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This is a pre print version of the following article:
Original Citation:
Availability:
This version is available http://hdl.handle.net/2318/1589130 since 2016-09-01T14:23:36Z
Published version:
DOI:10.1021/acs.jpca.6b05220
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(Article begins on next page)





This is the author's final version of the contribution published as:

Capobianco, Amedeo; Borrelli, Raffaele; Landi, Alessandro; Velardo, Amalia; Peluso, Andrea. Absorption Band Shapes of a Push-Pull Dye Approaching the Cyanine Limit: A Challenging Case for First Principle Calculations. JOURNAL OF PHYSICAL CHEMISTRY. A, MOLECULES, SPECTROSCOPY, KINETICS, ENVIRONMENT, & GENERAL THEORY. 120 (28) pp: 5581-5589. DOI: 10.1021/acs.jpca.6b05220

The publisher's version is available at: http://pubs.acs.org/doi/abs/10.1021/acs.jpca.6b05220

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# The Absorption Band Shapes of a Push-Pull Dye Approaching the Cyanine Limit: A Challenging Case for First Principle Calculations

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#### Abstract

The absorption band shapes of a solvent tunable donor-acceptor dye have been theoretically investigated by using Kubo's generating function approach, with minimum energy geometries and normal modes computed at DFT level of theory. The adopted computational procedure allows to include in the computation of Franck-Condon factors the whole set of normal modes, without any limitation on excitation quanta, allowing for an almost quantitative reproduction of the absorption band-shape when the equilibrium geometries of the ground and the excited states are well predicted by electronic computations. Noteworthy, the functionals which yield more accurate band shapes also provide good prediction of the dipole moment variations upon excitation; since the latter quantities are rarely available, theoretical simulation of band shapes could be a powerful tool for choosing the most appropriate computational method for predictive purposes.

### Introduction

Organic donor-acceptor dyes approaching the cyanine limit, i.e molecules with a  $\pi$ -conjugated backbone exhibiting vanishing bond length alternation (BLA),<sup>?</sup> have found applications in several technological fields, such as dyesensitized and bulk-heterojunction solar cells, <sup>1-4</sup> optical storage media, <sup>5,6</sup> and nonlinear optics.<sup>7–10</sup> Highly reliable computational methods are therefore needed for a deeper understanding of their electronic structures and for the rational design of new ones, which should logically include the estimation of the rates of the photochemical processes which occur in devices.<sup>11</sup> However, the easily polarizable electronic clouds of those compounds make them problematic and challenging systems for most of the computational approaches for large size molecules.<sup>12,13</sup> Time dependent (TD) density functional theory (DFT) predicts vertical excitation energies which significantly disagree with those corresponding to the observed absorption peaks in solution.<sup>14</sup> The discrepancy can be partly ascribed to an insufficient description of double-excitations at TDDFT level,<sup>12,15–17</sup> and partly imputed to nuclear relaxations upon excitation, which makes vertical transitions not corresponding to maximum absorption wavelengths.<sup>18,19</sup> Simulations of the whole absorption band shapes are therefore necessary for better assessing the reliability of electronic computations.<sup>11</sup>

Herein we report first principle calculations of the band shape of the donor-acceptor (Z)-2-(5-((5-(dibutylamino)thiophen-2-yl)methylene)thiophen-

2(5H)-ylidene)malononitrile (1) dye, see Scheme 1, which represents a very interesting test case because its absorption spectrum strongly depends on the polarity of the solvent, exhibiting not only a red-shift of the absorption maximum but also a remarkable change of the whole band shape.<sup>20</sup>



Scheme 1: The neutral (left) and zwitterionic (right) resonance forms of compound **1**. The path used to compute bond length alternation has been highlighted in red.

By employing a wide palette of density functional theory (DFT) approaches, we will show that those functionals which provide a satisfying reproduction of the absorption band shape are the only ones capable of reproducing the observed dipole moment change upon transition. Since the latter quantities are difficult to obtain and are rarely available in the literature,<sup>?</sup> the simulation of band shapes can represent an alternative feasible way for choosing the most appropriate computational method for a given class of molecules.

