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Absorption and Fluorescence Signatures of 1,2,3-Triazole Based Regioisomers: Challenging Compounds for TD-DFT †

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In the continuous quest for improving TD-DFT methodologies as a tool to predict the photophysical features of solvated chromophores, we herein investigate two model regioisomers based on the 1,2,3-triazole moiety. Starting from their experimental absorption and emission spectra, key energy differences highlighting the main trends between the two isomers are extracted and used to gauge the accuracy of several levels of theory. RI-CC2 and EOM-CCSD calculations allow us to ascertain that the low energy spectra is not linked to double excitations. In vacuum, neither these methods nor any of the implemented TD-DFT levels of theory, ranging from global hybrids (PBE0, B3LYP) to range-separated functionals without (CAM-B3LYP, ωB97X) or with dispersion corrections (ωB97X-D), is able to capture the key features that differentiate the two chromophores. Accounting for solvent within a specific PCM model allows to recover experimental trends, but the dramatic changes occurring when moving from toluene to THF and/or for different PCM approaches (LR, cLR, SS) suggest that this agreement is probably fortuitous. Even so the ωB97X-D functional combined to the SS-PCM scheme leads to quantitative agreement with experiment, TD-DFT results obtained for 1,2,3-triazole based chromophores need to be treated with caution. We also show that the SS-PCM approach may be useful to test current and novel exchange-correlation functionals against the charge transfer failure.

1 Introduction

The computation of accurate excitation energies of medium and large organic molecules has been at the centre of continuous attention from the Time Dependent Density Functional Theory (TD-DFT) community.1–3 In the past decade, TD-DFT has shown to be an economical yet pretty accurate tool for simulating the excited-states (ES) of chromophores. Besides the widespread applications afforded by organic and inorganic luminophores, we underline that three methodological breakthroughs have clearly boosted TD-DFT’s popularity. First, the development and implementation of new exchange-correlation functionals,4–7 dispersion corrected functionals,8–11 and double hybrid functionals12–14 allowed to overcome well-known pitfalls of conventional DFT kernels that is the appearance of spurious charge transfer states,15–17 the inability to describe weak-interactions and the simulation of doubly excited-states,18 respectively. The possibility to compute vibrationally resolved optical spectra19–23 stands as a second important progress as it allows, at least, to capture the band asymmetries if the underlying multi-peak structure of the gas phase spectra becomes smeared out by inhomogeneous broadening in solution. As steady-state optical spectroscopy in solution is routinely used to characterise the actual dipole of the excited-state. This can be done ei-
ther within a perturbative scheme, namely the corrected LR scheme (cLR),\textsuperscript{26,28} or self-consistently using the state-specific scheme (SS)\textsuperscript{27,29} or the vertical excitation model (VEM).\textsuperscript{30–32} Despite these developments, including large benchmarking, there is still need for further investigations in the quest of designing theoretical schemes able to accurately predict spectroscopic data as some unexpected failures of TD-DFT can still be found, and we tackle such problem in the present contribution. Among the most simple test examples one can imagine, regio-isomers (position isomers) of chromophores are good candidates for TD-DFT assessment due to their chemical similarity. 1,2,3-triazole based regio-isomers have recently been suggested as a kind of tutorial example to test novel exchange-correlation kernels.\textsuperscript{33} Indeed, while the absorption features of one of the isomers showed to be qualitatively well described when using the B3LYP global hybrid functional,\textsuperscript{34,35} the same level of theory was deficient for its isomeric counterpart.\textsuperscript{33} At the same time, triazole chemistry has gained renewed interest as it allows for click chemistry reactions, making triazole-based molecules interesting beyond an academic viewpoint.\textsuperscript{36} Indeed, with four attachment points, 1,2,3-triazole paves the way to different regio-isomers of various sizes and shapes, ranging from small push-pull chromophores\textsuperscript{33,37–43} to polymers and branched molecular architectures.\textsuperscript{33,41–48} These molecular compounds have shown diverse optical responses. For example, position isomerism has been demonstrated to dramatically change absorption features of push-pull chromophores\textsuperscript{38,39} and ligation of triazole on coumarins has been reported to increase their fluorescence quantum yield by more than an order of magnitude.\textsuperscript{37} Triazole based multipolar structures have also been reported to develop large nonlinear optical responses, in particular for electro-optic activity and two-photon absorption.\textsuperscript{33,42–48} Moreover, triazole is recognised as a versatile chemical moiety, able to act either as an electron donor\textsuperscript{37} or acceptor,\textsuperscript{36,42,44} as well as an active \(\pi\)-linker\textsuperscript{38,39} or as a site-isolation moiety,\textsuperscript{45–47} while affording metal ligation in some cases.\textsuperscript{40} This prompts the need for theoretical methodologies able to provide an accurate description of their ES structure or, at least, chemically sounded.

