔE_{H-L}), which has found widespread applications in solar cell design,[1] long-wavelength absorbing photosensitizers,[2] near-infrared (NIR) fluorophores in bioimaging of deep tissues,[3] and NIR emitters in organic electronics.[4] Decreasing the ΔE_{H-L} in organic compounds has been attributed to extending π conjugation,[5] enhancing planarity,[6] or adding donor–acceptor motifs[7] to conjugated structures (see Scheme 1). By quantifying their interplay in the ΔE_{H-L}, we recently developed design principles for NIR absorbing dyes.[8] Despite the fact that the abovementioned concepts are all efficient in decreasing the ΔE_{H-L}, there

Scheme 1. Previous approaches (top)[5–7] and our proposed distortion-controlled approach (bottom) for decreasing and tuning the HOMO–LUMO gap in organic dye molecules. D = donor, A = acceptor.

[a] A. K. Narsaria, Prof. Dr. C. Fonseca Guerra, Dr. T. A. Hamlin, Prof. Dr. K. Lammertsma, Prof. Dr. F. M. Bickelhaupt
Department of Theoretical Chemistry and Amsterdam Center for Multiscale Modeling (ACMM), Vrije Universiteit Amsterdam
De Boelelaan 1083, 1081 HV Amsterdam (The Netherlands)
E-mail: t.a.hamlin@vu.nl
k.lammertsma@vu.nl	f.m.bickelhaupt@vu.nl
[b] Prof. Dr. J. Poater
ICREA, Pg. Lluís Companys 23, 08010 Barcelona (Spain)
[c] Prof. Dr. J. Poater
Departament de Química Inorgànica i Orgànica & IQCTUB
Universitat de Barcelona, Martí i Franquès 1-11, 08028 Barcelona (Spain)
[d] Prof. Dr. C. Fonseca Guerra
Gorlaeus Laboratories, Leiden Institute of Chemistry
Leiden University, Einsteinweg 55, 2333 CC, Leiden (The Netherlands)
[e] Dr. A. W. Ehlers
van’t Hoff Institute for Molecular Sciences, University of Amsterdam
Science Park 904, 1098 XH Amsterdam (The Netherlands)
[f] Dr. A. W. Ehlers, Prof. Dr. K. Lammertsma
Department of Chemistry, University of Johannesburg
Auckland Park, Johannesburg, 2006 (South Africa)
[g] Prof. Dr. F. M. Bickelhaupt
Institute of Molecules and Materials (IMM)
Radboud University Nijmegen, Heyendaalseweg 135
6525 AJ Nijmegen (The Netherlands)

Supporting information and the ORCID identification number(s) for the author(s) of this article can be found under:
https://doi.org/10.1002/chem.201905355.
is still a lack of design concepts and a systematic framework to rationalize such concepts. Therefore, we aim to utilize the concept of structural distortion as a tool to modulate the $\Delta E_{\text{UL}}$ in pursuit of the rational design of intense NIR absorbing organic dyes.

By controlling the distortion of aromatic $\pi$ systems, their electronic structure can be tuned (see Scheme 1). We have already shown that incorporating a heavy heteroatom into the aromatic core of benzene ($Y = \text{Si, Ge, P}$) decreases the $\Delta E_{\text{UL}}$ due to in-plane structural distortion, which causes a red-shift in the lowest excitation.[6] We further showed that distortion into a boat-like conformation accelerated the Diels–Alder cycloaddition, relative to that of benzene (1). [9] A related example concerns hoop-shaped benzene rings, of which the para positions are covalently connected.[10] Reducing the number of benzene rings (n) decreases the hoop size of these [n]cycloparaphenylenes and results in smaller $\Delta E_{\text{UL}}$ values; this has been attributed to enhanced conjugation among the neighboring $\pi$ orbitals due to a decrease in torsional angles. Recently, structural distortion of a Sm@C88 carbon cage was found to affect the $\Delta E_{\text{UL}}$.[11] With increased pressure, the $\Delta E_{\text{UL}}$ becomes smaller as the cage is distorted from its equilibrium ellipsoid shape to spherical, to peanut-shaped, followed by eventual collapse. Other carbon nanostructures have shown similar behavior.[12] There are also extensive studies on distorted porphyrin rings that exhibit red-shift compared with their planar counterparts.[13] Collectively, these studies suggest a causal relationship between increased structural distortion and decreased $\Delta E_{\text{UL}}$, and therefore, a sound rationalization of the effect of controlled distortion on $\Delta E_{\text{UL}}$ is warranted.

Cyclophanes,[14–17] with their strained benzene rings, are ideal candidates to enhance our understanding of the influence of structural distortion on electronic properties. Increased strain in the cyclophane is known to decrease its $\Delta E_{\text{UL}}$, thereby red-shifting the absorption spectrum,[16] but the underlying mechanism is far from trivial. Recently, we delineated this relationship by bending benzene into a boat conformation to examine the Diels–Alder reactivity of [5]cyclophanes.[18] Herein, we expand the insights gained from the reactivity study toward tuning of the optoelectronic structure. The focus is on how distortion of the benzenoid core controls the modulation of $\Delta E_{\text{UL}}$ and how it can be leveraged to rationally design a distortion-controlled NIR absorbing organic dye. In addition to the proper absorption wavelength, a large oscillator strength is fundamental for the efficacy of a NIR chromophore. Therefore, we also undertake a rational approach to tune the oscillator strength. In addressing these questions, we explore the effect of $\pi$-$\pi$ stacking, heteroatom functionalization, and donor–acceptor architectures on the electronic and optoelectronic properties. We use a fragment-based approach to obtain mechanistic insights into the decrease of $\Delta E_{\text{UL}}$ upon subsequent introduction of various modes of structural distortion. The novelty of the present work is contained not only in the concrete applications, but in the fact that we combine various distortion-based tuning handles for tuning both the excitation energy and the oscillator strength, and trace the underlying physics, such that the emerging insights can be applied in rational design. All calculations have been performed by using the Amsterdam Density Functional (ADF) 2017 quantum chemistry package developed by SCM[19] and the optimized structures have been illustrated by CYLview.[20]

Computational Details

Structure optimization

The electronic ground-state geometry optimizations were performed with the dispersion-corrected GGA-BLYP-D3(BJ) functional[21] in combination with the TZ2P[22] Slater-type basis set and a small frozen core. The BLYP-D3(BJ) functional has been shown to satisfactorily reproduce the structural parameters of stacked systems.[21] Scalar-relativistic effects were included by using the scalar zeroth-order regular approximation (ZORA).[23] All model and benchmark molecules exhibit (local) energy minima, as confirmed by vibrational analysis to have zero imaginary frequencies.

Kohn–Sham molecular orbital (KS-MO) analysis

KS-MO[24] analyses have been performed at the same level of theory by using the all-electron TZ2P basis set (i.e., no frozen core approximation). To uncouple the effect of different structural distortions from each other, we performed KS-MO analyses on relaxed structures. Our quantitative molecular orbital interaction analyses, based on fragments, provides accurate and detailed insights into and easy-to-interpret models of the response of the electronic structure towards structural and electronic perturbations.