# Spectral Band Shapes

According to first order time dependent perturbation theory the absorption cross section per unit time for a radiative transition between the electronic states  $\left|i\right\rangle$  and  $\left|f\right\rangle$  is:  $^{21,22}$ 

$$\sigma(\nu) = \frac{8\pi^{3}\nu}{3nc}I_{fi}(\nu)$$

$$I_{fi}(\nu) = \sum_{l}\sum_{m} |\langle fm|\,\mu\,|il\rangle|^{2}\,e^{-\beta E_{il}}\delta(E_{fm} - E_{il} - h\nu)/Z_{i}$$
(1)

where c and n stand for the light velocity in vacuum and the index of refraction,  $\mu$  denotes the dipole moment operator, m and l specify the whole set of vibrational quantum numbers of  $|f\rangle$  and  $|i\rangle$ , respectively, and  $Z_i$  is the partition function of the initial state  $|i\rangle$ . The infinite summations appearing in Eq. ?? poses computational problems which, following the seminal works of Lax and Kubo,<sup>21,22</sup> can be conveniently avoided by introducing the integral representation of Dirac's  $\delta$  function:

$$\delta(E_{fm} - E_{il} - h\nu) = h^{-1} \int_{-\infty}^{\infty} exp\left[i\left(E_{fm} - E_{il} - h\nu\right)\tau/\hbar\right] d\tau \qquad (2)$$

By substituting Eqn 2 in Eqn 1 and integrating over the electronic coordinates:

$$I_{fi}(\nu) = h^{-1} \int d\tau e^{-i\omega\tau} I_{fi}(\tau), \qquad (3)$$

where the dummy variable  $\tau$  has units of time.

Eqn 3 allows to obtain the frequency distribution function  $I_{fi}(\nu)$ , i.e. the spectral band shape, as the inverse Fourier transform of the function  $I_{fi}(\tau)$ :

$$I_{fi}(\tau) = Tr\left[\langle l | \mu_{if} e^{\lambda \mathcal{H}_f} \mu_{fi} e^{-(\beta+\lambda)\mathcal{H}_i} | l \rangle\right]$$
(4)

where  $\lambda = it/\hbar$ ,  $\beta = 1/k_BT$ , and  $\mathcal{H}_i, \mathcal{H}_f$  are the Hamiltonian operators of the initial and final electronic states, respectively. If the latter ones are expressed in harmonic approximation, the infinite summation of eqn 4 can be evaluated into a closed form by a formula formula due to Melher,<sup>?</sup>? which allows for an analytical integration over the vibrational coordinates,<sup>19,21? -26</sup> making use of Duschinsky's transformation, which allows to express the vibrational coordinates of one electronic states in terms of those of the other:

$$\mathbf{Q}_{\mathbf{i}} = \mathbf{J}\mathbf{Q}_{\mathbf{f}} + \mathbf{K} \tag{5}$$

where  $\mathbf{J}$  is a rotation matrix and  $\mathbf{K}$  the vector of the equilibrium position displacements.

The GF approach provides several advantages with respect to the standard recursive calculation of Franck-Condon factors,<sup>27? -29</sup> inasmuch as it allows to include in computations the whole set of the molecular normal modes, taking into account both the effects due to changes of the equilibrium positions and of vibrational frequencies, as well as the effects due to normal mode mixing. Remarkably, the GF approach does not pose any limitation on the number of modes which can be excited and on their highest quantum numbers. Thus the main limitation consists in neglecting anhar-

monic effects; in principle they can be included, but the high computational costs pose a limit to the number of vibrational modes which can be treated anharmonically.<sup>30–32</sup> Apart from the use of an apodization function in the fast Fourier transform, <sup>24,25,33</sup> no external adjustable parameters are needed, so that spectral band shapes obtained by the GF approach ensure an effective test of the performances of the electronic calculations.

# Computational procedure

Absorption spectra were computed by using a local development version of MolFC program.<sup>33,34</sup> The curvilinear coordinate representation of the normal modes has been adopted to prevent that displacements of angular coordinates could result into unrealistic shifts of stretching coordinates upon excitation.<sup>30,35,36</sup> That problem is unavoidable in the rectilinear Cartesian coordinate representation and requires the use of high order anharmonic potentials for its correction.<sup>31,32,37</sup>

The minimum energy geometries of the ground and the first excited states of **1** have been computed at density functional theory level, by using a wide palette of methods, including pure and global hybrid GGA and meta GGA exchange-correlation potentials, see Table 1.

We have also used range separated hybrid functionals in which the exchange component is divided into short-range (SR) and long-range (LR) terms by splitting the Coulomb operator via the error function.<sup>45</sup> Adopting

Functional	$Type^{a}$	$\%~{\rm HF}~{\rm exchange}^b$	ref
PBE	GGA	0	38
M06-L	mGGA	0	39
B3LYP	GH-GGA	20	$40,\!41$
PBE0	GH-GGA	25	42
CAM-B3LYP	RSH-GGA	$19 \xrightarrow{0.33} 65$	43
M06-2X	GH-mGGA	54	39
$\mathrm{CAM}^{*\ c}$	RSH-GGA	$30 \xrightarrow{0.33} 90$	43
$\omega B97X$	RSH-GGA	$16 \xrightarrow{0.30} 100$	44

Table 1: Functionals used in the present work.