In that framework, we investigate here the photophysical properties of two model regio-isomers based on the 1,2,3-triazole moiety (Figure 1), labelled \(\text{da}\) and \(\text{dai}\). The motivation for choosing these two specific regio-isomers are two-fold. First, comparison of experimental photophysical data and TD-B3LYP results of multi-branched analogues of these two specific chromophores (e.g., quadrupoles \(\text{2da}\) and \(\text{2dai}\) obtained by gathering respectively two \(\text{da}\) and \(\text{dai}\) units in a head-to-tail architecture) was already available. It revealed that TD-B3LYP results fit nicely for \(\text{dai}\) analogues,\textsuperscript{43} but fail for \(\text{da}\) analogues,\textsuperscript{33} with large overestimation of both the oscillator strength ratio and the band splitting between the two-first absorption bands. Next, working with dipoles instead of quadrupoles, or even larger multi-branched chromophores, reduces considerably the computational cost, thus allowing implementation of higher levels of theory. To ensure relevant comparison with experimental data, the chromophores have been synthesised, crystallised and their fluorescence and absorption spectra were measured in both dichloromethane (DCM) and tetrahydrofuran (THF). From the theoretical point of view, we used several quantum chemical approaches. Among those, \textit{ab initio} RI-CC2 and EOM-CCSD allowed to assess the importance of double excitations in the low-energy spectra. TD-DFT calculations are performed with different approximations, first in vacuum, next for solvated chromophores using the LR-, cLR- and SS-PCM methods (see above) combined with a panel of exchange-correlation functionals including both global (B3LYP,\textsuperscript{35} and PBE\textsuperscript{0,80}) and range-separated (LC-BLYP,\textsuperscript{3} CAM-B3LYP,\textsuperscript{6} \(\omega\)B97X-D\textsuperscript{9}) hybrids. After the Section devoted to computational and experimental details, we discuss molecular geometries. Next, we present the experimental phosphophysical properties and extract relevant information for comparisons with theoretical results. Then, we successively inspect the ES properties related to absorption, emission, 0-0 energy and Stokes shift.

2 Methodology

2.1 Computations

We employed a blend of \textit{ab initio} approaches to model the two chromophores of interest. A locally modified version of GAMESS was used to carry out TD-DFT calculations with both the LC-BLYP,\textsuperscript{4} (\(\omega = 0.3\)) and B3LYP,\textsuperscript{35} functionals using the same convergence criteria described in a previous study.\textsuperscript{51,52} The TURBOMOLE code was used to carry out wave-function based RI-CC2 calculations. These gas phase calculations relied on the SVP basis set and used geometries optimised in an earlier work at the HF/6-31G level.\textsuperscript{33} All other computations were performed with the latest version of the Gaussian09 package (D.01 revision),\textsuperscript{53} using default algorithms except when noted. Ground and excited state geometries, subsequent excited state structures were computed with

![Fig. 1 Molecular structures of 1,2,3-triazole based model chromophores: \(\text{da}\) (left) and the corresponding \(\text{dai}\) position isomer (right).](Image)
different functionals: PBE0\textsuperscript{49,50}, CAM-B3LYP\textsuperscript{6}, ωB97X\textsuperscript{7} and ωB97X-D\textsuperscript{9}, employing the 6-31+G(d) atomic basis\textsuperscript{54,55}. Additional EOM-CCSD calculations were performed using the 6-31G(d) atomic basis in vacuum. All geometries reported here correspond to true minima of the potential energy surface with absence of imaginary frequencies and correspond to a so-called tight optimisation with residual mean forces smaller than 0.00001 a.u.

Solvent effects were modeled thanks to the PCM approach\textsuperscript{25} as it provides an accurate description of solvation effects as long as no specific interactions come into play. The three different PCM approaches implemented in Gaussian09 have been applied, namely LR, cLR and SS schemes (see Introduction). Within the LR model the PCM reaction field has a contribution which is related to the ground to excited state transition density rather than to the change in electron density upon excitation. This can be corrected with more advanced schemes either pertubatively (cLR) or self-consistently (SS), but at the prize of an increased computational effort, especially in the SS model. As stated in the introduction, slow (fast) processes should be described in an equilibrium (non-equilibrium) limit irrespective of the selection of LR, cLR or SS schemes. Therefore, geometry optimisations and adiabatic transition energies correspond to the equilibrium regime, whereas vertical transition energies do not, as they require the equilibrium limit for the initial state and the non-equilibrium one for the final state. Furthermore, to accurately compute the crossing point of fluorescence and absorption bands, known as the experimental 0-0 reference ($E_00$), the equilibrium adiabatic energies should be corrected for both zero-point vibrational energy (ZPVE) and non-equilibrium effects. We redirect the interested reader to Ref. 56 for details of the selected approach than 0.00001 a.u.

Table 1 Selected structural data of da and dai: the rings are labeled A, B and C from left to right. Distances are given in Å and angles in degrees. All calculations in gas phase with 6-31+G(d) but for HF (6-31G). Additional data is available as ESI.

