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# Accurate Simulation of Optical Properties in Dyes

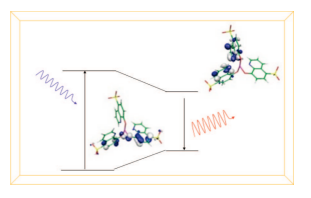
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## CONSPECTUS

**S** ince Antiquity, humans have produced and commercialized dyes. To this day, extraction of natural dyes often requires lengthy and costly procedures. In the 19th century, global markets and new industrial products drove a significant effort to synthesize artificial dyes, characterized by low production costs, huge quantities, and new optical properties (colors). Dyes that encompass classes of molecules absorbing in the UV—visible part of the electromagnetic spectrum now have a wider range of applications, including coloring (textiles, food, paintings), energy production (photovoltaic cells, OLEDs), or pharmaceuticals (diagnostics, drugs). Parallel to the growth in dye applications, researchers have increased their efforts to



design and synthesize new dyes to customize absorption and emission properties. In particular, dyes containing one or more metallic centers allow for the construction of fairly sophisticated systems capable of selectively reacting to light of a given wavelength and behaving as molecular devices (photochemical molecular devices, PMDs).

Theoretical tools able to predict and interpret the excited-state properties of organic and inorganic dyes allow for an efficient screening of photochemical centers. In this Account, we report recent developments defining a quantitative ab initio protocol (based on time-dependent density functional theory) for modeling dye spectral properties. In particular, we discuss the importance of several parameters, such as the methods used for electronic structure calculations, solvent effects, and statistical treatments. In addition, we illustrate the performance of such simulation tools through case studies. We also comment on current weak points of these methods and ways to improve them.

## 1. Introduction

Since Antiquity, dyes have been intensively produced and commercialized by men. Archaeological evidences found for Tyrian purple (TP, an indigo derivative) date its use back to 1900 BC, while Pliny the Elder in his *Naturalis Historia* gives a detailed description of the process of TP extraction from *Murex* sea snails. For many natural dyes still, extraction often required lengthy and costly procedures. The development of a global market and new industrial products induced a significant effort to synthesize artificial dyes, characterized by low production costs, huge quantities, and new optical properties (colors). While in 1914 96% of the indigo world production was of artificial origin, the first synthetic dye (that is, human-designed), mauveine, appeared as early as 1856. Nowadays, dyes encompassing classes of molecules absorbing in the UV—visible part of the electromagnetic spectrum have a wider range of applications, including coloring (e.g., textile, food, painting), energy production (e.g., photovoltaic, OLED), or medical purposes (e.g., diagnosis, drugs).<sup>1</sup>

Parallel to the growing number of applications, a great effort has been spent in the design and synthesis of new dyes, in order to customize absorption and emission properties (wavelengths and intensities). In particular, inorganic dyes containing metallic center(s) have recently become fashionable systems since developments in design, synthesis, and characterization of supramolecular architectures<sup>2,3</sup> nowadays allow for the construction of fairly sophisticated systems capable of selectively reacting to a given external input (light) and behaving as devices at the molecular level (photochemical molecular devices, PMDs).<sup>4</sup>

In this context, theoretical simulations can act at two different levels: the interpretation and the prediction of optical properties. Indeed, a great precision (of the order of half a nanometer) in the simulation of such properties is required for both technological and coloring applications. In conjunction, an accurate prediction often means a rapid screening of photocenters and consequently allows the establishment of structure—property relationships that help in the design of new compounds.

In the present Account, we report recent developments setting up a quantitative ab initio protocol for modeling dye spectral properties. We first discuss the various parameters, such as the method used for the electronic calculations or for solvation effects, as well as the possible statistical treatments, that enter our global scheme. Then some case studies are detailed with the aim of giving a flavor of the actual performance of our protocol. Besides the large part that is devoted to the calculation of the absorption spectra, we briefly comment on the emission (here fluorescence) spectra and give practical examples. Some remarks concerning the current drawbacks of the methods, as well as possible improvements, are finally listed.