Atomic Voronoi overlap (AVO)

To assess more quantitatively in which part of the molecule overlap arises between the fragment molecular orbitals (FMOs) that constitute the pertinent overall molecular orbital, we present the concept and method of the AVO, which has been inspired by the concept of Voronoi deformation density (VDD) atomic charges.[25] The AVO $A_i^{\text{AVO}}$ on atom A is computed as the numerical integral of the overlap density $\rho_i(r)$ between FMO $\phi_i(r)$ on fragment I and FMO $\phi_j(r)$ on fragment J (Eq. [1]) in the volume of the Voronoi cell of atom A [Eq. (2)], which is defined as the compartment of space bound by the bond mid-planes perpendicular to all bond axes between nucleus A and its neighboring nuclei, that is, the region in real space that is closer to a particular nucleus than to any other nucleus (c.f., the Wigner–Seitz cells in crystals). The computations have been carried out with ADF and the associated tool DENSIF.[19]
\[ \rho_1(r) = \bar{\rho}_1(r) \varphi_{1}(r) \]  
\[ S_{AA}^{AVO} = \int_{\text{Voronoi cell } A} \rho_1(r) \, dr \]

The interpretation of the AVO is straightforward and transparent: \( S_{AA}^{AVO} \) constitutes that part of the overlap, \( S_{ij} = \int \bar{\rho}_i(r) \varphi_j(r) \, dr \), that arises in the Voronoi cell of atom A. Equivalently, it can be considered as the amount of charge associated with the overlap density, \( \rho_1(r) \), located in Voronoi cell A.

Benchmark and excited-state calculations

Linear response time-dependent density functional theory (TD-DFT),\(^{[26]}\) within its adiabatic framework, has become the primary tool to simulate excited states of small- to medium-sized molecules.\(^{[27]}\) Despite its general applicability, there are instances in which it can fail to accurately describe the excited-state properties of low-lying \( \pi^* \pi^* \) states. For example, TD-DFT may yield an imbalanced description of the two lowest singlet states of our reference model system benzene 1 (oligoacenes, in general).\(^{[27]}\) Therefore, a benchmark study was carried out to select the appropriate TD-DFT functional among the hybrid PBE0,\(^{[28]}\) the meta-hybrid M06-2X,\(^{[29]}\) and the range-separated hybrid (RSH) functional CAMY-B3LYP\(^{[30]}\) to reproduce the experimentally measured \( \lambda_{\text{max}} \) (for details, see Section S1 and Table S1 in the Supporting Information). CAMY-B3LYP emerged as the best candidate in terms of accuracy (mean absolute deviation (MAD) relative to \( \lambda_{\text{max}} = 0.39 \) eV) and systematic consistency (\( R^2 = 0.99 \)), in line with a study by Jacquemin and coworkers.\(^{[31]}\) Recent studies by Hopf and co-workers also highlighted the accuracy of RSH functionals to accurately reproduce the absorption spectra of cyclophanes.\(^{[32]}\) Furthermore, it is known that RSH functionals improve the long-range asymptotic behavior of the exchange-correlation potential, enabling them to accurately compute strong charge-transfer excitations compared to other standard TD-DFT functionals.\(^{[33]}\) Compounds 7’ and 7” (see Figure S1 in the Supporting Information) are representative examples of charge-transfer-based excitation, for which we observe that CAMY-B3LYP (MAD = 0.12 eV) is more accurate than M06-2X (MAD \( \approx 0.22 \) eV). Benchmark studies by Tozer et al. showed that local excitations were also well reproduced by RSH functionals.\(^{[34]}\) In light of the highly delocalized and charge-transfer model systems investigated in our study, we have selected CAMY-B3LYP as the appropriate TD-DFT functional, in combination with the TZ2P basis set for obtaining accurate and reliable trends. All electrons are treated variationally (no frozen core), whereas scalar-relativistic effects are considered through the ZORA formalism. All of the calculations of model systems have been performed in vacuum to remove the solvatochromic effect on the absorption spectra. An additional TD-DFT calculation was performed on only 18 (see Figure 7) with the nonequilibrium continuum solvation model COSMO\(^{[35]}\) in the linear-response framework by using CHCl\(_3\) (DCM) as the solvent. Hereafter, we denote the \( S_1 \rightarrow S_1 \) excitation energy as \( E_1(S_1) \), which is composed of a predominant HOMO \( \rightarrow LUMO \) excitation (Table S2 in the Supporting Information), except for 1–4 and 9, in which \( S_1 \rightarrow S_0 \) excitation matches the nature of \( S_1 \rightarrow S_0 \) excitation, and thus, \( E_1(S_1) \) is discussed for a fair comparison.

Results and Discussion

The paper is organized as follows. The first section outlines the individual roles of various tuning parameters on the \( \Delta E_{\text{UL}} \) value of 1 (see Scheme 2), namely, 1) out-of-plane bending of the aromatic core (\( \alpha \)), 2) bending of the cyclophane bridge with respect to the aromatic core (\( \beta \)), 3) electronic nature of the bridge (X), 4) stacking of the \( \pi \)-conjugated cores, and 5) inter-core distance (\( d \)). All mechanistic studies have been performed on benzene model systems with the aim of understanding and quantifying the effects of the aforementioned structural and electronic tuning parameters on \( \Delta E_{\text{UL}} \). The results are illustrated in Figures 1–4. The second section investigates a series of small cyclophane systems to understand the relationship between the tuning parameters (1–5) and \( \Delta E_{\text{UL}} \). Then, we combine all of our findings for the design of more complex cyclophane-based model systems and rationally engineer a NIR absorbing cyclophane-like organic dye molecule. The TD-DFT results of the target system are presented in Figure 5 and Table 1. We conclude the study by tuning the oscillator strength of the lowest singlet excitation. The framework that emerges explains the decrease of \( \Delta E_{\text{UL}} \) upon increased structural distortion and can serve as a tool to design novel dyes with tailored optical properties, which is illustrated for a distortion-controlled NIR absorbing cyclophane.