<table>
<thead>
<tr>
<th></th>
<th>X-Ray</th>
<th>HF</th>
<th>PBE0</th>
<th>CAM-B3LYP</th>
<th>ωB97X</th>
<th>ωB97X-D</th>
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<td>1.281</td>
<td>1.292</td>
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<td>1.293</td>
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<tr>
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<td>1.416</td>
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<td>B-C</td>
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<td>1.466</td>
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<tr>
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<td>31</td>
<td>26</td>
<td>29</td>
<td>30</td>
<td>31</td>
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<tr>
<td>$\beta$C</td>
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<td>2</td>
<td>4</td>
<td>3</td>
<td>8</td>
<td>11</td>
</tr>
</tbody>
</table>

2.3 Crystal structures

Both da and dai crystals: (C$_{16}$H$_{16}$N$_4$): M = 264.33. APEXII, Bruker-AXS diffractometer, Mo-K$_\alpha$ radiation ($\lambda = 0.71073$ Å), T = 150(2) K; da is triclinic P-1 (I.T.#2), a = 5.8977(6), b = 7.4177(7), c = 30.865(3) Å, $\alpha = 91.949(4)$, $\beta = 92.578(5)$, $\gamma = 94.343(4)^\circ$, V = 1344.1(2) Å$^3$, Z = 4, $\mu = 1.306$ g·cm$^{-3}$, $\mu = 0.081$ mm$^{-1}$; dai is monoclinic P$2_1$/n (I.T.#14), a = 5.7857(5), b = 16.921(2), c = 13.7060(17), $\beta = 98.453(4)^\circ$, V = 1327.2(3) 3, Z = 4, $\mu = 1.323$ g·cm$^{-3}$, $\mu = 0.082$ mm$^{-1}$. The structures were solved by direct methods using the SIR97 program, and then refined with full-matrix least-square methods based on $F^2$ (SHELXL-97)\textsuperscript{59} with the aid of the WINGX program. All non-hydrogen atoms were refined with anisotropic atomic displacement parameters. H atoms were finally included in their calculated positions. A final refinement on $F^2$ with 6089 unique intensities and 365 parameters converged at $\omega R(F^2) = 0.139$ (R(F) = 0.0589) for 4195 observed reflections with $I > 2\sigma(I)$ for the da crystal. For dai, refinement on $F^2$ with 3038 unique intensities and 183 parameters converged at $\omega R(F^2) = 0.1065$ (R(F) = 0.0444) for 2515 observed reflections with $I > 2\sigma(I)$.

2.4 Optical measurements

UV/Vis/IR absorption spectra were recorded on a Jasco V-570 instrument. Steady-state fluorescence spectra were recorded on an Edinburgh FL920 instrument. Measurements were performed at room temperature using freshly-prepared air-equilibrated dilute solutions (ca. $10^{-6}$ M, optical density < 0.1) contained in standard 1 cm quartz cuvettes. Emission spectra were obtained by exciting samples at the wavelength of the absorption maximum. Fluorescence quantum yields were measured according to literature procedures using fluorescein in 0.1 N NaOH as a standard (quantum yield

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\( \phi = 0.90 \).

3 Results and discussion

3.1 Ground and excited state molecular geometries

The ground state geometries obtained from X-Ray diffraction and calculated optimized geometries lead to non-planar molecular conformations for both regio-isomers (Table 1). Experimental conformations show sizeable dihedral angles on both sides of the triazole ring (Figure 2). The DFT structures are in overall good agreement with experiment, though one notices a systematically too small diazo distance (ca. 0.02 Å), a previously reported trend.\(^{61}\) The length of the interring bonds is estimated accurately by all theoretical levels. The dihedral angles obtained after geometry optimisation more significantly depend on the level of theory in use but test calculations revealed that this has an only small impact on the estimated optical properties (vide infra). They are systematically larger for the ring connected to the nitrogen atom of the triazole ring. Given the fact that solid-state packing effects might significantly tune the dihedral angles, it is not straightforward to pinpoint a preferred level of theory from the data listed in Table 1. We have also investigated solvent effects (Table 2), considering only the \( \omega B97X-D \) functional. By comparing the gas phase and THF data for the ground-state structure, one notices that solvation induces a small elongation of the diazo bonds, and slight increase of the dihedral angles. Let us now turn towards the first excited-state. While, ground state of both regio-isomers showed benzene rings with a clear aromatic character and a slightly quinoidal nature for rings A and C (Figure 1), the quinoidal character increases in the excited-states and becomes very significant for ring C. Concomitant decreases of inter-ring bond lengths, with a larger effect for the one involving the nitrogen atom of the triazole ring, and planarisation of the geometries are observed when moving to the electronically excited-state. Such a phenomenon is quite common for chromophores having low dipolar character in their ground state.

![ORTEP representation of the crystal structures of da (two molecules on top) and dai (molecule at bottom).](image)

| Table 2 Comparison between ground and (first bright) excited states \( \omega B97X-D/6-31+G(d) \) geometrical parameters computed in THF with the PCM model. See Table 1 for more details. |
|---|---|---|---|---|---|
| | GS | ES | GS | ES |
| N\(_{\text{a}}\) | 1.391 | 1.34 | 1.391 | 1.34 |
| A-B | 1.426 | 1.379 | 1.471 | 1.46 |
| B-C | 1.466 | 1.409 | 1.425 | 1.34 |
| \( \bar{\delta} \) | 16 | 1 | 45 | 0 |

3.2 Experimental optical spectra

UV/Vis and fluorescence spectrum of da and dai measured in both THF and DCM are shown Figure 3. Consistently with earlier findings on related chromophores,\(^{41,42}\) both regio-isomers have an intense absorption band in the near UV region and display a strong photoluminescence in the violet part of the visible spectrum. In THF, the absorption extinction coefficients attain 27000 and 21000 cm\(^{-1}\).mol\(^{-1}\).L whereas the fluorescence quantum yields amount to 0.87 and 0.18, for da and dai, respectively. da absorption and emission spectra are clearly not mirror images of each other (See in the ESI). This hints that at least one low-lying ES with significantly smaller oscillator strength than the higher lying state at the origin of the main absorption band is present. A crude deconvolution leads to an energy splitting between these two states of ca. 0.3–0.5 eV range (See in the ESI).

