First Principles Design of Dye Molecules with Ullazine Donor for Dye Sensitized Solar Cells

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ABSTRACT: We design a series of metal-free donor- π -bridge molecules (denoted VB0–VB4) based on a new donor group ullazine donor—as sensitizers for dye sensitized solar cell (DSSC) applications. Density functional theory (DFT) and timedependent DFT calculations reveal that the physical properties of dyes, including spectral response, light harvesting efficiency, and electron injection rate, are systematically improved by combining ullazine donor to a series of length changing π bridges. Dye VB2 is the best candidate thanks to its outstanding performance on key parameters and achieving a balance between competing factors. Compared to two other series of molecules—L and M dyes, which differ from VB dyes by only the donor group—VB dyes have the largest light harvesting efficiency and the largest number of electrons injected to the conduction band of TiO₂. These results suggest that the ullazine group can serve as an excellent donor for future DSSC applications.

INTRODUCTION

Since the enlightening paper published in Nature by O'Regan and Grätzel in 1991,¹ dye-sensitized solar cells (DSSCs) have drawn comprehensive attention as one of the most promising renewable energy devices. Typical DSSCs are composed of dye molecules anchored on nanocrystalline TiO2 deposited on transparent conducting oxide glass, platinized counter electrode, and redox electrolyte. The highest energy conversion efficiency of 12.3% under AM 1.5 irradiation has been achieved.² DSSCs based on conventional Ru-based chromophores such as N3/N719³ and the black dyes⁴ have been investigated intensively. However, heavy metal ions employed in metal-based dyes are scarce in nature and not environmentally friendly. Meanwhile, the synthesis and purification processes of these dyes are complicated, which prevent them from large-scale fabrication. Fortunately, metal-free organic dyes provide a prospective alternative to Ru-based dyes for their benign environmental effects, high molecular extinction coefficients, and cheap preparation processes.⁵ These advantages meet the technical requirements toward utilizing a thinner TiO₂ layer to reduce energy loss during electron transport, making metal-free organic dyes ideal candidates in DSSC devices.

Most metal-free organic dyes adopt a donor- π -acceptor (D- π -A) structure, with the acceptor (A) ligand anchored on the semiconductor surface. This structure will favor efficient charge

transfer of excited electrons to the conduction band of semiconductor and the regeneration of excited dyes to the ground state by the redox shuttle. Arylamine and 2-cyanoacrylic acid are the typical donor and acceptor groups employed, respectively.⁶ To reach high energy conversion efficiency, the dyes must meet a series of severe requirements.⁷ There should be a large driving force to ensure efficient electron injection, and the dyes' ground state oxidation potential should be higher than that of the redox couple in electrolyte to guarantee fast regeneration of the oxidized dyes. The dye acceptor, also acting as the anchor to semiconductor substrate, should be strongly adsorbed to provide stable charge transfer channels and fast electron injection. Finally, the bridge group is carefully selected to maximize sunlight absorption by shifting the spectrum from UV light to visible and near-infrared light⁸ and to facilitate charge separation upon photoexcitation.

Chemical modification on individual dye units is an effective way to tune the redox potential and absorption properties of D- π -A organic dyes.^{5,9,10} For instance, by changing different π spacers and electron acceptors, a red-shift in absorption spectra and significant differences in the redox potential are achieved for triphenylamine dyes.¹⁰ Theoretical methods based on

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Figure 1. The donor units, R, of the three series of dyes (VB, L, M) are shown in the upper row. With R replaced by R(VB) in the lower row we could get VB0, VB1, ..., VB4 dyes.

density functional theory (DFT) and time-dependent DFT (TDDFT) calculations were also employed to rationally optimize optical and electrochemical properties of dye molecules.^{8,11–13} Examples include identifying dye structures by optimizing the π spacer⁸ and the acceptor group of dyes.¹¹ However, despite all these extensive works, there were few studies on the effects of systematic modifications of the donor group. The geometry, electrophilicity, and polarizability will directly influence the dye adsorption, spectra response, and electrodynamics. In this work we will investigate the influence of different donor groups on the performance of DSSCs by systematically evaluating the key DSSC factors determining solar cell efficiency. Rational design of dye structures based on first-principles can significantly facilitate test and development of new dyes, reducing the cost and time consumed in conventional trial-and-error experimental approaches.^{14,15}