<sup>*a*</sup>GGA = generalized gradient approximation, GH = global hybrid, RSH = range-separated hybrid, mGGA = meta-GGA. <sup>*b*</sup> The percentages of HF exchange in the short (left,  $\alpha$  in eq. 6) and long range (right,  $\alpha + \beta$  in eq. 6) limits are reported for RSH functionals; the parameter  $\omega$  (bohr<sup>-1</sup>) is reported over the arrow. <sup>*c*</sup> CAM-B3LYP with  $\alpha$  and  $\beta$  set to 0.30 and 0.60, respectively.

the formalism of ref 43:

$$\frac{1}{r_{12}} = \underbrace{\frac{1 - \left[\alpha + \beta \operatorname{erf}\left(\omega r_{12}\right)\right]}{r_{12}}}_{SR} + \underbrace{\frac{\alpha + \beta \operatorname{erf}\left(\omega r_{12}\right)}{r_{12}}}_{LR},\tag{6}$$

with  $0 \le \alpha \le 1$ ,  $0 \le \beta \le 1$ ,  $0 \le \alpha + \beta \le 1$ .

The DFT exchange interaction is given by the first term of eq. 6, while the long-range orbital-orbital exchange interactions are expressed as Hartree-Fock (HF) exchange integrals. The parameter  $\alpha$  is the fraction of HF exchange which contributes over the whole range; the parameter  $\beta$  incorporates the DFT counterpart by a factor of  $1 - \alpha - \beta$  and acts in such a way that  $\alpha + \beta$  is the fraction of HF exchange in the asymptotic limit;  $\omega$  defines the range of the separation: the larger its value, the sharper is SR-LR separation. Among long range-corrected functionals, we have considered  $\omega$ B97X which recovers the exact 1/r asymptotic behavior of the exchange potential and the 'Coulomb attenuating' CAM-B3LYP functional.<sup>43,44</sup> We have also introduced CAM\*, a modification of CAM-B3LYP in which a larger fraction of HF exchange has been imposed by increasing the  $\alpha$  and  $\beta$  parameters up to 0.30 and 0.60 respectively, while leaving  $\omega$  at its default value (0.33 bohr<sup>-1</sup>), see Table 1.

Effects due to solvent polarization were included by the polarizable continuum model (PCM).<sup>46,47</sup> The non-equilibrium solvation scheme, in which the effects of the slow motion of the solvent are not included, has been used for excited states.<sup>48</sup> The 6-31+G(d,p) basis set was employed for DFT and TDDFT computations because it provides reliable ground and excited state equilibrium geometries and dipole moments.<sup>49–51</sup> Indeed calculations of ground state equilibrium geometries and vertical excitation energies carried out by using the 6-311++G(2df,2p) basis set gave essentially the same results as those obtained by the smaller basis set, see Tables S1-S3 in the Supporting Information.

(PCM)MP2/6-31+G(d) geometry optimizations and computations of ground state dipole moments were also performed to have reference values for the solvents in which electrooptical absorption (EOA) measurements are not available. MP2 was chosen because it is known to give reliable dipole

moments,  $^{52}$  even for donor-acceptor systems.  $^{50,51}$  The Gaussian package was used for MP2, DFT and TDDFT calculations.  $^{53}$ 

Coupled cluster computations including single and approximate double excitations (CC2) in conjunction with the resolution of identity approximation and the aug-cc-pVDZ basis set were carried out by using the TURBO-MOLE suite of programs.<sup>54–58</sup> The frozen core approximation was used in MP2 and CC2 computations. The butyl chains of **1** (Scheme 1) have been replaced by methyl groups in all the calculations.

# Results

In its most stable Z configuration, chromophore **1** can assume the *s*-trans and the *s*-cis conformations shown in Scheme 2.