Role of various tuning parameters

Out-of-plane bending (\( \alpha \))

Out-of-plane (boat-like) bending eliminates the degeneracy of the frontier orbitals of benzene as the symmetry is reduced from \( D_{6h} \) to \( C_2v \) (see Figure S3 in the Supporting Information). Upon increasing \( \alpha \) from 0 to 40°, the \( \pi \)-HOMO is destabilized, whereas the \( \pi^* \)-LUMO is stabilized, leading to a decrease in \( \Delta E_{\text{UL}} \). A more detailed explanation and analysis are given than that previously reported.\(^{[9]}\) It starts by combining two allylic \( C_2H_4 \) triradical fragments to form the two \( \sigma \) bonds and one \( \pi \) bond of 1 (Figure 1a). Bending destabilizes \( \pi \)-HOMO as the antibonding interaction increases between the two allylic \( \pi \)-HOMO FMOs, whereas the \( \pi^* \)-LUMO is stabilized due to a favorable interaction between the \( \pi^* \)-LUMO of one allylic fragment and the nuclei of the other. Increasing \( \alpha \) from 0 to 25° enhances the antibonding interaction due to an increase in overlap between the allylic \( \pi \)-HOMOs (\( S_{\text{out}-\text{in}} \) Table S3 in the Supporting Information). Figure 1b schematically depicts this increase in overlap to originate from 1) a favorable spatial orientation, leading to a larger through-bond overlap of the \( p_\pi \) lobes of the allylic \( \pi \)-HOMO on C2 and C3 (similarly for C6 and

\[ \text{Scheme 2. Schematic illustration denoting the various structural and electronic parameters studied systematically in benzene (ring in black) out-of-plane bending (in red); bending of the bridge with respect to the core (in green), bridge substituent (X in blue), stacking with another } \pi \text{ core (in orange), and inter-core distance (in brown).} \]
C5); and 2) a larger through-space overlap between the far-lying $p_a$ lobes of the terminal C1 and C4 atoms (see Figure 1b for atom numbering) due to a decrease in the C1–C4 distance from 2.80 to 2.68 Å. To quantify the extent to which these different modes of overlap contribute to the total overlap, we use the AVO method, which identifies the overlap between two FMOs in the Voronoi cell of a particular atom. For $\alpha = 25^\circ$, the AVO for the C1 and C4 atoms is significant, 0.022 each (25% of $\pi$-$\pi$ overlap, $S_{\pi\pi}^\alpha$, Figure 2c), upon forming the $\pi$-HOMO. Counterintuitively, the overlap between the allylic $\pi^a$-LUMOs ($S_{\pi\pi}^\alpha$, Figure 2c) decreases upon bending (see Table S3 in the Supporting Information) as the in-phase overlap ($S_{\pi\pi}^\alpha$ depicted in pink in Figure 2c) is compounded by the out-of-phase overlap ($S_{\pi\pi}^\alpha$ depicted in green in Figure 2c) as illustrated in Figure 1c. The pronounced lowering in energy of the $\pi^a$-LUMO of benzene is caused primarily by the stabilization of the two allylic $\pi^a$-LUMOs due to the positive potential that one diffuse FMO experiences upon penetrating the nuclei of the other fragment (see Figure 1a and Table S4 in the Supporting Information). This phenomenon of stabilizing interactions upon spatial proximity of molecular fragments, popularly referred to as charge penetration,[37] is well established for bonding in nonpolar molecules.[24a,38] The extent of this effect is evident upon plotting the contours of the allylic $\pi^a$-LUMOs wave functions for the flat and bent geometries (Figure 1d). If $\alpha = 25^\circ$, the $\pi^a$-LUMO of one allylic fragment penetrates more into the nuclei of the other (faint blue contour lines crossing the C4 nuclei, see Figure 1c), relative to planar benzene (same holds for the other allylic $\pi^a$-LUMO). These small-amplitude contour lines penetrating the C4 (or C1) nucleus of one allylic fragment stem from the bent $p_a$ orbitals on C2 (or C3) and C6 (or C5) of the other fragment.

The large change in electronic structure upon bending, specifically in the context of a smaller $\Delta E_{H-L}$, has a direct impact on the photophysical properties. An increase in $\alpha$ from 0 to 40° redshifts $E_0(S_0)$ by nearly 1.20 eV (see Table S5 in the Supporting Information). Interestingly, in all cases, except if $\alpha = 40^\circ$, the $S_1\rightarrow S_0$ transition is composed of two singly excited configurations, HOMO$\rightarrow$LUMO + 1 and HOMO$\rightarrow$1–LUMO, and all of them exhibit zero oscillator strength. However, if $\alpha = 40^\circ$, the large decrease in $\Delta E_{H-L}$ changes the nature of the lowest singlet excitation to a predominant HOMO$\rightarrow$LUMO transition, with an increased oscillator strength from 0.00 to 0.04 (see Table S5 in the Supporting Information).

**Bending the bridge ($\beta$)**

The structural deformation of benzene can also be accomplished by bending of the cyclophane bridge, $\beta$, to induce a decrease in the $\Delta E_{H-L}$ (Figure 3a and Figure S5 in the Supporting Information), in accordance with our study on aromatic Diels–Alder reactions.[10] The effect can be analyzed by considering the interaction between [H–H] and [C2H2]; triplet diradical fragments to form benzene.

Symmetry breaking of the bent system allows for mixing of the otherwise orthogonal $\sigma$ and $\pi$-FMOs and overlap of the $s$ lobes of the hydrogen atoms with the $\pi$ orbitals of the aromatic core. The frontier MOs of benzene are composed of three or-

---

**Figure 1.** (a) Schematic $\pi$-FMO interaction diagram between two equivalent C3H6 quartet triradical fragments based on quantitative KS-MO analysis depicting the effect of out-of-plane bending ($\alpha$) on $\Delta E_{H-L}$ in comparison with flat benzene (in gray). Schematic representation of (b) different modes of $S_{\pi\pi}$ between allylic $\pi$-HOMOs and (c) $S_{\pi\pi}$ between allylic $\pi^a$-LUMOs. (d) Side view of the allylic $\pi^a$-LUMO wave function plotted on a cut plane through the [H–C1–C4–H] moiety and perpendicular to the planes of the two allylic fragments (−0.03–0.03 au range). The contour of only one allylic $\pi^a$-LUMO is plotted for clarity; the other is symmetrical.
and from of d transition changes to ap redominantly T (left) a overlap causes greater s overlap and ad ecrease in overlap (see Fig-ure 3b), and consequently, gives a smaller value of $\Delta E_{\text{exc}}$.

This decrease in the $\Delta E_{\text{exc}}$ also influences the optoelectronic structure. Similar to the case of $\alpha$, a gradual increase of $\beta$ from 0 to 40$^\circ$ also redshifts $E_{\text{L}}(S_1)$, although the degree of redshift is smaller in the case of increasing $\beta$ compared with that of increasing $\alpha$ (red-shift of 1.17 eV versus 0.85 eV as $\alpha$ and $\beta$ go from 0 to 40$^\circ$, respectively; see Tables S5 and S6 in the Supporting Information). Again, exactly the same as $\alpha$, if $\beta = 40^\circ$, the nature of the $S_1\rightarrow S_2$ transition changes to a predominantly HOMO$-$LUMO transition and the oscillator strength also becomes non-negligible.