![Experimental photophysical data of the two dyes in eV.](image)

| Table 3 Experimental photophysical data of the two dyes in eV. \( \omega B97X-D/6-31+G(d) \) |
|---|---|---|---|
| | DCM | THF |
| \( \omega_{\text{abs}}^{\text{max}} \) | 4.18 | 4.22 |
| FWHM | 0.7 | 0.7 |
| \( \Delta \omega_{\text{abs}} \) | 0.3 | 0.4 |
| \( \omega_{\text{em}}^{\text{max}} \) | 2.81 | 2.78 |
| FWHM | 0.52 | 0.71 |
| \( E_{00} \) | 3.30 | 3.30 |
| \( \Delta \omega \) | 1.37 | 1.39 |

On the contrary, absorption and emission of dai are close to be mirror images. In this case, would the first ES be a (nearly) dark state, the splitting with the bright state would be much smaller than in da. Both regio-isomers have higher lying ES
Table 4 Calculated photophysical data (eV) in gas phase. Excited state number is given in ket notation and $\Delta \omega^{\text{abs}}$ and $\delta$ have been defined in the caption of Table 3. In the rightmost column, $\delta(\omega_{\text{max}}^{\text{abs}})$ and $\delta(\Delta \omega^{\text{abs}})$ are given for each method. $\delta(\omega_{\text{max}}^{\text{abs}})$ are derived from convoluted gaussians with experimental FWHM (0.5 eV) when the two bright states relevant for absorption are close in energy. Relevant experimental values in THF are also shown to facilitate comparison.

<table>
<thead>
<tr>
<th>Method</th>
<th>$\omega$ (eV)</th>
<th>$\Delta \omega^{\text{abs}}$</th>
<th>$\delta(\omega_{\text{max}}^{\text{abs}})$</th>
<th>$\delta(\Delta \omega^{\text{abs}})$</th>
</tr>
</thead>
<tbody>
<tr>
<td>EOM-CCSD/6-31G(d)$^a$</td>
<td>1</td>
<td>4.67 (0.07)</td>
<td>0.36</td>
<td>-0.19</td>
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<tr>
<td>RI-CC2/SVP$^a$</td>
<td>1</td>
<td>4.43 (0.21)</td>
<td>0.27</td>
<td>-0.27</td>
</tr>
<tr>
<td>LC-BLYP/SVP$^b$</td>
<td>1</td>
<td>4.59 (0.23)</td>
<td>0.23</td>
<td>-0.23</td>
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<tr>
<td>oB97X-D/SVP$^b$</td>
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<td>4.56 (0.36)</td>
<td>0.23</td>
<td>-0.24</td>
</tr>
<tr>
<td>oB97X-D/6-31G(d)$^b$</td>
<td>1</td>
<td>4.64 (0.33)</td>
<td>0.21</td>
<td>-0.22</td>
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<tr>
<td>oB97X-D/6-31+G(d)$^b$</td>
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<td>4.45 (0.33)</td>
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<td>-0.17</td>
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<tr>
<td>CAM-B3LYP/6-31G+G(d)$^e$</td>
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<td>4.35 (0.38)</td>
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<td>-0.18</td>
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<tr>
<td>B3LYP/SVP$^a$</td>
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<td>4.35 (0.38)</td>
<td>0.33</td>
<td>-0.18</td>
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<tr>
<td>PBE0/6-31+G(d)$^b$</td>
<td>1</td>
<td>3.71 (0.19)</td>
<td>0.86</td>
<td>-0.12</td>
</tr>
</tbody>
</table>

$^a$CAM-B3LYP geometry
$^b$HF geometry
$^c$oB97X-D geometry

Fig. 3 Normalised fluorescence (left) and absorption (right) spectra of da (top) dai (bottom) in DCM (straight lines) and THF (dotted lines).

with significant oscillator strength leading to an increase of the absorption intensity near 5.1 eV. Unfortunately, the existence of several ES with significant oscillator strength hinders straightforward deconvolution of the spectra (see in the ESI). Nevertheless, to highlight the main trends, different energies and energy differences ($\delta$, corresponding to energy shifts between da and dai) have been extracted from the spectra and are summarised Table 3. $\Delta \omega^{\text{abs}}$ corresponds to an estimate of the energy shift between the first bright excited state (ES$_{\text{max}}$) and the first excited state (ES$_1$). These energy shifts reveal clear trends between the two isomers. Indeed, both $\omega_{\text{max}}^{\text{abs}}$ and $\delta(\Delta \omega^{\text{abs}})$ are positive with a larger value for the latter, evidencing a clear separation between the first and the bright ES of da. On the contrary, $\delta(\omega_{\text{max}}^{\text{abs}})$ are negative indicating a larger stabilisation of the relaxed ES in da than in dai for both solvents. This also shows up through larger Stokes shifts for the former dye and consequently positive values for $\delta(\Delta \omega_{\text{SS}})$. These different behaviours in absorption and emission also yield redshifted crossing points ($E_{\text{SS}}$) for da and therefore negative $\delta(E_{00})$.