Scheme 2: *s-trans* and *s-cis* conformations of **1**. They differ for the torsion coordinate highlighted in red. S S S CN Bu CN Bu The predicted relative energies are reported in Table 2: *s-trans* confor-Bu s-transion is more stable than *s-ciscis*ne according to all the tested methods, in line with <sup>1</sup>H-NMR data.<sup>59</sup>

	gas	DX	DE	AN
PBE	1.3	1.4	1.4	1.5
B3LYP	1.0	1.2	1.3	1.4
CAM-B3LYP	0.9	1.1	1.2	1.4
M06-2X	1.4	1.6	1.8	2.0
$CAM^*$	0.8	1.0	1.2	1.5
$\omega B97X$	0.9	1.1	1.3	1.6
MP2	1.4	1.5	1.6	1.8

Table 2: Predicted energy differences (kcal/mol) between *s-cis* and *s-trans* conformers of compound 1. DX = 1,4-dioxane, DE = diethyl ether, AN = acetonitrile.

Dipole moments Experimental and computed ground and excited state dipole moments of 1 are reported in Table 3. Only functionals possessing a high fraction of HF exchange yield dipole moment changes upon excitation ( $\Delta \mu = \mu_e - \mu_g$ ) in reasonable agreement with experimental results and MP2 computations, which are known to give reliable results.<sup>50,51,61</sup> In dioxane, CAM\* and  $\omega$ B97X predict  $\Delta \mu$ 's of 4.6 and 4.8 D, respectively, in good agreement with EOA measurements (5 D), whereas pure functionals and functionals with lower fraction of HF exchange, refer to Table 1, yield  $\Delta \mu$  as low as 0.1 and  $\approx$ 1 D, respectively. Upon increasing the fraction of HF exchange in the functional, predicted ground state dipole moments decrease whereas excited state ones increase, leading to a significant increase of  $\Delta \mu$ 's.

Absorption band shapes In non-polar solvents, the absorption spectrum of compound **1** is characterized by two well resolved peaks, falling in 1,4-dioxane at 15385 and 16491 cm<sup>-1</sup> and possessing roughly the same intensity, see Figure 1.<sup>20</sup> As the solvent polarity increases, the longer wavelength absorption gains intensity and shifts to lower frequencies (ca 15110 cm<sup>-1</sup> in acetonitrile, Figure 1, orange line), whereas the shorter wavelength peak becomes a shoulder.

Figure 1: UV/Vis absorption spectra of dye **1** in solvents of different polarity at 298 K ( $c = 10^{-5}$  M). The thick black line is for the least polar solvent diethyl ether (in 1,4-dioxane the spectra is almost identical), blue line: ethyl acetate, purple line: tetrahydrofuran, red line: acetone, and orange line: acetonitrile. The arrow indicates the spectral shift with increasing solvent polarity. Reproduced with permission from ref 20. Copyright (2008) John Wiley and Sons.

Since absorption spectra are known to exhibit only a marginal dependence on conformational degree of freedom, we have first focused attention on the most stable *s*-trans conformer, c.f. Table 2. The computed absorption spectra at T=298 K in low polar environment (1,4-dioxane) are reported in Figure 2, together with the experimental spectrum (dashed line). Intensities have been scaled in such a way that all the spectra possess the same area. For a proper comparison of band-shapes, spectra have been aligned by

making coincident the computed and the experimental wavenumbers of the minimum absorption between the two peaks, but for panel a) of Figure 2 where the alignment has been done on the longer wavelength peak. Panel a) of Figure 2 shows that pure functionals as well as functionals with low fraction of HF exchange are not able to reproduce the essential features of the experimental spectrum. The predicted band shapes exhibit only one peak instead of two, with a small shoulder at shorter wavelength. Furthermore, the predicted bandwidths are significantly narrower than the experimental one.

The agreement between observed and predicted band shapes improves significantly by employing equilibrium geometries and normal modes predicted by functionals with higher fraction of HF exchange. Figure 2b shows that CAM-B3LYP yields in 1,4-dioxane two well resolved peaks, whose intensity ratio is however significantly different from the observed (1:1) one. Further improvement is obtained by increasing the amount of Hartree-Fock exchange in the functional: the CAM\*, our modification of CAM-B3LYP discussed above, see Table 1, yields an absorption band shape characterized by two peaks with intensity ratio 1.2:1, in satisfying agreement with the experimental one, see Figure 2c. The energy difference between the two absorption peaks is slightly overestimated (1430 vs  $\approx$ 1100 cm<sup>-1</sup>) and the simulated spectrum is slightly broader than the observed one. Those small discrepancies can be attributed to anharmonic effects, which can be simply corrected –without affecting peak intensity ratios– by using standard scal-





Figure 2: Predicted UV/absorption spectra of 1 (*s*-trans conformer) in low polarity solvents. The experimental spectrum recorded in DE has been superimposed as a dashed line. Panel a): PBE(DX), red line; M06-L(DX), blue line; B3LYP(DX), green line. Panel b): CAM-B3LYP(DX). Panel c): CAM\*(DX). Panel d): CAM\* in the gas phase.