**Electronic nature of the bridge (X)**

The third modification concerns the electronic effect of the bridge itself, which we have evaluated by comparing bent benzene (Figure 3a, left) with substituted bent 1,4-dimethylbenzene (Figure 3c), in which the two methyl groups serve as a simplification for the (CH$_3$)$_2$桥 bridge of paracyclophane. The two $\sigma$-donating methyl groups destabilize mostly the HOMO ($\Delta E = 0.53$ eV) and less the LUMO ($\Delta E = 0.25$ eV), resulting in a net decrease of the $\Delta E_{\text{exc}}$. The destabilization of the HOMO results from admixing the rather high-energy $\sigma$-HOMO of the [CH$_3$:CH$_3$]: triplet diradical fragment with the $\pi$-HOMO of the [CH$_3$:CH$_3$]: triplet diradical fragment in an antibonding fashion ($S_{p\pi} > 0$, Figure 3c and Figure S7 in the Supporting Information). This is in sharp contrast to the parent system because the [H$-$H]: triplet diradical fragment has no $\sigma$/$\pi$-HOMO (Figure 3a and Figure S7 in the Supporting Information).

**$\pi$-$\pi$ Stacking**

The final parameter to analyze in this section is the effect of stacking of the benzene rings, which we have studied for the co-facial benzene dimer ($D_{2h}$). There are many studies which show that $\pi$-$\pi$ stacking can radically change the electronic structure of the stacked system, especially the $\Delta E_{\text{exc}}$, and consequently, the optoelectronic properties relative to the monomer. Indeed, we also observe that the introduction of a second molecule of benzene decreases $\Delta E_{\text{exc}}$ (gray dotted lines in Figure 4a), in agreement with experimental findings.

Stacking destabilizes the $\pi$-HOMO (out-of-phase combination of the $\pi$-HOMO FMOs) and stabilizes the $\pi^*$-LUMO (in-phase combination of $\pi^*$-LUMO FMOs), leading to a net decrease in $\Delta E_{\text{exc}}$ (see Figure S8 in the Supporting Information). The frontier $\pi$-MO levels of benzene lose their degeneracy in the dimer to split into bonding and antibonding MOs. The extent of this splitting is governed by the degree of overlap of the $\pi$-FMOs of the benzene fragments, which is maximal for a co-facially stacked arrangement, giving a maximum decrease in $\Delta E_{\text{exc}}$.

Intermolecular distances, $d$, of 3.0–4.0 Å of $\pi$-stacked molecules are known to induce profound changes in their electronic structure. Decreasing the inter-core distance ($d = 3.79$ Å)
for the equilibrium geometry of the benzene dimer increases the through-space $\pi-\pi$ overlap, resulting in a destabilization of $\pi$-HOMO and stabilization of $\pi^*$-LUMO, and consequently decreasing the $\Delta E_{\text{H-L}}$. The inter-core distance of 3.09 Å obtained from the crystal structure of [2.2]paracyclophane\(^{43}\) is much smaller than that of the ideal $\pi$-stacking distance between aromatic rings of 3.40 Å in graphite\(^{44}\) and reflects the potential for strong overlap between the $\pi$ orbitals of the aromatic entities. We found the same effect as that of $\pi-\pi$ overlap between the $C_6H_4$ fragments, which increases significantly ($\Delta S_{\pi,\pi}$ and $\Delta S'_{\pi,\pi} \approx 280\%$; Figure 4b) if $d$ is reduced from 3.79 Å to 3.00 and 2.50 Å (Figures S8 and S9 in the Supporting Information), leading to a large splitting between the bonding and antibonding combinations (black solid lines in Figure 4a), and consequently, a decrease in $\Delta E_{\text{H-L}}$ from 4.6 eV to 3.6 and 2.2 eV.

Interplay of tuning parameters

Complex model systems

Having addressed the effect of individual structural distortions on the electronic structure of benzene, we continue by investigating their combined effect on the change in $\Delta E_{\text{H-L}}$ for a series of cyclophanes (Figure 5 and Table 1), with the aim of ra-
The isomeric [6]- and [5]metacyclophanes (5 and 6, respectively), in which the bridge connects two meta carbon centers, show a similar trend with $E_{\pi}(S_0)$ being 0.40 eV smaller for shorter bridged 6 (4.55 eV) than that for 5 (4.96 eV).[43] Moreover, $E_{\pi}(S_0)$ is systematically blue-shifted in the case of [n]metacyclophanes compared with their corresponding [n]paracyclophane counterparts. We note that the nature of the participating MOs in the transition differ because the HOMO and HOMO−1 are inverted due to a different mechanism of out-of-plane bending than that of paracyclophane (Figure S9 in the Supporting Information).[39] Introducing a second benzene ring in a sandwich-shaped motif held together by para bimethylene bridges, to give [2,2]paracyclophane (7), an $E_{\pi}(S_0)$ of 4.41 eV is obtained, that is, as expected, red-shifted by 1.77 eV from 1, but only marginally blue-shifted (0.03 eV) from that of 4. Apparently, the significant red-shift of $E_{\pi}(S_0)$, resulting from intermolecular $\pi$–$\pi$ stacking, is compensated for by the blue-shift that has its origin in the much smaller bending caused by the eight-membered bridge of 7 that than of the pentamethylene bridge of 4 ($\Delta \alpha = 12^\circ$). The strong $\pi$–$\pi$ interaction results from the short transannular distance of 3.10 Å (3.09 Å in the crystal structure),[16] which also lengthens the C–C bond of the bimethylene bridges to 1.62 Å. Replacing these bridges for methylene ones, as in [1,1]paracyclophane (8), reduces $\Delta E_{\text{HL}}$ by a significant 41%, as a result of the shorter inter-core distance and the larger $\alpha$ and $\beta$ angles. Our calculated absorption maximum of $\lambda = 419$ nm for 8 compares well with the reported value of $\lambda = 377$ nm measured at 77 K in an inert matrix.[42] Bending induced by connecting the close-lying stacked aromatic cores ($d = 3.0 \text{ Å}$) by a short para hexamethylene bridge reduces $\Delta E_{\text{HL}}$ from 3.6 (co-facially benzene dimer, see the previous section) to 2.2 eV. Such a large decrease in $\Delta E_{\text{HL}}$ reflects a (non-linear) cumulative effect of bending and inter-core distance on $\Delta E_{\text{HL}}$. However, the counterpart of 8, [1,1]metacyclophane (9) shows a rather large blue-shift of 1.24 eV from 8. Although bending is similar in both 8 and 9, the loss of $\pi$–$\pi$ stacking in the latter prohibits red-shift of the absorption relative to that of 8. Adding another methylene bridge to 8 pulls the stacked aromatic cores closer to each other ($\Delta d = 0.3 \text{ Å}$) in [1,1,1][1,4,5]cyclophane (10), leading to a marginal red-shift of about 0.20 eV of $E_{\pi}(S_0)$ from that of 8. Because the length of the bridge could not be reduced further, we decided to enhance the red-shift by strengthening $\pi$–$\pi$ stacking by adding one more methylene bridge to 10 to obtain 11 or [1,1,1][1,2,4,5]cyclophane. As predicted, compound 11 undergoes a large red-shift of 0.79 eV.
from 10. The large red-shift stems from a stronger π–π stacking due to an extremely short $d$ of 2.4 Å, which is 0.3 Å shorter than that of 10. This indicates that the inter-core distance decreases with an increase in the number of bridges. Relative to reference 1, compound 11 exhibits an extremely large red-shift in absorption of about 4.20 eV due to a combined effect of large bending due to multiple short bridges, along with a strong π–π stacking ($d=2.40$ Å versus 3.79 Å in benzene dimer), giving rise to a highly distorted system. The inherent strain in 11 is reflected in its significantly distorted benzene Figure 5. Equilibrium structures of model cyclophane systems (out-of-plane bending and bending of the bridge, $\alpha$ (in red) and $\beta$ (in green), respectively, in $^\circ$ and inter-core distance, $d$, in Å) analyzed in this study. C: gray, H: white, Si: gold, P: orange, N: blue. Only the maximum $\alpha$ and $\beta$ (see Scheme 1) and minimum $d$ (see Scheme 1) are illustrated for each molecule to ensure a consistent comparison, except 10, in which the average inter-core distance is mentioned. $\alpha$ and $\beta$ are not assigned in 10 because they are not congruent with the definitions in Scheme 1.
rings and elongated sp²–sp³ C–C bonds of 1.56 versus 1.50 Å in propene. Although the synthesis of 11 has eluded synthetic chemists due to challenges with stability, its higher homologue, (2,2.2.2)1,2,4,5)cyclophane, has been characterized by Boekelheide and co-workers. Overall, it is evident that shortening the length of the bridge and including multiple bridges enhances the bonding interaction between the carbon π and π stacking interaction to cumulatively, but non-linearly, decrease the ΔE_{HL} and consequently, red-shift E_{0}\text{LUMO}.