3.3 Modelled absorption

In order to make a direct comparison with gas phase TD-DFT results, we have carried out simulations with both RI-CC2 and EOM-CCSD wavefunction approaches (see Table 4). We underline that these methods indicate that none of the four first excited-states of both dyes presents a significant weight for double excitations. Overall, the TD-DFT and wavefunction results provide the same general qualitative trends, though the TD results are, of course, sensitive to the selected exchange-correlation functional. The agreement with the experimental data reported in Table 3 looks reasonable for da: the splitting between the first and bright state, $\Delta \omega^{\text{abs}}$, is well repro-
Table 5 Natural transition orbitals of da and dai calculated in gas phase highlighting the appearance of low-lying charge-transfer states for da when using global hybrids (PBE0 for instance). Panels quote excited state number, associated eigenvalue, transition energy and oscillator strength. Geometries are optimized at the CAM-B3LYP/6-31+G(d) level and TD calculations use the 6-31+G(d) basis set. Orbitals were plotted using XCrySDen.

<table>
<thead>
<tr>
<th></th>
<th>hole</th>
<th>electron</th>
<th>da</th>
<th>electron</th>
<th>dai</th>
</tr>
</thead>
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<tr>
<td>1</td>
<td>w=0.99, ω01 = 3.71, f01 = 0.19</td>
<td>w=0.98, ω01 = 4.30, f01 = 0.62</td>
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<td>w=0.99, ω02 = 4.26, f02 = 0.04</td>
<td>w=0.93, ω02 = 4.41, f02 = 0.09</td>
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<tr>
<td>1</td>
<td>w=0.95, ω01 = 4.35, f01 = 0.38</td>
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<tr>
<td>CAM-B3LYP</td>
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<td></td>
<td>2</td>
<td>w=0.90, ω02 = 4.68, f02 = 0.41</td>
<td>w=0.89, ω02 = 4.72, f02 = 0.85</td>
</tr>
</tbody>
</table>

Transition energies and corresponding oscillator strengths relevant for absorption (vertical approximation) have been calculated within the non-equilibrium regime and are reported Table 6 for four different functionals and three different solvents. Interestingly, the data obtained for dai are quite insensitive to the solvent (toluene, THF or DCM) and this holds for the three PCM model (LR, cLR or SS). This vastly contrasts with da for which large variations of the relative oscillator strengths of the two first excited states take place, especially with PBE0. As the two states are quite well separated with PBE0 (at least with LR and cLR), this result is puzzling as normally intensity borrowing takes only place between states that are closer on the energetic scale. This leads to drastic solvent effects with PBE0, e.g., the LR-PBE0 Δωabs goes from 0.00 to 0.62 when shifting from toluene to THF, and this consequently reflects in the sign change of δ(ωmax) characterizing the two systems. Whist, this brings the LR-PCM PBE0 results in quite good agreement with experimental absorption spectra in THF, such radical variations are probably calculation artifacts. Impressively, while the cLR-LR difference are already sizable with PBE0, the SS-LR shift appears unrealistically large. Indeed, the cLR correction to the transition energy of the first ES of da amounts to more then 0.3 eV whereas the SS corrections lead to huge shifts and dramatic decrease of oscillator strengths for both compounds. This reveals appearance of low lying spurious charge transfer states and makes the
Table 6 Calculated photophysical data using the continuum PCM model in its non-equilibrium limits. All data used the 6-31+G(d) atomic basis set and the geometries are optimized at the same level of theory as the one used for the TD-DFT part. See caption of Table 4 for more details. Relevant experimental values in THF are also shown to facilitate comparison.