By taking advantages of theoretical approaches, we investigate an electron-rich molecular unit, ullazine donor, which has been recently synthesized experimentally¹⁶ (structure shown in Figure 1), to be used as part of dye sensitizers for DSSC application. The nitrogen-containing heterocycle of ullazine group (Figure 1, top left) with 16 π -electrons will facilitate a strong intramolecular charge transfer. It could serve as both an electron donor and an acceptor under different conditions, which indicates a good potential for future optoelectronic applications.¹⁶ The ullazine group has a large planar structure which will benefit patterning on the surfaces. The structure is relatively simple that could be easily manipulated in experiments and modeled in first principles calculations. The resultant dyes with ullazine donor, denoted as VB series of dyes, show near-red optical absorption and high extinction coefficients. We expect this group of dyes with ullazine donor act as high efficient candidates for DSSCs.

To systematically tune the dye's photoabsorption and electrochemical properties, we choose 2-cyanoacrylic acid as electron acceptor and anchoring group and alkene/thiophene fragments as the π spacer. By inserting alkene and thiophene between the donor and the acceptor groups, we can gradually change the energy levels of the highest occupied molecule orbital (HOMO) and the lowest unoccupied molecular orbital (LUMO) of the sensitizer and extend major absorption peak from the UV-light region to near-infrared. This results in a series of dye molecules VB0, VB1, ..., VB4, with distinct optical and electrochemical characters (Figure 1). The dye's structure and performance are optimized in terms of achieving a balance among competing factors such as light harvesting, spectral response, electronic driving force, and open circuit voltage for DSSC applications. For comparison, we take L dyes with a diphenylaminophenyl donor (R(L) in Figure 1), which were intensively studied in the literature,¹⁷ and M dyes with a dimethylaminophenyl donor (R(M) in Figure 1), as control groups. Both L dyes and M dyes own the same π spacer and acceptor as VB dyes--they only differ by the donor group. We found that VB dyes have higher extinction coefficients, and the number of electrons injected into the semiconductor conduction band is almost 2 times larger for VB dyes than for L or M dyes, resulting from stronger electronic interactions between VB molecules and the TiO₂ surface.

METHODS

The sunlight-to-electricity conversion efficiency (η) of solar cell devices is determined by the open-circuit photovoltage (V_{OC}), short-circuit current density (J_{SC}), and the fill factor (FF), as compared to incident solar power (P_{inc}):

$$\eta = FF \frac{V_{\rm OC} J_{\rm SC}}{P_{\rm inc}}$$
(1)

In DSSCs, we can calculate the $V_{\rm OC}$ as¹⁷

The Journal of Physical Chemistry C

$$V_{\rm OC} = \frac{E_{\rm C} + \Delta {\rm CB}}{q} + \frac{kT}{q} \ln \left(\frac{n_{\rm c}}{N_{\rm CB}}\right) - \frac{E_{\rm redox}}{q}$$
(2)

Here q is the unit charge, E_c is the conduction band edge (CBE) of the semiconductor substrate, k is the Boltzmann constant, T is the absolute temperature, n_c is the number of electrons in the conduction band, $N_{\rm CB}$ is the density of accessible states in the conduction band,⁸ and $E_{\rm redox}$ is the reduction—oxidation potential of electrolyte. Δ CB is the shift of E_c when the dyes are adsorbed on substrate and can be expressed as¹⁸

$$\Delta CB = \frac{q\mu_{\text{normal}}\gamma}{\varepsilon_0\varepsilon}$$
(3)

In this expression, $\mu_{\rm normal}$ denotes the dipole moment of individual dye molecules perpendicular to the surface of the semiconductor substrate and γ is the dyes' surface concentration. ε_0 and ε represent the vacuum permittivity and the dielectric permittivity, respectively. It is obvious from eqs 2 and 3 that $n_{\rm c}$ and $\mu_{\rm normal}$ exert crucial influences on $V_{\rm OC}$.