To enhance the red-shift of E_{0}\text{LUMO} of the cyclophanes even further, with the aim of achieving NIR absorption we introduced main-group heteroatoms into the aromatic rings. A heteroatom in a (poly)cyclic aromatic compound can serve as an auxiliary electron donor and/or acceptor and changes energy levels, shapes of the frontier orbitals, and the geometry of the aromatic core. For instance, incorporating heavier main-group elements into benzene rings has been shown to lower ΔE_{HL}. Upon replacing one of the unsubstituted aromatic carbon atoms of 11 for either a silicon (12) or a phosphorus (13) atom, ΔE_{HL} decreases due to both destabilization of the π-HOMO and stabilization of the π*-LUMO. This is consistent with studies by Wisor and Czuchajowski, who investigated pyridinophanes (Y = N) and found a small red-shift in the absorption maximum relative to that of the all-hydrocarbon analogue. The effect of heteroatom substitution is significant because E_{0}\text{LUMO} is red-shifted by 0.28 and 0.36 eV upon going from 11 to 12 and 13, respectively, and increases to 0.38 eV for the di-P-substituted derivative 14. Distortion of the hexagonal structure of 1 due to the larger Si and P atoms (Figure 6a) reduces the bonding interaction between the carbon π bond, which, in turn, leads to a destabilization of the π-HOMO adjacent to the heteroatom, which, in both of which result in a decreased ΔE_{HL}. Additionally, the out-of-plane distortion caused by the enhanced steric repulsion between the larger heteroatom and its neighboring carbon atoms (Δσ(11–13) = 4°) adds to a decrease in ΔE_{HL}.

Table 1. Orbital energy (H = HOMO, L = LUMO) and ΔE_{HL} gap, first and second lowest singlet excitation energies (E_{i}(S_1) and E_{i}(S_2)), along with the oscillator strength (f) corresponding to the S_{1}→S_{0} transition for the model cyclophane systems.

| Compound | H^{H} | L^{H} | ΔE_{HL}^{[b]} | E_{i}(S_1)^{[b]} | E_{i}(S_2)^{[b]} | E_{0}\text{LUMO} (nm)^{[c]} | f^{[d]} |
|----------|-------|-------|-------------|-----------------|-----------------|----------------------------|-----|
| 1        | −6.1  | −1.1  | 5.0         | 5.46            | 6.18 HL         | 227                       | 0.000 |
| 2        | −5.3  | −1.2  | 4.1         | 4.91            | 5.36 HL         | 235                       | 0.004 |
| 3        | −5.3  | −1.3  | 4.0         | 4.69            | 4.98 HL         | 264                       | 0.005 |
| 4        | −5.3  | −1.6  | 3.2         | 4.38            | 4.42 HL         | 283                       | 0.007 |
| 5        | −5.5  | −1.1  | 4.4         | 4.96 HL         | 5.65            | 250                       | 0.009 |
| 6        | −5.3  | −1.3  | 4.0         | 4.55 HL         | 5.34            | 272                       | 0.010 |
| 7        | −5.2  | −1.5  | 3.7         | 4.41 HL         | 4.83            | 281                       | 0.000 |
| 8        | −4.8  | −2.6  | 2.2         | 2.96 HL         | 3.37            | 419                       | 0.000 |
| 9        | −5.1  | −2.0  | 3.1         | 4.06            | 4.20 HL         | 305                       | 0.000 |
| 10       | −4.8  | −2.8  | 2.0         | 2.77 HL         | 3.39            | 448                       | 0.002 |
| 11       | −4.8  | −3.4  | 1.4         | 1.98 HL         | 2.99            | 626                       | 0.000 |
| 12       | −4.7  | −3.7  | 1.0         | 1.70 HL         | 1.85            | 729                       | 0.000 |
| 13       | −4.9  | −3.9  | 1.0         | 1.62 HL         | 2.09            | 765                       | 0.000 |
| 14       | −5.0  | −4.0  | 1.0         | 1.60 HL         | 1.75            | 775                       | 0.000 |
| 15       | −4.9  | −4.1  | 0.8         | 1.45 HL         | 1.54            | 655                       | 0.008 |

[a] Energies (in eV, unless stated otherwise). (b) Computed at the BLYP-D3(BJ)/TZ2P level. (c) Computed at the CAMB3LYP/TZ2P//BLYP-D3(BJ)/TZ2P level. See Table S2 in the Supporting Information for details on the MO composition. The excitation energy corresponding to a HOMO–LUMO transition is marked by HL.

Figure 6. a) Structural distortion (in red) upon heteroatom substitution (Si) in benzene. b) Change in π-HOMO energy due to distortion of the aromatic core (r) upon introducing Si. The blue filled circles represent the in-phase p bond of C. The energies were calculated at the BLYP-D3(BJ)/TZ2P level.
Finally, to even further reduce $\Delta E_{11i}$ we applied the push-pull concept$^{[32]}$ to di-P cyclophane 14 in the expectation of a favorable contribution from the charge-transfer excitation effect. To this end, the para position of one of the phosphabenzene rings of 14 was substituted with an electron-donating (D) amine group and the other phosphabenzene ring with an electron-withdrawing (A) cyanide group to give 15. These two substituents cause a red-shift of $E_{1}(S_0)$ by 0.15 eV, leading to a NIR absorption at 1.45 eV ($\lambda = 855$ nm, see Table 1). The decrease in $\Delta E_{11i}$ results from symmetry-allowed mixing of the high-lying amine lone pair that destabilizes the $\pi$-HOMO of 14 and the low-lying cyano $\pi^*$ orbital that stabilizes the overall $\pi^*$-LUMO. This decrease in $\Delta E_{11i}$ upon D/A substitution is in qualitative agreement with previous studies.$^{[5,53]}$ However, in all these modeled systems, the oscillator strength corresponding to the NIR absorption is negligible rendering them to be poor chromophores. Hence, in the next section, we provide a systematic and rational approach to tune the oscillator strength.