<table>
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<th>Functional (medium)</th>
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<td></td>
<td></td>
<td>da</td>
<td></td>
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<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>( \omega_1 (f_1) )</td>
<td>( \omega_2 (f_2) )</td>
<td>( \Delta \omega_{abs} )</td>
<td>da</td>
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<tr>
<td></td>
<td></td>
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<td>4.01 (0.75)</td>
<td>0.00</td>
<td>( \omega_1 (f_1) )</td>
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<td>2.44 (0.10)</td>
<td>0.07</td>
<td>2.72 (0.19)</td>
</tr>
<tr>
<td>PBE0 (THF)</td>
<td>LR</td>
<td>3.67 (0.17)</td>
<td>4.29 (0.42)</td>
<td>0.62</td>
<td>4.07 (0.70)</td>
</tr>
<tr>
<td></td>
<td>cLR</td>
<td>3.34 (0.17)</td>
<td>4.19 (0.42)</td>
<td>0.85</td>
<td>3.91 (0.70)</td>
</tr>
<tr>
<td></td>
<td>SS</td>
<td>2.59 (0.08)</td>
<td>2.68 (0.01)</td>
<td>0.00</td>
<td>2.64 (0.16)</td>
</tr>
<tr>
<td>CAM-B3LYP (Toluene)</td>
<td>LR</td>
<td>4.34 (0.45)</td>
<td>4.59 (0.54)</td>
<td>0.25</td>
<td>4.57 (0.08)</td>
</tr>
<tr>
<td></td>
<td>cLR</td>
<td>4.32 (0.45)</td>
<td>4.64 (0.54)</td>
<td>0.32</td>
<td>4.57 (0.08)</td>
</tr>
<tr>
<td></td>
<td>SS</td>
<td>3.90 (0.27)</td>
<td>4.58 (0.50)</td>
<td>0.68</td>
<td>4.47 (0.73)</td>
</tr>
<tr>
<td>CAM-B3LYP (THF)</td>
<td>LR</td>
<td>4.37 (0.33)</td>
<td>4.58 (0.65)</td>
<td>0.21</td>
<td>4.56 (0.06)</td>
</tr>
<tr>
<td></td>
<td>cLR</td>
<td>4.35 (0.33)</td>
<td>4.62 (0.65)</td>
<td>0.27</td>
<td>4.55 (0.06)</td>
</tr>
<tr>
<td></td>
<td>SS</td>
<td>4.08 (0.23)</td>
<td>4.54 (0.59)</td>
<td>0.48</td>
<td>4.35 (0.72)</td>
</tr>
<tr>
<td>CAM-B3LYP (DCM)</td>
<td>LR</td>
<td>4.37 (0.32)</td>
<td>4.57 (0.66)</td>
<td>0.20</td>
<td>4.55 (0.06)</td>
</tr>
<tr>
<td></td>
<td>cLR</td>
<td>4.35 (0.32)</td>
<td>4.62 (0.66)</td>
<td>0.27</td>
<td>4.55 (0.06)</td>
</tr>
<tr>
<td></td>
<td>SS</td>
<td>4.05 (0.22)</td>
<td>4.54 (0.60)</td>
<td>0.49</td>
<td>4.33 (0.71)</td>
</tr>
<tr>
<td>( \omega )B97X (THF)</td>
<td>LR</td>
<td>4.63 (0.16)</td>
<td>4.78 (0.92)</td>
<td>0.15</td>
<td>4.70 (0.06)</td>
</tr>
<tr>
<td></td>
<td>cLR</td>
<td>4.64 (0.16)</td>
<td>4.84 (0.92)</td>
<td>0.20</td>
<td>4.70 (0.06)</td>
</tr>
<tr>
<td></td>
<td>SS</td>
<td>4.58 (0.08)</td>
<td>4.78 (0.83)</td>
<td>0.20</td>
<td>4.62 (0.04)</td>
</tr>
<tr>
<td>( \omega )B97X-D (THF)</td>
<td>LR</td>
<td>4.46 (0.20)</td>
<td>4.63 (0.87)</td>
<td>0.17</td>
<td>4.52 (0.09)</td>
</tr>
<tr>
<td></td>
<td>cLR</td>
<td>4.47 (0.20)</td>
<td>4.68 (0.87)</td>
<td>0.21</td>
<td>4.55 (0.09)</td>
</tr>
<tr>
<td></td>
<td>SS</td>
<td>4.40 (0.12)</td>
<td>4.61 (0.75)</td>
<td>0.21</td>
<td>4.48 (0.04)</td>
</tr>
</tbody>
</table>

SS-PCM PBE0 values completely off (experimental) target.

With CAM-B3LYP and the LR-PCM model, all three solvents provide comparable energy shifts. With CAM-B3LYP, the cLR correction remains moderate, if not negligible, while SS corrections are significant for both regioisomers, ranging from 0.2 to 0.4 eV. At the SS-CAM-B3LYP level of theory, the oscillator strength ratio of the first two ES of da improves compared to LR-CAM-B3LYP but at the expense of an increased energy splitting between the two states. For dai, SS corrections induce a larger stabilization for the bright excited state for all three solvent, the first states becoming the brightest. Consequently, the SS-PCM CAM-B3LYP energy shifts are in qualitatively satisfying agreement with the reference data, though the \( \delta (\Delta \omega_{abs}) \) is slightly overestimated. Both Head-Gordon’s range-separated hybrids, namely \( \omega \)B97X and \( \omega \)B97X-D, give similar results, that is comparable energies and oscillator strengths with the three solvation models. Moreover, both cLR and SS corrections remain small (\( \leq 0.15 \) eV). Subsequently, in contrast with the CAM-B3LYP situation, no inversion of the nature of the two first excited states is found for dai. As can be seen in Table 6 the best agreement with experiment for both \( \delta (\omega_{abs}^{\text{max}}) \) and \( \delta (\Delta \omega_{abs}) \) is obtained with a method combining \( \omega \)B97X-D and SS-PCM.

3.4 Emission

Vertical fluorescence energies and oscillator strengths are listed in Table 7 for geometries optimized for the lowest lying excited-states, following Kasha’s rule. These optimizations have been performed in the equilibrium limit of the PCM model. At the LR-PCM level, only PBE0 restores the correct trend for \( \delta (\omega_{abs}^{\text{max}}) \) with little differences between toluene and THF. However, the LR model is known to be rarely the most suited for emission properties. The SS-PCM PBE0 fluorescence energies are completely off the experimental value, as already noted for the absorption. Indeed, for da, PBE0 transition energies decrease by 1 and 2 eV for toluene and THF, respectively, and the oscillator strengths are reduced by a factor of ca. 2. For dai, while corrections introduced by the SS scheme remain modest in toluene, they become huge in THF and attain about the same amplitude as for da. These dramatic effects reveal that combining the SS-PCM model to the PBE0 functional is inadequate in the present case.