The intensity ratio between the two absorption peaks approaches the observed 1:1 value by using predicted CAM\* geometries and normal modes in the gas-phase, Figure 2d, suggesting that solvent polarization effects modeled by PCM could be slightly overestimated.<sup>18</sup> M06-2X and  $\omega$ B97X func-

tionals give band shapes very similar to that obtained by CAM-B3LYP and CAM\*, respectively (see Figure S1 in the Supporting Information), thus showing that the amount of exact exchange is the key factor to achieve a good accuracy for the vibronic spectra of compound **1**.

In weakly polar environments *s*-*cis* is predicted to be less stable than *s*-trans conformer by ca. 1 kcal/mol according to CAM\* and  $\omega$ B97X functionals (Table 2). Given that exiguous energy difference, close to calculation accuracy, it is possible that *s*-*cis* is sufficiently populated to be detected at room temperature. We have therefore also computed the spectrum of that conformer in low polar environments (gas and dioxane) by using the best performing functionals (CAM\* and  $\omega$ B97X). Figure 3 reports the gas-phase spectra obtained at the CAM\* level for *s*-trans and *s*-*cis* conformers. As expected, both conformers exhibit very similar spectra, at variance with 1/s-trans the 1/s-*cis* exhibits the shorter wavelength peak slightly more intense than the longer wavelength one. The same also holds in 1,4-dioxane, see Figure S3 in the Supporting Information.

Computed (T=298 K) and experimental absorption spectra in acetonitrile are reported in Figure 4. They have been aligned by making coincident the wavenumber of the absorption maximum; intensities have been properly normalized, so that all the spectra possess the same area. As the polarity of solvent increases, bandwidths are predicted to be narrower and less structured than those obtained in low polar solvents (compare with Figure 2), in line with experimental results. With the exception of PBE, all the function-



Figure 3: Predicted (CAM\*, gas) UV/absorption spectra of 1, *s*-trans conformer, red line; *s*-cis conformer, black line.

als yield a reasonably accurate spectrum, but still high-exchange functionals (Figure 4, panels c and d) ensure the best agreement with the experiment.

The assignments of the two peaks in low polar solvents is not a simple matter because, apart from one low frequency mode which is significantly more displaced than all the others, there is a huge number (30-40) of slightly displaced modes. The components of the equilibrium position displacement vector **K** predicted by two prototypical functionals, B3LYP and CAM<sup>\*</sup>, in the gas-phase and in acetonitrile, are reported in Table ??. In both environments and according to both functionals, the most displaced mode is an in plane low frequency bending mode, which involves the whole molecular backbone. The frequency of that mode is too low for being responsible of the two peaks observed in low polar environments. At room temperature that mode plays an important role, inasmuch as it broadens the absorption bands by providing vibronic progressions which, because hot vibronic states



Figure 4: Predicted and observed (dashed line) UV/absorption spectra of **1** in acetonitrile.

are populated, fall around the most intense transitions, both at shorter and longer wavelengths. As concerns the higher frequency modes, the comparison between the **K** components predicted by B3LYP and CAM\* indicates that the most significant differences are found in the region 1500-1600 cm<sup>-1</sup>, where CAM\* predict three modes, whose equilibrium position displacement is higher than 0.4 Åuma<sup>-1/2</sup>, whereas no displaced modes are predicted by B3LYP in that wavenumber region. The computed CAM\* and B3LYP spectra of **1** at 0 K, obtained by using the usual recursive FC calculation, are

displayed in figure ??. Only the first ten most displaced vibrational modes are allowed to change their quantum number upon electronic transition in those computations. While the B3LYP spectrum extends up to 1400 cm<sup>-1</sup>, showing only two major absorption regions, one in the range 0-500 cm<sup>-1</sup>, the other extending from 1000 to 1300 cm<sup>-1</sup>, the CAM\* spectrum exhibits a third absorption region, from 1500 up to 1800 cm<sup>-1</sup>. In that region at least three vibronic progressions can be distinguished, involving  $0 \rightarrow 1$  transition of the displaced modes at  $\tilde{\nu} = 1521, 1574, 1601$  cm<sup>-1</sup>, accompanied by several  $0 \rightarrow n$  transitions of the most displaced low frequency mode, with n from 1 at least to 5, the maximum excitation number used in computations. The density of states provided by those transitions is significantly higher than that computed at 1000-1300 cm<sup>-1</sup>, giving rise to the second peak at shorter wavelength appearing in low polar solvents. The three displaced modes predicted by CAM\* at  $\tilde{\nu} = 1521, 1574, 1601$  cm<sup>-1</sup> correspond to stretching and bending vibrations of the whole molecular skeleton.