**Tuning the $E_{1}(S_0)$ oscillator strength**

The transition probability and oscillator strength ($f$) of the vertical $S_1\rightarrow S_0$ excitation must both be nonzero for light-harvesting molecules. This necessitates fine-tuning of the ground and excited states of our model cyclophanes because all have negligible oscillator strengths (Table 1) due to electric dipole-forbidden transitions, vanishing transition dipole moments, and/or cancellation of transition dipole moment vectors.$^{[54]}$ To circumvent this, we focused on modifying the nature of the MOs involved in the lowest electronic transitions by introducing aromatic substituents.

Although silacyclophane 12 and phosphacyclophane 13 are both asymmetric, their oscillator strengths are vanishingly small for the $S_1\rightarrow S_0$ excitation, but the next lowest excitation, $S_1\rightarrow S_2$, which is essentially a HOMO\(-1\)--LUMO transition, exhibits a non-negligible oscillator strength of about 0.02. Analysis of the all-carbon homologue 11 provides insight into how to move forward. HOMO of 11 has a nodal plane passing along C1 (or C1') and C4 (or C4'), leading to zero electronic coupling between the substituents at these positions (see Figure 7a for atom numbering), but this is not the case for its HOMO--1. Therefore, we aimed to invert the order of HOMO and HOMO--1 by judicious choice of substituents. Such perturbations might enable the oscillator strength to be adjusted and the light-harvesting properties altered.$^{[55]}$ Addition of a single benzene substituent at the C4 position of phosphacyclophane 13 to give 16 (Figure 7) inverts the HOMO and HOMO--1, but $f$ remains negligible for the $S_1\rightarrow S_0$ transition, which is now instead predominantly composed of the HOMO\(-1\)--LUMO transition (cf., HOMO of 11)--LUMO transition (Figure 7b) and, as expected, results in a poor $f$. This change in the nature of the $S_1\rightarrow S_0$ transition from a HOMO--LUMO transition is due to a large contribution of the coupling matrix, $K$, in describing this excitation, which becomes clear from Eqs. (3) and (4).$^{[56a,56]}$

$$\Omega_{\text{int}, j\rightarrow i} = \delta_{ij} \delta_{pq} \Delta_{\text{int}}^2 + 2 \sqrt{\Delta_{\text{int}}} K_{\text{int}, j\rightarrow i} \sqrt{\Delta_{\text{int}}} \quad (4)$$

in which $\Omega$ is the four-index matrix and the excitation energies, $\omega_{ij}$, and oscillator strength, $f$, are obtained by solving the eigenvalue in Eq. (3). $\Delta$ represents the orbital energy gap between occupied orbital $i$ and unoccupied orbital $j$ ($\epsilon_i - \epsilon_j$) and $K$ is the coupling matrix, which includes the contribution to the change in density, and thus, other linear response molecular properties if an electron is excited from an occupied to an unoccupied orbital. In a zeroth-order approximation, in which $K$ is neglected, the lowest excitation is the pure HOMO--LUMO transition. However, the inclusion of $K$, which is necessary for accurately predicting excitation properties, couple other transitions, such as HOMO--1--LUMO (in our case) with HOMO--LUMO. In 16, a large magnitude of this coupling changes the nature of the lowest excitation and this results again in a negligible $f$. A similar negligible value of $f$ is obtained for diphenyl-substituted 17. Interestingly, the second lowest excitation ($S_1\rightarrow S_2$) is close-lying (0.06 eV) relative to the first ($S_1\rightarrow S_0$), and is composed of the desired HOMO (HOMO--1 of 11)--LUMO transition, which unexpectedly results in a nonzero $f$. From our previous analysis, we showed that the large decrease of the $\Delta E_{11i}$ when $\alpha$ and $\beta = 40^\circ$ completely changed the nature of the $S_1\rightarrow S_0$ transition from being dipole-forbidden to dipole-allowed. We hypothesize that we can use this same concept and decrease the $\Delta E_{11i}$ further, which might lead to a bright lowest singlet excited state. Furthermore, we know that the orbital energy gap is the leading term in linear response TD-DFT [Eq. (4)]. So, a further decrease in the HOMO--LUMO gap might reduce the dominant effect of the coupling matrix, $K$, and revert the nature of the $S_1\rightarrow S_0$ transition to a predominant HOMO--LUMO transition, and thereby, increase the magnitude of $f$. To this end, we introduced $\pi$-conjugated fluorene groups at the C1 and C4 positions to give 18 (Figure 7c). Fluorene exhibits desirable absorption properties and has found extensive application in co-polymers to enhance their light-harvesting character.$^{[57]}$ Gratifyingly, this resulted in an intense NIR absorption at $\lambda = 785$ nm (1.58 eV, see Figure 8a) composed of the HOMO--LUMO transition, along with a large oscillator strength of 0.22. This result is consistent with the work of Wang et al., who found a red-shifted emission for fluorenedithia[3.3]paracyclophane in both solution and the solid-state, along with an increase in photoluminescence quantum yield$^{[58]}$. The large $f$ for 18 results mainly from 1) an increase in the transition dipole moment due to an increased spatial overlap between the delocalized HOMO and LUMO on the fluorene moieties (see Figure S11 in the Supporting Information), and 2) a large configuration interaction coefficient, corresponding to the HOMO--LUMO transition (0.94; Figure 8b). Introduction of the effect of solvation $(\text{CH}_2\text{Cl}_2)$ with the continuum solvation model COSMO to stabilize the charge-separated excited state shifts the longest wavelength peak of 18 even further into the NIR region to $\lambda = 821$ nm.

Additional benefits of introducing the two large $\pi$-conjugated substituents are more pronounced absorptions with non-zero intensity in the UV/Vis region (see Figure 8a). The most in-
tense one for 18 is the \( S_1 \rightarrow S_0 \) transition, which represents nearly exclusively a transition from the HOMO to the LUMO + 1 (0.93), which corresponds to the LUMO + 2 of 11 highlighted in green in Figure 7b. The strong bathochromic shift \( (\Delta E_0(S_1) = 0.55 \text{ eV}) \) of this excitation compared to that of 11 and its much higher oscillator strength \( (\Delta f = 0.68) \) stand out and suggest that the outlined strategy might be of use for the design of light absorbers with a wide spectral response.