## 2. The Computational Approach

Electronic methods constitute undoubtedly the main ingredient for a reliable protocol to evaluate optical properties of dyes, and therefore, the choice of the related computational parameters must be carefully performed to avoid the introduction of significant bias. Environmental effects must be introduced in simulations if a quantitative agreement between theory and experiments is targeted. All these aspects, combined with a statistic treatment, allow definition of an error bar on the calculation results. As a consequence, the protocol is able not only to interpret optical properties of experimentally characterized systems but also to predict the behavior of new chromophores.

**2.1. Electronic Aspects: DFT, TD-DFT, and Atomic Basis Set.** Nowadays, it is well established that the Kohn–Sham approach to density functional theory (DFT) can provide an accurate description of a large number physico-chemical properties for the ground state.<sup>5</sup> Furthermore, the current performance/accuracy ratio tends to exceed that of more sophisticated post-Hatree–Fock (post-HF) approaches.

In contrast, the situation for excited electronic states is often more involved. In fact, from one side, fast and cheap purposely tailored semiempirical approaches lack consistency when applied to families of molecules not included in the original training sets. From the other side, more reliable theoretical tools such as post-HF methods are, in most cases, too expensive to afford the study of the large solvated systems of chemical and industrial interest.

The development of the time-dependent density functional approaches (TD-DFT),<sup>6</sup> including the recent implementation of their analytical energy gradients,<sup>7,8</sup> allows for the extension of the ground-state DFT efficiency to excited states. However, as for the ground state, the choice of the exchange-correlation functional form, which is the only approximated term in DFT, plays a key role in reaching chemically sound results. Concerning the UV-vis absorption spectra simulation, the local density approximation (LDA) tends to strongly underestimate valence excitation energies of most organic molecules, whereas better performances are obtained with the generalized gradient approximations (GGAs) or with meta-GGA approaches. The most accurate values are usually predicted by hybrid functionals, which explicitly include a fraction of HF exchange. Aiming for a classification, three different families of hybrid functionals can be defined. The first is composed by global hybrids (GHs), in which the percentage of HF exchange is constant at each point in space. To the second family belong the local hybrids (LHs), characterized by a mixing of HF exchange that depends on the spatial electronic coordinate. Finally, range-separated hybrids (RSHs) use a growing fraction of exact exchange as the interelectronic distance increases, giving a long-range correction (LC) to the original DFT scheme. The B3LYP,<sup>9</sup> *Lh*-BLYP,<sup>10</sup> and LC-PBE<sup>11</sup> functionals are respective prototypes of these three classes.

Here, we mainly focus on the most widespread GHs and, in particular, on B3LYP and PBE0.<sup>12</sup> On the one hand, B3LYP has de facto become a standard for DFT calculations. On the other hand, we have shown that the PBE0 functional, containing 25% HF exchange, generally outperforms other global hybrids in the computation of UV—vis spectra, so it can be

considered as a reference functional for this property (see below). Furthermore PBEO can be used as a "multipurpose" approach since it yields appropriate simulations for many other chemicophysical properties, thus providing a theoretically consistent description of both ground- and excited-state properties.<sup>12,13</sup>

Finally, it is worth mentioning that the choice of the basis set also plays a relevant role, even though TD-DFT appears to be less basis set dependent than post-HF methods. In our experience, a medium size basis set like valence double- or triple- $\zeta$  basis already gives converged results for valence transitions when both polarization and diffuse functions are added. In particular, the Pople's 6-311+G(2d,p) provides converged transition energies of low-lying states for the majority of investigated dyes, while the compact 6-31+G(d) basis represents a valuable compromise between accuracy and computational speed. Concerning metal atoms, the basis set effect seems even less pronounced so that standard pseudopotentials, such as those developed by Hay and Wadt, are often found to be well suited. However, we have shown that when higher energy states (e.g., Rydberg) are sought, larger basis sets, including very diffuse functions, are mandatory.<sup>14</sup>