In acetonitrile, CAM<sup>\*</sup> computations predict that only one high frequency mode (1694 cm<sup>-1</sup>) is displaced. Although the displacement is significant ( $\approx 0.4$ ) a single displaced mode is not enough for providing a high density of states in that region, so that the second peak disappears and the whole spectrum becomes narrower.

Because of the high number of displaced modes, it could be useful to try to relate the absorption band shapes to some sort of collective coordinate, such as the bond length alternation (BLA) or the bond order alternation

(BOA). BLA is defined as the difference between the average lengths of the nominally single and double bonds along a  $\pi$ -conjugated backbone; it is an important physicochemical parameter for understanding the electronic properties of dyes, which, together with the bond order alternation, has proven particularly useful in designing organic materials with high nonlinear optical properties.<sup>?</sup> Here, we will consider only BLA, since absorption band shapes are more directly related to molecular geometry changes. Computed BLA are reported in Table 4, both for the ground and the excited states. For all the environments, the BLA increases upon increasing the fraction of exact exchange in the functional,<sup>63,64</sup> as seen e.g. by B3LYP and CAM\* estimates in dioxane, amounting to 0.02 and 0.05 Å, respectively. Noteworthy, only functionals giving BLA  $\geq 5 \times 10^{-2}$  Å are capable of reproducing both the observed  $\Delta \mu$  and the absorption band shape in low-polarity environments.

Absorption band shapes are very sensitive to BLA. In dioxane, B3LYP and CAM<sup>\*</sup> predict BLA changes upon transition of 0.03 and 0.05 Å, respectively, but band shapes are significantly different.

#### Absorption energies

Table 5 shows that vertical transition energies are systematically overestimated with respect to experimental maximum absorption energies; the poor performance on the prediction of reliable excitation energies for cya-

nines is a well documented issue, which is known to affect the quality of almost all the computational methods, not only TDDFT.<sup>65</sup> Noteworthy, our calculations have shown that maximum absorptions occurs, within 0.01 eV (80 cm<sup>-1</sup>), at the energy of the  $0 \leftarrow 0$  transition rather than at that corresponding to the vertical one. The energy differences between the vertical transition and the  $0 \leftarrow 0$  one can be as large as 0.24 eV, see Table 5, so that the computation of vibronic band shapes appears a necessary task to make a meaningful comparison of experimental and predicted transition energies. Indeed, a better agreement between theoretical and experimental maximum absorption energies,<sup>12,19,66</sup> see Table 5. Notwithstanding, predicted transition energies are still significantly overestimated; interestingly pure functionals or functional with a low ratio of HF exchange yield transition energies in better agreement with experimental ones, even though they fail in predicting the real electronic structure of **1**.

For the specific case under investigation, we have found that reliable transition energies can be obtained by using single point CC2 calculations with the minimum energy geometries obtained by functionals which better reproduce band shapes and estimating solvent polarization effects from DFT computations. Adopting the minimum energy geometries from  $\omega$ B97X optimizations of the ground and the first excited state, the electronic transition in the gas-phase is predicted at 2.21 eV by CC2 computations. Since the ground and excited state dipole moments predicted at CC2 level (12.3 and

18.5 D, respectively) are very similar to those obtained by the  $\omega$ B97X functional (see Table 3), the effect of solvent polarization can be safely taken from DFT computations ( $\approx 0.27$  eV in going from the gas-phase to 1,4dioxane), or estimated by the Onsager model. Using the former choice and neglecting zero point energy changes upon transition, we obtained  $E_{0\to0}=$ 1.94 eV, in pretty good agreement with experimental result (1.92 eV).

# Conclusions

Donor-acceptor dyes possessing easily polarizable electronic clouds are difficult systems for DFT computations. Methods based on pure or low percentage of HF exchange functionals yield excitation energies more close to the experimental ones, but they erroneously assign a cyanine-like character to chromophore **1** in low polar solvents, predicting a dipole moment change upon electronic transitions which is largely underestimated with respect to the results of EOA measurements. Instead, functionals with high percentage of HF exchange correctly reproduce the dipole moment variation upon excitation and provide reliable absorption band shapes. Since EOA measurements are scarce in the literature and difficult to obtain in highly polar solvents, band-shape computations can be an alternative reliable criterion for choosing the best computational tool for difficult cases as that considered here. It is also a necessary effort for a meaningful comparison between computed and observed excitation energies.