With CAM-B3LYP, the SS corrections are more reasonable for dai in all solvents. On the other hand, for da transition energies and oscillator strengths are markedly reduced by SS corrections in polar medium. In toluene, the changes stay within a reasonable range and the \( \delta (\omega_{abs}^{\text{max}}) \) at the SS-CAM-B3LYP coincide nicely with the experimental value. Never-
3.5 0-0 energies

A meaningful choice to check the absolute agreement between experiment and theory is to determine the 0-0 energies \( E_{00} \) that can be compared to the crossing point between absorption and emission curves. To accurately determine this crossing point, we have used a computational protocol defined elsewhere,\(^{56}\) and that includes corrections for both zero-point vibrational energy (ZPVE) and non-equilibrium solvation effects. The obtained values are reported in Table 8 together with corresponding energy shifts between the two isomers, in order to allow easy comparison with experimental data. Overall the obtained trends are in line with the conclusions drawn for absorption and emission namely: i) within the LR-PCM scheme the global hybrid PBE0 gives \( \delta(E_{00}) \) in agreement with experiment while range separated hybrids perform poorly, giving either small negative shifts (CAM-B3LYP) or a deficient \( \text{da} \) and \( \text{dai} \) ranking (\( \omega \)B97X and \( \omega \)B97X-D); ii) with the SS-PCM model, PBE0 completely fails as a result of the huge overstabilization of states having a charge transfer character; iii) the same occurs for CAM-B3LYP in THF while it does not in toluene; and iv) \( \omega \)B97X-D provides the correct ordering at SS-PCM level. It is also noteworthy that the SS-PCM \( \omega \)B97X-D \( E_{00} \) energies are 3.16 and 3.62 eV for \( \text{da} \) and \( \text{dai} \), respectively, in reasonable match with the experimental values of 3.30 and 3.55 eV. Indeed, error of ca. a quarter of an eV are typical for such TD-DFT calculations.\(^{56}\)

Table 8 Computed 0-0 reference, \( E_{00} \), and the corresponding energy shifts between \( \text{da} \) and \( \text{dai} \), \( \delta(E_{00}) \). All data used the 6-31+G(d) atomic basis set and the geometries are optimized at the same level of theory as the one used for the TD-DFT part. Relevant experimental values in THF are also given to facilitate comparison.

<table>
<thead>
<tr>
<th></th>
<th>( \text{da} )</th>
<th>( \text{dai} )</th>
<th>( \delta(E_{00}) )</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>LR</td>
<td>SS</td>
<td>LR</td>
</tr>
<tr>
<td>PBE0 (Tol)</td>
<td>3.24</td>
<td>3.52</td>
<td>3.17</td>
</tr>
<tr>
<td>PBE0 (THF)</td>
<td>3.19</td>
<td>3.42</td>
<td>2.26</td>
</tr>
<tr>
<td>CAM-B3LYP (Tol)</td>
<td>3.82</td>
<td>3.40</td>
<td>3.79</td>
</tr>
<tr>
<td>CAM-B3LYP (THF)</td>
<td>4.06</td>
<td>3.91</td>
<td>4.04</td>
</tr>
<tr>
<td>( \omega )B97X (THF)</td>
<td>3.88</td>
<td>3.16</td>
<td>3.85</td>
</tr>
<tr>
<td>( \omega )B97X-D (THF)</td>
<td>3.30</td>
<td>3.30</td>
<td>3.55</td>
</tr>
</tbody>
</table>

3.6 Stokes shifts

Last, let us discuss the Stokes shifts reported Table 9 as they provide a complementary path to \( E_{00} \) for relevant theory–experiment comparisons. Of course, one could also wish to compute vibrationally resolved spectra, so to compare band shapes, but the experimental absorption and emission spectra are broad and structureless, making the vibronic simulations relatively pointless. Contrary to the energies of Table 8, the Stokes shifts reveal the dramatic effects of solvatochromic effects with PBE0. This marked change is the direct consequence of the relative oscillator strength of the excited state contributing to the first absorption band of \( \text{da} \) and brings the LR-PCM(THF) PBE0 result in good agreement with the experimental Stokes shifts and \( \delta(\Delta_{SS}) \). All other LR calculations fail to reproduce the correct sign of \( \delta(\Delta_{SS}) \), but, as stated above, LR is not a very good model for emission nor, consequently, Stokes shifts. With the SS-PCM model, the unphysical large SS corrections obtained for PBE0 absorption and emission prevents any rationale analysis of the Stokes shifts.

Table 7 Vertical emission wavelengths computed for the two dyes in eV, the number between brackets are the corresponding oscillator strengths (in a.u.). See caption of Table 6 for more details. At the rightmost part of the Table, the \( \delta(E_{\text{max}}) \) are reported. Relevant experimental values in THF are also shown to facilitate comparison.