2.2. Modeling Solvent Effects. Electronic transitions energies are affected by the surroundings, inclusion of which rapidly appears to be crucial for their quantitative simulations, especially if polar groups are directly exposed to the solvent. However, calculations performed on isolated molecules can often provide qualitative trends. The magnitude of the solventinduced shift strongly depends not only on the nature of the solvent (polar or not, protic or not) but also on the nature of the excitation under investigation. In organic molecules,  $n-\pi^*$ transitions are more affected than  $\pi - \pi^*$  transitions because the variation of the electronic distribution is larger in the former case; it is also enhanced in protic solvents, which obviously tend to interact with the lone pair. Several reliable approaches have been developed for solvent modeling in ab initio simulations. Accurate simulation of bulk effects for both nonpolar and polar solvents can be obtained by polarizable continuum models (PCMs),<sup>15</sup> possibly including nonequilibrium effects in TD-DFT calculations.<sup>16</sup> More problematic is the simulation of protic solvents, where specific H-bonding interactions can modify the electronic structure of the solute. In that case, the best compromise is represented by mixed models, where all molecules involved in the interactions (e.g., the first solvation shell) are explicitly considered in the electronic calculations and then embedded in a PCM to simulate the bulk effect. A precise determination of the orientation of the solvent molecules has usually a negligible effect with respect to their inclusion and can be simply determined by standard structure minimization procedures without going for more expensive dynamic approaches.

**2.3. Defining an Error Bar: The Role of a Statistical Treatment.** To improve the agreement between theory and experiment, the results of several approaches can be advantageously combined. To obtain the most efficient combination, multiple linear regression (MLR), based on the numerical technique of least-squares fitting and analyzing the relationship between one dependent variable (experimental value) and *p* independent variables (theoretical values), is a method of choice. To test the significance of a regression curve, the total sum of squares (MSS) and the residual sum of squares (RSS)

$$TSS = MSS + RSS$$
(1)

$$\sum_{i=1}^{n} [y_i - \bar{y}]^2 = \sum_{i=1}^{n} [y_i(x_i) - \bar{y}]^2 + \sum_{i=1}^{n} [y_i - y_i(x_i)]^2$$
(2)

with *n* being the number of points (here dyes) considered,  $y_i$  the experimental quantities,  $y(x_i)$  the regression data, and  $\bar{y}$  the average values (here transition energies or wavelengths). If the fitted curve passes through all the original data points, MSS is equal to TSS, and RSS is zero. This can be the case if there are numerous descriptors (*p*) and only a few data points (*n*). Therefore, one uses an adjusted correlation coefficient ( $R_{adi}^2$ ):

$$R_{adj}^{2} = 1 - \frac{n-1}{n-p-1} \left( 1 - \frac{RSS}{MSS} \right)$$
 (3)

Reliability limits for the regression parameters, measuring the adequacy of each independent variable in the model are also determined, allowing to step-by-step removoval of the less significant independent variables. MLR provides not only the usual mean absolute error (MAE) but also a standard deviation,  $d_{R}$ :

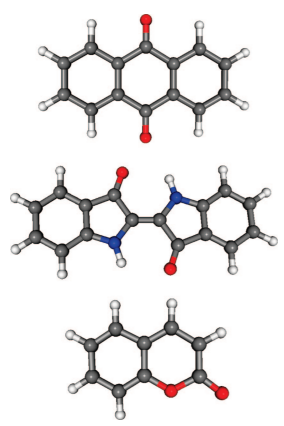
$$d_{\rm R} = \sqrt{\frac{\rm RSS}{n-p-1}} \tag{4}$$

which is useful for the prediction of properties of compounds not included in the training set.

#### 3. Cases Studies

In the past few years, we have studied a large number of dyes, including both organic and inorganic compounds. Here, we will report only representative examples of transition types both in organic molecules and in coordination complexes.