#### Acknowledgement

The financial supports of PON2007-2014 (Relight project) and of the University of Salerno are gratefully acknowledged.

#### Supporting Information Available

Spectra not shown in the article (Figures S1-S5). Dipole moments and excitation energies evaluated by using the 6-311++G(2df,2p) basis set (Tables S1-S3). Predicted BLA's for excited states (Table S4). This material is available free of charge via the Internet at http://pubs.acs.org/.

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	gas		D	DX		D	E	-	AN	
	$\mu_{ m g}$	$\mu_{ m e}$	$\mu_{ m g}$	$\mu_{ m e}$		$\mu_{ m g}$	$\mu_{ m e}$	$\mu_{ m g}$	$\mu_{ m e}$	
PBE	14.6	15.0	18.5	18.6		21.0	20.8	24.4	23.6	
M06-L	14.4	14.9	18.2	18.3		20.6	20.4	23.8	23.0	
B3LYP	14.4	15.5	18.2	19.1		20.8	21.2	24.4	23.9	
PBE0	14.2	15.5	18.0	19.0		20.6	21.1	24.2	23.6	
CAM-B3LYP	13.3	16.3	16.9	19.9		19.5	21.9	23.9	24.1	
M06-2X	12.7	16.0	16.2	19.6		18.7	21.6	22.7	23.7	
CAM*	12.6	16.6	15.7	20.3		18.1	22.4	22.8	24.5	
$\omega B97X$	12.4	16.4	15.5	20.3		18.1	22.5	22.9	24.5	
MP2	11.1	—	13.6	—		15.5	_	19.1	—	
$exper^b$	11.2	16.7	14.0	19.0		_	_	_	_	

Table 3: Computed and experimental ground  $(\mu_g)$  and excited state  $(\mu_e)^a$  dipole moments (D). DX = 1,4-dioxane, DE = diethyl ether, AN = acetonitrile. Computations refer to *s*-trans conformation.

<sup>*a*</sup>Vertical approximation, see ref 60. <sup>*b*</sup>Ref 20; gas phase values were obtained by using the Onsager model, see ref 59.

Table 4: Predicted ground (left) and excited (right) state BLA ( $10^{-2}$  Å).

	~	1.0	-			-		
	G	AS	DX		DE		A	N
PBE	2	-1	2	-2	1	-3	0	-4
M06-L	3	-1	2	-2	1	-3	0	-4
B3LYP	4	0	2	-1	2	-2	0	-2
PBE0	4	0	2	-1	2	-1	0	-2
CAM-B3LYP	5	0	4	0	3	0	0	0
M06-2X	5	1	4	0	3	0	0	0
$CAM^*$	6	0	5	0	4	0	2	0
$\omega B97X$	6	0	5	-1	4	0	2	0

		$\operatorname{gas}$			DX			AN	
	$E_{\rm m}$	$E_{00}$	$E_{\mathbf{v}}$	$E_{\rm m}$	$E_{00}$	$E_{\rm v}$	$E_{\rm m}$	$E_{00}$	$E_{\rm v}$
PBE	2.20	2.21	2.36	2.03	2.03	2.17	2.10	2.05	2.20
M06-L	2.34	2.25	2.50	2.17	2.17	2.31	2.17	2.19	2.33
B3LYP	2.36	2.38	2.53	2.17	2.17	2.31	2.19	2.19	2.32
PBE0	2.42	2.43	2.58	2.22	2.22	2.36	2.24	2.24	2.37
CAM-B3LYP	2.52	2.53	2.73	2.27	2.28	2.45	2.29	2.29	2.38
M06-2X	2.50	2.51	2.70	2.25	2.26	2.42	2.25	2.25	2.34
$CAM^*$	2.66	2.66	2.95	2.39	2.39	2.63	2.38	2.39	2.46
$\omega B97X$	2.60	2.60	2.88	2.33	2.32	2.56	2.31	2.32	2.38
exper.	_	_	—	1.92	_	—	1.87	_	—

Table 5: Predicted maximum absorption  $(E_{\rm m})$ , vertical  $(E_{\rm v})$  and  $E_{00}$  transition energies. All data are expressed in eV.