**Conclusion**

The HOMO–LUMO energy gap of model cyclophanes can be reduced and systematically tuned by structural distortion that invokes 1) out-of-plane bending of the aromatic core; 2) bending of the bridge with respect to the aromatic core; 3) nature of the bridge substituents; and 4) \( \pi \rightarrow \pi \) stacking, of which out-of-plane bending, bending of the bridge, and \( \pi \rightarrow \pi \) stacking are most prominent. The various parameters are coupled by the length and number of bridges and have a non-linear cumulative effect on the decrease of the HOMO–LUMO energy gap, and thus, the absorption wavelength. Complex molecules in-

---

**Figure 7.** a) Atom numbering of C atoms (red filled circles) in 11. b) Five frontier orbitals (HOMO–1, HOMO, LUMO, LUMO + 1, and LUMO + 2) of 9, along with the shift in MO ordering, moving from model system 11 to 13 and 16–18, together with the \( S_1 \rightarrow S_0 \) transition [in brown, see also the text and Eqs. (3) and (4)]. c) Photophysical data of model systems 13 and 16–18 (out-of-plane bending of the aromatic core \( \alpha \) in Å and the inter-core distance in Å) computed at the CAMY-B3LYP/TZ2P//BLYP-D3(BJ)/TZ2P level.
Corporating these design parameters are modeled in a systematic and rational approach to achieve intense NIR absorption. Cyclophanes with shorter bridges are subject to more bending of the aromatic core and bridge and, when multiple π-conjugated cores are involved, a tighter π–π stacking. The effect of out-of-plane bending could be quantified by dissecting benzene into two allylic fragments. This revealed an antibonding contribution to the π-HOMO (and thus destabilization) with increasing out-of-phase overlap of the allylic π-HOMOs and stabilization of the π*-LUMO because the fragment π*-LUMOs penetrate favorably into each other. Bending the bridge amplifies the decrease of the HOMO–LUMO gap by mixing the orthogonal σ- and π-FMOs of the bridge and aromatic core, respectively. However, the largest change in the MO energies originates from π–π stacking because decreasing the intercore distance by reducing the bridge length or increasing the number of bridges enables significant through-space π–π orbital overlap. Inclusion of a heavy main-group heteroatom, such as silicon or phosphorus, and adding a push–pull effect to a heavily distorted cyclophane augmented the decrease of the HOMO–LUMO gap and strongly red-shifted the HOMO–LUMO excitation relative to that of benzene.

These insights, along with a rational approach to tune the oscillator strength, were applied in a proof-of-concept design that led to distortion-controlled cyclophane-like model dye molecule 18, which had a bright TD-DFT-calculated absorption at \( \lambda = 785 \) nm and at \( \lambda = 812 \) nm if the effect of CH$_2$Cl$_2$ (DCM) as the solvent was included. These absorptions are well into the NIR range and reflect a large redshift of about 600 nm from the HOMO–LUMO excitation of reference benzene. The contribution of the two fluorene groups in 18 is crucial because they cause the needed switching of the highest occupied MOs to convert the dark state into a bright excited state. This finding establishes distortion-control as an alternative design principle for tuning the optoelectronic structure of organic dye molecules, especially for, but not limited to, red-shifting absorption into the NIR, with possible applications ranging from bent organic semiconductors to molecular switches.

Acknowledgements

This work is part of the Industrial Partnership Program (IPP) “Computational sciences for energy research” (project 14CSER011), which is part of the Netherlands Organization for Scientific Research (NWO) and co-financed by Shell Global Solutions International B.V. J.P. thanks the Spanish MINECO (CTQ2016-77558-R and MDM-2017-0767) and the Catalan Government (2017SGR348). A.K.N. thanks Souloke Sen for insightful discussions on TD-DFT.

Conflict of interest

The authors declare no conflict of interest.

Keywords: cyclophanes · density functional calculations · dyes/pigments · HOMO–LUMO gap · near-infrared absorptions

[1] G. Qian, Z. Y. Wang, Chem. Asian J. 2010, 5, 1006.
[2] a) A. Burke, L. Schmidt-Mende, S. Ito, M. Grätzel, Chem. Commun. 2007, 234; b) P. Y. Reddy, L. Giribabu, C. Lyness, H. J. Snith, C. Vijaykumar, M. Chandrasekharan, M. Lakshminarayanan, J.-H. Yum, K. Kalyanasundaram, M. Grätzel, M. K. Nazeeruddin, Angew. Chem. Int. Ed. 2007, 46, 373; Angew. Chem. 2007, 119, 377.
[3] G. Hong, A. L. Antaris, H. Dai, Nat. Biomed. Eng. 2017, 1, 0010.
[4] a) E. L. Williams, J. Li, G. E. Jabbour, Appl. Phys. Lett. 2006, 89, 083506; b) C. Borek, K. Hanson, P. I. Djurovich, M. E. Thompson, K. Aznavour, R.
due to structural distortion, which
Inorg. Chem. 46, 119.
J. Chem. Phys. 120, 119.

[20] E. R. Darzi, R. Jasti, A. K. Narsaria, T. A. Hamlin, K. Lammertsma, F. M. Bickelhaupt, J. Am. Chem. Soc. 2011, 133, 1253.

[21] a) G. te Velde, F. M. Bickelhaupt, E. J. Baerends, C. Fonseca Guerra, S. J. A. van Gisbergen, J. G. Snijders, E. van Lenthe, J. A. Groeneveld, J. G. Süsser, J. Chem. Soc. Rev. 1999, 110, 5029.

[22] E. van Lenthe, E. J. Baerends, J. G. Süsser, J. Am. Chem. Soc. 1999, 121, 10356.

[23] a) F. M. Bickelhaupt, E. J. Baerends in Reviews in Computational Chemistry (Eds.: K. B. Lipkowitz, D. B. Boyd), Wiley, Hoboken, 2000, pp. 1–86; b) T. Ziegler, A. Rauk, Inorg. Chem. 1979, 18, 1755.

[24] C. Fonseca Guerra, J. W. Hardgraaf, E. J. Baerends, F. M. Bickelhaupt, J. Comput. Chem. 2004, 25, 189.

[25] a) S. J. A. van Gisbergen, J. G. Süsser, E. J. Baerends, Comput. Phys. Commun. 1999, 118, 119; b) A. Rosa, E. J. Baerends, S. J. A. van Gisbergen, E. van Lenthe, J. A. Groeneveld, J. G. Süsser, J. Am. Chem. Soc. 1999, 121, 10356.

[26] T. B. Demisiss, H. Dodziuk, J. Waluk, R. Kudr, M. Pietrzak, V. Vetokhina, S. Szymanski, J. Žažiwek, H. Hopf, J. Phys. Chem. A 2016, 120, 724.

[27] a) R. Grimm, M. Parac, PhysChemPhys 2003, 4, 292; b) R. M. Richard, J. M. Herbert, J. Chem. Theory Comput. 2011, 7, 1296.