**3.1. Absorption of Organic Molecules.** 9,10-An-thraquinone (AQ) derivatives (Figure 1) represent about 1/3 of



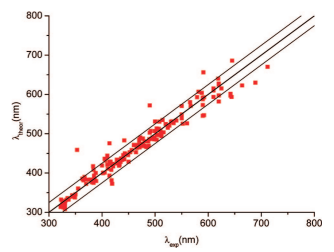
**FIGURE 1.** Sketches of the basic organic dyes investigated. From top to bottom: 9,10-anthraquinone; prototypical indigo; prototypical coumarin.

today's world organic dye production. This success originates in the stability, easy synthesis, and versatile applications of AQs.<sup>1</sup> In a series of works,<sup>17–20</sup> we have shown that combining the results obtained with the B3LYP and PBE0 functionals provides very accurate estimates of the  $\lambda_{max}$ . In general, the B3LYP hybrid tends to be more efficient for long-wavelength AQ derivatives that typically display a strong intramolecular hydrogen bond between the carbonyl and the side amino groups, whereas PBEO works best for the hydroxy-AQs.<sup>17,18</sup> It was also shown that extending the basis set further than 6-31G(d,p) is useless;<sup>18,21</sup> that is, in this peculiar case, diffuse functions do not provide more accurate (before fitting) or consistent (after fitting) transition energies. Using all the data from refs 18-20, we designed a set of 189 AQ derivatives solvated in various media (CH<sub>2</sub>Cl<sub>2</sub>, MeOH, and EtOH), including a large panel of standard substitutions as well as complex laser dyes. Prior to fitting, the mean absolute deviations (MAEs) are 0.10 eV for both functionals. It is worth pointing out that such an accuracy exceeds the generally accepted error for TD-DFT calculations (gas-phase = 0.3 eV) and, of course, that obtained with a CIS approach, 0.58 eV.<sup>21</sup> Optimizing the MLR coefficients for this 189-AQ set leads to

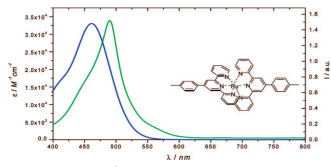
$$\lambda_{\max}^{\exp} = -33.48 - 0.6845 \lambda_{\max}^{B3LYP} + 1.80535 \lambda_{\max}^{PBE0}$$
(5)

which provides average deviations limited to 0.08 eV, that is, 20% better than the raw B3LYP or PBE0 values. From the comparison of simulated and experimental wavelengths of Figure 2, it is indeed obvious that there is an excellent agreement between theory and experiment. Indeed, less than 15% of the cases present unacceptable deviations (>0.15 eV). This means that the absorption spectra of substituted AQ can be accurately foreseen at an affordable computational cost.

Another striking example of the efficiency of TD-DFT for simulating the absorption spectra of organic dyes is provided by the indigoid derivatives.<sup>22–24</sup> From a methodological point of view, indigoids are the typical systems for which a second set of polarization functions is required to reach  $\lambda_{max}$  convergence. Studies have also clearly demonstrated that B3LYP is to be favored over PBEO only when at least one internal NH-O=C hydrogen bond is formed.<sup>24</sup> Using the PCM-TD-DFT/6-311+G(2d,p)//PCM-DFT/6-311G(d,p) approach, we have determined a  $\lambda_{max}$  of 304 indigoid derivatives, including compounds with several heteroatoms (indigo, thioindigo, selenoindigo), various substitutions on the outer phenyl rings (Tyrian purple, nitro derivatives), differently linked (indirubin, isoindigo), or conformers (cis/trans), and solvated in a wide panel of solvents. For this large set, which covers the full width of the visible part of the electromagnetic spectrum, 22-24 we obtain a MAE limited to 0.04 eV, which is basically 1 order of magnitude smaller than the expected 0.3 eV average deviation, although absolutely no statistical correction was performed. Because similar accuracy levels have been found for



**FIGURE 2.** Comparison between experimental and theoretical absorption wavelengths for 189 AQ dyes. The theoretical values are obtained using the MLR definition on the basis of PCM-TD-(B3LYP/PBE0)/6-31G(d,p)//B3LYP/6-31G(d,p) raw data. The central line indicates a perfect theory/experiment match, whereas the two side lines are boundaries for  $\pm 25$  nm deviation.