Table 6: Frequencies  $(\nu, \text{cm}^{-1})$  and equilibrium position displacements  $(K, \text{ Å uma}^{-1/2})$  of the most displaced normal modes of cyanine 1 in gas and acetonitrile, respectively

B3LY	$P E_{reorg} = 60.68$	35	CAM	* E <sub>reorg</sub> = 205.68	9	B3LYP E <sub>reorg</sub> =37.133 CAM <sup>*</sup> E <sub>reorg</sub>			$M^* E_{reorg} = 56.3$	28	
$\nu_{exc}$ (cm <sup>-1</sup> )	$\nu_{ground} (\text{cm}^{-1})$	K	$\nu_{exc}$ (cm <sup>-1</sup> )	$\nu_{ground} (\text{cm}^{-1})$	K	$\nu_{exc}$ (cm <sup>-1</sup>	) $\nu_{ground}$ (cm <sup>-1</sup> )	K	$\nu_{exc}$ (cm <sup>-1</sup> )	$\nu_{ground}$ (cm <sup>-1</sup>	) K
28.82	35.80	-0.216	20.50	32.19	-0.255	17.34	28.93	-0.116	46.35	44.09	1.724
42.56	41.89	1.925	33.05	33.48	0.126	33.36	38.78	-0.196	75.36	71.95	0.160
62.68	69.03	-0.190	45.87	44.26	2.250	42.79	41.85	1.542	78.53	88.42	0.288
67.15	82.34	0.159	66.00	68.41	-0.297	80.11	81.55	0.459	96.14	96.62	-0.182
75.99	89.17	0.337	84.71	95.94	-0.273	100.78	101.67	0.296	102.85	104.69	-0.332
93.75	95.02	0.141	132.00	147.18	-0.454	124.94	124.13	-0.264	126.95	126.53	-0.266
127.44	126.84	0.236	137.22	157.77	-0.124	239.08	243.54	-0.166	136.71	140.03	-0.257
143.04	146.53	-0.159	189.71	193.53	0.317	248.91	254.53	0.173	179.16	178.35	-0.112
185.14	186.01	-0.133	241.89	246.94	-0.249	284.39	283.45	0.398	295.31	291.33	0.396
234.83	241.80	-0.218	298.81	298.40	0.311	377.30	432.37	0.178	339.44	343.20	0.148
247.74	251.85	0.192	309.17	319.74	0.109	430.01	435.29	-0.231	397.17	443.26	0.112
288.13	287.50	0.312	338.25	343.79	-0.124	622.06	638.73	-0.105	455.84	457.90	-0.280
321.86	330.57	-0.149	396.42	440.69	0.406	669.52	681.10	-0.124	527.36	525.25	-0.102
379.47	422.53	0.251	454.13	453.77	-0.524	677.38	689.83	0.211	643.01	645.13	-0.113
431.30	436.42	-0.318	485.39	487.03	0.234	1007.74	1076.80	0.346	710.81	725.09	0.265
464.79	468.74	0.124	505.44	508.48	-0.023	1189.81	1182.88	0.157	717.07	730.71	-0.110
553.43	562.82	-0.146	523.83	518.00	0.107	1234.86	1257.83	-0.122	978.46	1008.43	-0.129
608.14	606.04	-0.106	578.73	582.86	-0.293	1247.11	1281.85	-0.133	1129.06	1113.42	0.133
671.32	680.27	-0.182	628.08	625.70	-0.146	1335.30	1392.52	-0.132	1237.58	1221.12	-0.146
681.96	694.10	-0.180	673.09	701.53	0.211	1471.21	1483.80	-0.156	1299.55	1304.80	-0.154
1036.51	1080.10	0.423	713.30	724.25	0.295				1328.90	1326.55	-0.100
1091.47	1109.40	-0.129	974.59	1010.10	-0.129				1358.08	1369.26	0.182
1187.40	1179.48	-0.236	1121.97	1117.62	0.358				1604.27	1620.18	0.103
1264.60	1274.25	-0.201	1195.50	1194.92	0.238				1833.20	1694.90	0.413
1278.18	1295.86	-0.169	1235.10	1216.53	-0.335						
1316.53	1351.37	-0.117	1264.05	1270.42	0.246						
1341.27	1399.44	-0.208	1329.54	1326.93	0.069						
1461.64	1489.94	0.140	1345.69	1342.32	0.293						
1466.45	1496.96	-0.181	1359.89	1379.70	0.206						
			1471.69	1467.24	0.176						
			1508.61	1496.09	-0.184						
			1521.32	1527.15	0.569						
			1523.97	1533.84	0.268						
			1531.20	1548.54	0.134						
			1540.56	1576.79	-0.380						
			1550.51	1627.73	0.285						
			1574.42	1647.71	0.408						
			1601.41	1673.84	0.558						
			1643.87	1730.65	-0.165						
			2414.75	2448.73	0.114						