<table>
<thead>
<tr>
<th></th>
<th>( \text{da} )</th>
<th>( \text{dai} )</th>
<th>( \delta(E_{\text{max}}) )</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>LR</td>
<td>SS</td>
<td>LR</td>
</tr>
<tr>
<td>PBE0 (Tol)</td>
<td>3.11</td>
<td>2.09</td>
<td>3.30</td>
</tr>
<tr>
<td>PBE0 (THF)</td>
<td>3.04</td>
<td>1.03</td>
<td>3.24</td>
</tr>
<tr>
<td>CAM-B3LYP (Tol)</td>
<td>3.61</td>
<td>3.18</td>
<td>3.60</td>
</tr>
<tr>
<td>CAM-B3LYP (THF)</td>
<td>3.55</td>
<td>2.29</td>
<td>3.48</td>
</tr>
<tr>
<td>CAM-B3LYP (DCM)</td>
<td>3.56</td>
<td>2.20</td>
<td>3.46</td>
</tr>
<tr>
<td>( \omega )B97X (THF)</td>
<td>3.84</td>
<td>3.63</td>
<td>3.67</td>
</tr>
<tr>
<td>( \omega )B97X-D (THF)</td>
<td>3.66</td>
<td>2.79</td>
<td>3.54</td>
</tr>
</tbody>
</table>
computed with PBE0, and the same holds for CAM-B3LYP results in THF. In toluene, the CAM-B3LYP data reasonably match experiment, as SS-PCM CAM-B3LYP vertical energies were meaningful (see above). Once again, ωB97X-D provides accurate results, better matching experiment than their ωB97X counterparts.

Table 9 Computed Stokes shifts (derived from convoluted bands with experimental FWHM when the two bright states relevant for absorption are close in energy), and corresponding energy shifts δ(ΔSS) between da and dai. Relevant experimental values in THF are also given to facilitate comparison.

<table>
<thead>
<tr>
<th></th>
<th>da</th>
<th>dai</th>
<th>δ(ΔSS)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>LR</td>
<td>SS</td>
<td>LR</td>
</tr>
<tr>
<td>PBE0 (Tol)</td>
<td>0.55</td>
<td>0.44</td>
<td>0.81</td>
</tr>
<tr>
<td>PBE0 (THF)</td>
<td>1.25</td>
<td>1.56</td>
<td>0.83</td>
</tr>
<tr>
<td>CAM-B3LYP (Tol)</td>
<td>0.87</td>
<td>1.40</td>
<td>1.05</td>
</tr>
<tr>
<td>CAM-B3LYP (THF)</td>
<td>0.97</td>
<td>2.23</td>
<td>1.17</td>
</tr>
<tr>
<td>ωB97X (THF)</td>
<td>0.92</td>
<td>1.14</td>
<td>1.27</td>
</tr>
<tr>
<td>ωB97X-D (THF)</td>
<td>0.95</td>
<td>1.80</td>
<td>1.14</td>
</tr>
<tr>
<td>Experiment (THF)</td>
<td>1.39</td>
<td>1.39</td>
<td>1.00</td>
</tr>
</tbody>
</table>

4 Conclusions

Starting from the experimental absorption and emission spectra of two model chromophores based on the 1,2,3-triazole moiety, we have thoroughly investigated the ability of TD-DFT methodologies to describe their main photophysical features. Our findings show that these regio-isomers are particularly challenging compounds for TD-DFT and, consequently, can be viewed as interesting benchmarks to evidence potential deficiencies of both new exchange-correlation functionals and/or solvent models.

First, none of the implemented level of theories, ranging from global hybrids (B3LYP, PBE0) to range-separated functionals without (CAM-B3LYP, ωB97X) or with dispersion corrections (ωB97X-D), was able to recover all key features of the two regio-isomers in vacuum. Secondly, accounting for solvent effects within different PCM approaches may conduct to quantitative agreement with experiment, but assessing the potential of several models suggests that some theory-experiment match might be fortuitous. Indeed, within the LR approximation of the PCM model, none of the range-separated hybrids gives satisfactory trends whereas PBE0 results are rather good, especially for the absorption-fluorescence crossing points. However, LR-PCM PBE0 calculations lead to marked differences for absorption features of da when going from toluene to THF, with a better agreement between experiment and calculations for the latter. Moreover, both cLR and SS schemes of the PCM model lead to dramatic decreases of transition energies, most probably related to the appearance of spurious charge transfer states, an artifact common of global hybrids but apparently enhanced with SS-PCM, making the SS-PCM-PBE0 combination clearly out of control in the present cases.

SS-PCM corrections are larger for da than for dai and bigger in THF than in toluene when using range-corrected hybrids. While the magnitude of these corrections remain reasonable for both the absorption and emission of dai, they become unphysically large for most of the fluorescence wavelengths computed for da, e.g., SS-PCM(THF) CAM-B3LYP results are both large and leading to non satisfying match with experiment. When using the ωB97X exchange-correlation functional, that incorporates larger amount of exact exchange, no methodological catastrophe is to be found but the experimental differences between the two isomers are not restored. Though the SS-PCM corrections for the fluorescence of da remain large, ωB97X-D, that allows to recover most of the experimental trends for the systems investigated herein, could be the best available compromise approach.

In summary, our findings suggest that the SS-PCM approach may be useful to reveal the charge transfer failure of current and new exchange-correlation functionals, even though it may improves the description in some cases. Such tendency was noted before, though much less dramatically then here, as only systems with a strong intramolecular charge-transfer described with global hybrids were shown to exhibit an incorrect behaviour of the SS-PCM approach. Last but not least, this work also prescribes particular circumvention about TD-DFT results obtained for 1,2,3-triazole based chromophores.

Acknowledgement

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References


Photophysical signatures of 1,2,3-triazole regioisomers: a surprisingly challenging problem for TD-DFT.