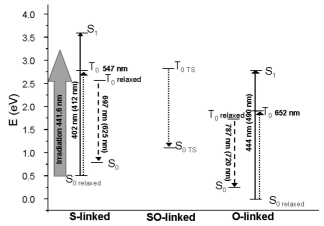


**FIGURE 3.** Calculated (blue, full width at half-maximum, fwhm = 0.1 eV) and experimental (green) absorption spectra of a typical Ru(II)–polypyridine complex.

many other dyes,<sup>25</sup> we can conclude that our computational protocol is reliable for quantitative color estimates of organic derivatives.

3.2. Absorption of Inorganic Dyes. There are few systematic studies dealing with the effect of the computational protocol on the modeling of given excited states in inorganic dyes. This is mainly due to their size (in terms of atom numbers and active electrons) and, contrary to organic dyes, to the difficulty to obtain large sets of similar compounds. Focusing on inorganic dyes including d transition metal atoms, the TD-DFT approach, in conjunction with the PBEO functional, well reproduces the metal to ligand charge transfer bands (MLCT) usually responsible for their strong absorption in the visible region. Illustrative examples are the complexes of the Ru-polypyridine family for which the experimental transitions and the band shape are both nicely reproduced (see Figures 3 and 4).<sup>26-32</sup> However, the errors are larger than for organic molecules, ranging between 0.15 and 0.40 eV. Furthermore, it is a matter of fact that for such complex systems, associated experimental facts are seldom straightforwardly rationalized. In this context, theoretical methods again constitute a valuable means to interpret and even anticipate photochemical behaviors.<sup>26,30,32,33</sup> A clear example is the phototriggered isomerization (PTI) process undergone by the [Ru(bpy)(tpy) $dmso]^{2+}$  complex (bpy = 2,2'-bipyridine; dmso = dimethyl sulfoxide) for which theory has amended and anticipated the experimental evidence of the pathway of photoisomerization. In this system, the sixth position on the octahedral coordination sphere of Ru(II) is constituted by dmso that can either bind to the sulfur atom (S-linked isomer) or to the oxygen (O-linked isomer).

The S-linked isomer of [Ru(bpy)(tpy)dmso](CF<sub>3</sub>SO<sub>3</sub>)<sub>2</sub> undergoes an immediate color change upon irradiation at 441.6 nm: the absorption is shifted from 412 to 490 nm, and the corresponding photoproduct is stable for days. Upon excitation, the complex also displays a weak luminescence ( $\lambda_{em} =$ 

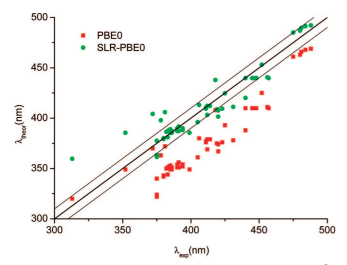


**FIGURE 4.** Schematic representation of the computed (TD-DFT) relative energy of the ground ( $S_0$ ) and first singlet ( $S_1$ ), and first triplet ( $T_1$ ) excited states of several conformers of [Ru(bpy)(tpy)dmso]<sup>2+</sup> and computed absorption/emission energies. Full line arrows correspond to computed allowed transitions, dotted arrows to nonallowed ones, and dashed arrows to the computed emission. Experimental values are reported in parenthesis; *relaxed* stands for computed at the corresponding optimized structure, and TS indicates a transition state.

720 nm) at room temperature, while a new additional emission feature appears at 625 nm upon cooling to 170 K.<sup>34</sup> These observations lead to the proposal of a rather complicated reaction path, where the isomerization mechanism takes place in the lowest triplet or singlet excited states and involves a hypothetical  $\eta_2$  species, characterized by concomitant S- and O- linkages to the metal cation. This  $\eta_2$  structure was supposed to be responsible for the weak emission at 625 nm.<sup>34</sup> The theoretical reaction pathway clearly appears different.<sup>35</sup> Indeed, while the  $S_0 \rightarrow S_1$  absorption of both S- and O-linked species, as well as the emission from the O-linked triplet state, are correctly reproduced, no evidence of the formation of the a stable  $\eta_2$  species can be found. Both in the ground and in triplet excited states,  $\eta_2$  truly corresponds to a transition state (TS), see Figure 4. Furthermore, the emission at 625 nm could be attributed to the decay of the triplet state of the S-bonded isomer computed at 697 nm.