[28] a) Z. Chen, M. Giorgi, D. Jacquemin, M. Elhabiri, O. Siri, J. Chem. Phys. 2004, 120, 17597.

[29] a) E. R. Darzi, R. Jasti, A. K. Narsaria, T. A. Hamlin, K. Lammertsma, F. M. Bickelhaupt, J. Am. Chem. Soc. 2011, 133, 1253.

[30] a) G. te Velde, F. M. Bickelhaupt, E. J. Baerends, C. Fonseca Guerra, S. J. A. van Gisbergen, J. G. Snijders, E. van Lenthe, J. A. Groeneveld, J. G. Süsser, J. Chem. Soc. Rev. 1999, 110, 5029.

[31] a) Y. Zhao, D. G. Trulhar, J. Chem. Phys. 2006, 125, 194101; b) Y. Zhao, D. G. Trulhar, Theor. Chem. Acc. 2008, 120, 215.

[32] a) R. Grimm, J. Comput. Chem. 2004, 25, 1463; b) M. Enzerhof, G. E. Scuseria, J. Chem. Phys. 1999, 110, 5029.

[33] a) Z. Chen, M. Giorgi, D. Jacquemin, M. Elhabiri, O. Siri, J. Chem. Phys. 2004, 120, 17597.

[34] a) E. R. Darzi, R. Jasti, A. K. Narsaria, T. A. Hamlin, K. Lammertsma, F. M. Bickelhaupt, J. Am. Chem. Soc. 2011, 133, 1253.

[35] a) G. te Velde, F. M. Bickelhaupt, E. J. Baerends, C. Fonseca Guerra, S. J. A. van Gisbergen, J. G. Snijders, E. van Lenthe, J. A. Groeneveld, J. G. Süsser, J. Chem. Soc. Rev. 1999, 110, 5029.
Ishibashi, A. Dargelos, C. Darrigan, A. Chrostowska, S.-Y. Liu, Organometallics 2017, 36, 2494; e) Z. Liu, J. S. A. Ishibashi, C. Darrigan, A. Dargelos, A. Chrostowska, B. Li, M. Vasiliu, D. A. Dixon, S.-Y. Liu, J. Am. Chem. Soc. 2017, 139, 6082; f) J. Böhne, H. Braunschweig, J. C. Jiménez-Halla, I. Krummenacher, T. E. Stennett, J. Am. Chem. Soc. 2018, 140, 848.

[50] a) P. D. Burrow, A. J. Ashe, D. J. Bellville, K. D. Jordan, J. Am. Chem. Soc. 1982, 104, 425; b) N. Nakata, N. Takeda, N. Tokitoh, J. Am. Chem. Soc. 2002, 124, 6914; c) T. Ishii, K. Suzuki, T. Nakamura, M. Yamashita, J. Am. Chem. Soc. 2016, 138, 12787; d) N. Tokitoh, K. Wakita, R. Okazaki, S. Nagase, P. von Ragüé Schleyer, H. Jiao, J. Am. Chem. Soc. 1997, 119, 6951; e) S. M. Draper, D. J. Gregg, R. Madathil, J. Am. Chem. Soc. 2002, 124, 3486; f) X. Wang, F. Zhang, K. S. Schellhammer, P. Machata, F. Ortmann, G. Cuniberti, Y. Fu, J. Hunger, R. Tang, A. A. Popov, R. Berger, K. Müllen, X. Feng, J. Am. Chem. Soc. 2016, 138, 11606.

[51] a) A. K. Wisor, L. Czuchajowski, J. Phys. Chem. 1986, 90, 1541. 

[52] a) G. L. Eakins, J. S. Alford, B. J. Tiegs, B. E. Breyfogle, C. J. Stearman, J. Phys. Org. Chem. 2011, 24, 1119; b) F. Burel, RSC Adv. 2014, 4, 58826; c) C. Duan, F. Huang, Y. Cao, J. Mater. Chem. 2012, 22, 10416.

[53] a) J. R. Mulder, C. Fonseca Guerra, J. C. Sloomweg, K. Lammertsma, F. M. Bickelhaupt, J. Comput. Chem. 2016, 37, 304; b) M. Ottonelli, M. Piccardo, D. Duce, S. Thea, G. Delegianni, J. Phys. Chem. A 2012, 116, 611; c) L. Ji, A. Lorbach, R. M. Edkins, T. B. Marder, J. Org. Chem. 2015, 80, 5658.

[54] E. B. Guidez, C. M. Aikens, J. Phys. Chem. C 2013, 117, 21466.

[55] a) A. G. Crawford, A. D. Dewyer, Z. Liu, A. Steffen, A. Beeby, L.-O. Pålsson, D. J. Tozer, T. B. Marder, J. Am. Chem. Soc. 2011, 133, 13349; b) L. Ji, R. M. Edkins, A. Lorbach, I. Krummenacher, C. Brückner, A. Eichhorn, H. Braunschweig, B. Engels, P. J. Low, T. B. Marder, J. Am. Chem. Soc. 2015, 137, 6750; c) I. Merz, J. Fink, A. Friedrich, I. Krummenacher, H. A. Mamari, S. Lorenzen, M. Haehnel, A. Eichhorn, M. Moos, M. Holzapfel, H. Braunschweig, C. Lambert, A. Steffen, L. Ji, T. B. Marder, Chem. Eur. J. 2017, 23, 13164.

[56] M. A. L. Marques, N. T. Maitra, F. M. S. Nogueira, E. K. U. Gross, A. Rubio, Fundamentals of Time-Dependent Density Functional Theory, Springer, Heidelberg, 1996.

[57] a) P. Najchalski, Y. Morel, O. Stéphan, P. L. Baldeck, Chem. Phys. Lett. 2001, 343, 44; b) D. C. Watters, Y. Hunan, A. J. Pearson, K. Amid, L. David, Macromol. Rapid Commun. 2013, 34, 1157; c) F. Huang, K.-S. Chen, H.-L. Yip, S. K. Hau, O. Acton, Y. Zhang, J. Luo, A. K. Y. Jen, J. Am. Chem. Soc. 2009, 131, 13886.

[58] a) W.-L. Wang, J. Xu, Z. Sun, X. Zhang, Y. Lu, Y.-H. Lai, Macromolecules 2006, 39, 2777; b) W. Wang, J. Xu, Y.-H. Lai, Org. Lett. 2003, 5, 2765.

[59] a) M. Ball, B. Zhang, Y. Zhong, B. Fowler, S. Xiao, F. Ng, M. Steigerwald, C. Nuckolls, Acc. Chem. Res. 2019, 52, 1068; b) B. Zhang, M. T. Trinh, B. Fowler, M. Ball, Q. Xu, F. Ng, M. L. Steigerwald, X. Y. Zhu, C. Nuckolls, Y. Zhong, J. Am. Chem. Soc. 2016, 138, 16426.

[60] D. Bléger, S. Hecht, Angew. Chem. Int. Ed. 2015, 54, 11338; Angew. Chem. 2015, 127, 11494.