This apparent discrepancy was finally conciliated when new experimental data<sup>36</sup> confirmed the validity of the theoretical outcomes, pointing out that the reaction actually proceeds within a direct one-step mechanism, and as predicted by theory, the SO-linked  $\eta_2$  species is the TS. This example, though based on a quite simple light triggered phenomenon, clearly illustrates the essential role that computational approaches can play in the interpretation of complex photoinduced phenomena.

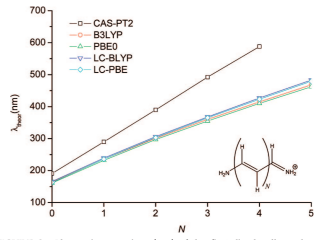
**3.3. Fluorescence.** The theoretical framework allowing for the computation of geometric gradients at the TD-DFT level<sup>8</sup>



**FIGURE 5.** Comparison between theoretical and experimental  $\lambda_{max}^{fluo}$  of substituted coumarins. The central line indicates a perfect theory/experiment match, whereas the two side lines are boundaries for  $\pm 10$  nm deviation.

has paved the way toward the calculation of the excited-state structures of large molecules, hence, the determination of fluorescence. Nevertheless, solvent effects in a PCM framework were only recently included in a TD-DFT approach.<sup>9</sup> Consequently, the number of works assessing the efficiency of TD-DFT for the calculation of fluorescence wavelengths of a statistically significant set of dyes is still limited, because the largest investigations were performed for neutral and charged coumarins (Figure 1),<sup>37,38</sup> for which PCM-TD-DFT correctly reproduces the modifications induced by auxochromic effects for the emission properties. Indeed, a simple linear regression applied to PCM-TD-DFT results allows one to obtain a perfect match with measurements whatever the form of the coumarin dye (enol, keto, cation, or anion). This approach returns absolute estimates with very small average errors (0.07 eV, Figure 5). More importantly, the absorptionfluorescence Stokes shift for a given compound is only slightly underestimated. Furthermore, the direct TD-DFT optimization of the excited-state structure allows one to avoid the use of CIS structures to estimate fluorescence wavelengths. This circumvents a problematic approximation as electron-correlation effects differ for the ground and excited state. In addition, the symmetry of the excited-state geometries and the solvatochromic effects appears to be not always consistently reproduced by CIS.38

In order to test the performance of TD-DFT for the description of the fluorescence properties of inorganic dye, the spectral properties of an Al(III) complex of 8-hydroxyquinoline-5sulfonic acid (Al(III)(8-HQS)<sub>3</sub>) were computed.<sup>39</sup> This latter was considered due to its strong fluorescence both in solution and in solid state, making it a reference compound for applica-



**FIGURE 6.** Absorption maxima (nm) of the first dipole-allowed electronic transition as a function of the chain length (*N*) in model cyanines.

tion as emissive material in the emerging display technology based on organic light-emitting diodes (OLEDs).<sup>40</sup> Ligand-centered  $\pi - \pi^*$  transitions of HOMO–LUMO type are responsible for the absorption of the complex in the near-UV. Its fluorescence ( $S_1 \rightarrow S_0$ ), observed between 475 and 485 nm,<sup>41</sup> is computed at 488 nm, in excellent agreement with the experimental data. Indeed, calculations not only nicely reproduce the measured bands and Stoke shift but also give access to the geometrical structure of the excited state. In particular, the analysis of the geometrical parameters corresponding to the optimized  $S_1$  structure validates the experimental assumption that within the complex the AI cation plays the same role as a proton for the isolated ligand, the structural change in the complex being similar to those obtained for the photoinduced tautomerization observed for the isolated ligand.

#### 4. Breakdowns and Perspectives

The computational protocols above-described can generally be applied to the calculation of optical properties of dyes. However, it is clear that when dealing with TD-DFT approaches, the recognized failures of DFT for the ground state due to the use of approximate functionals (such as the lack of nondynamical electronic correlation or the self-interaction error) sum up to others specific to the excited states model (LDA kernel, missing of double excitations, response treatment limited to linear term, lack of the 1/*r* asymptotical behavior). From a practical point of view, the two most problematic cases are represented by systems characterized by multireference electronic states or when dealing with transitions of charge transfer (CT) type.

An example of the former breakdown can be illustrated by the cyanine derivatives (Figure 6), for which standard DFT

		without fit				with linear correction	
family	method	MSE	MAE	rms	$R^2$	MAE	rms
AQ	HF	127	127	132	0.97	11	14
	PBE	-71	74	80	0.78	28	36
	PBEO	12	19	27	0.96	12	15
	LC-PBE <sup>b</sup>	67	67	71	0.98	8	10
	LC- $\omega$ PBE <sup>b</sup>	85	85	89	0.99	8	9
	CAM-B3LYP	53	33	58	0.98	8	10
AB	HF	64	64	75	0.89	15	21
	PBE	-90	90	103	0.83	22	27
	PBEO	-20	25	27	0.93	12	17
	LC-PBE	33	33	43	0.95	10	14
	LC- $\omega$ PBE	46	46	54	0.96	10	14
	CAM-B3LYP	15	20	28	0.94	10	16
IG	HF	170	170	174	0.89	19	26
	PBE	-96	96	116	0.71	34	42
	PBEO	6	19	23	0.93	17	21
	LC-PBE	70	70	72	0.97	10	13
	LC- $\omega$ PBE	99	99	101	0.97	12	14
	CAM-B3LYP	58	58	60	0.98	10	12
all	HF	116	116	127	0.76	37	45
	PBE	-86	87	102	0.81	30	40
	PBEO	-3	22	29	0.91	22	28
	LC-PBE	52	52	58	0.94	18	23
	LC- $\omega$ PBE	71	71	78	0.94	18	23
	CAM-B3LYP	37	38	46	0 0.93	20	25

**TABLE 1.** Statistical Analysis for 9,10-Anthraquinone (AQ), Azobenzene (AB), and Indigoid (IG) Dye Series<sup>a</sup>

<sup>*a*</sup> All data are in nanometers and are from ref 25. <sup>*b*</sup> Please note that LC-PBE and LC- $\omega$ PBE differ both in how the short-range contribution is derived and in the range separation parameter,  $\omega$  (0.33 b<sup>-1</sup> in LC-PBE and 0.40 b<sup>-1</sup> in LC- $\omega$ PBE).<sup>44</sup>

approaches fail in predicting the evolution of  $\lambda_{max}$  with increasing chain length.<sup>42</sup>

The CT problem was nicely underlined in model systems, where the electron donor and acceptor are fully electronically decoupled.<sup>43</sup> In such conditions (zero overlap between the two moieties), both GGA and GH approaches significantly underestimate the electronic transition energies. This is no longer the case for RSH methods, such as LC- $\omega$ PBE<sup>44</sup> or CAM-B3LYP,<sup>45</sup> for which the variation (but not always the absolute value) of the CT bands with respect to the donor-acceptor distance can be adequately reproduced.<sup>46</sup> However, if the two parts have an even minimal orbital overlap, CT transitions can correctly be reproduced even using standard GH, as demonstrated by several studies on coordination chemistry compounds<sup>26,30,33</sup> and on typical organic dyes.<sup>47</sup> More generally, RSH models represent a further step toward a greater accuracy,<sup>48</sup> even if the improvement with respect to the PBEO results is not systematic, as nicely summarized in the data of Table 1, where the  $\lambda_{max}$  of several families of organic dyes are reported.25

Finally, it is worth mentioning that if a calculation protocol for vertical transition energies in organic dyes can be well defined, still in order to get a realistic simulation of absorption spectra (band shape and observed transition energies) of even very simple systems, another ingredient is still missing: the calculation of vibronic effects. Recent advances in the calculation of Franck–Condon factors, also feasible for relatively medium-sized compounds, pave the way toward even more realistic computation of dye spectra.<sup>49,50</sup>

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#### **BIOGRAPHICAL INFORMATION**

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