The J_{SC} in DSSCs is determined by the following equation:⁸

$$J_{\rm SC} = \int LHE(\lambda) \Phi_{\rm inject} \eta_{\rm collect} \, d\lambda \tag{4}$$

where LHE(λ) is the light harvesting efficiency at a given wavelength, Φ_{inject} evinces the electron injection efficiency, and $\eta_{collect}$ denotes the charge collection efficiency. In the systems which are only different in sensitizers, $\eta_{collect}$ can be reasonably assumed to be constant. LHE(λ) can be calculated with

$$LHE = 1 - 10^{-f}$$
(5)

where f represents the oscillator strength of adsorbed dye molecules. Φ_{inject} is related to the driving force ΔG_{inject} of electrons injecting from the excited states of dye molecules to the semiconductor substrate. It can be estimated as¹⁹

$$\Delta G_{\rm inject} = E^{\rm dye*} - E_{\rm CB} = E^{\rm dye} + E_{0-0} - E_{\rm CB} \tag{6}$$

where $E^{\text{dye}*}$ is the oxidation potential of the excited dye, E^{dye} is the redox potential of the ground state of the dye, E_{0-0} is the vertical transition energy, and E_{CB} is the conduction band edge of the semiconductor. So J_{SC} can be well estimated through fand ΔG_{inject} .

On the basis of eqs 1-6, we could roughly predict the efficiency of novel dyes without intensive calculations. More insightful theoretical estimations could be achieved using DFTbased real-time evolution of electron dynamics as in our previous works.^{20,21} However, the calculations are rather expensive (with thousands of CPU hours per trajectory). We believe the present approach will be interesting and valuable for assisting experimentalists in searching for high performance dyes. All the calculations are performed based on DFT and TDDFT.²² Ground-state geometry optimization is carried out with SIESTA code. During the optimization, we use Troullier-Martins pseudopotentials²³ to represent the atomic cores, generalized gradient approximation (GGA) as exchange-correlation functional,²⁴ a double- ξ plus polarization basis sets of localized orbitals with an energy cutoff of 100 Ry, and the optimization will stop when the force on each atom is smaller than 0.02 eV/Å in magnitude. We also use the Gaussian09 program package to simulate the absorption spectra of all dyes in vacuum with CAM-B3LYP exchange-correlation functional and the energy levels of the dyes' molecule orbitals with B3LYP

exchange-correlation functional. The 6-31G(d) basis sets were used for C, H, O, N, and S. This set of parameters is illustrated to produce nice agreements with experiment.¹¹

RESULTS

Structure and Properties of VB Dyes. The optimized structure of VB series of dyes is shown in Figure 1. Dye VB0 is composed by ullazine donor directly connected to the cyanoacrylic acid acceptor group. Dye VB1 has an additional thiophene group inserted as a linker. The linker group in dye VB2 is further elongated by adding an alkene group. Dyes VB3 and VB4 were formed by extending the π spacer by another thiophene group or by a thiophene and a new alkene group, respectively. This series of VB dyes, from VB0 to VB4, are gradually elongated by extending the π spacer.

To demonstrate their characteristic electronic structure, we choose three representative VB dyes—VB0, VB2, and VB4— and plot the distribution of molecular orbital wave functions in Figure 2. Wave functions of VB1 and VB3 dyes have similar



Figure 2. Frontier molecular orbitals of VB0, VB2, and VB4 dyes. A typical $D-\pi$ -A character is shown.

features. As expected, the distribution of molecular orbitals of VB dyes show a strong D- π -A character; namely, the HOMO orbitals are localized on the donor part while the LUMO orbitals are mainly localized on the cyanoacrylic acceptor group of the molecule. The HOMO-1 orbitals of all dyes are more or less delocalized over the entire molecule. The π -characters of these orbitals will contribute to high extinction coefficients of the VB series of dyes. The D- π -A charge distribution favors a strong electron-hole separation on TiO₂ surface, which is

Chemical modifications in the linker group would tune photoabsorption of VB dyes, since it is well-known that longer π linkers favor red-shifted absorption in D- π -A dyes. We then calculate the UV-vis absorption spectrum of VB dyes, shown in Figure 3, to analyze one of the key factors in DSSC, LHE.



Figure 3. Calculated absorption spectra of VB dyes.

The extinction coefficient is evaluated following the procedures in ref 11. Table 1 shows that the wavelength for maximum absorption, λ_{max} is shifted from ultraviolet range (315.3 nm for VB0) to blue-green range (501.8 nm for VB4) as the linker gets longer. The red-shifted absorption spectrum matches better solar spectrum, which has a maximal intensity at ~500 nm. Higher efficiencies could be achieved with dyes showing more red-shifted light absorbance. Experiments also reveal that dyes (JD21) with the acceptor group linked to the side position of ullazine (site 5 instead of site 6) yield additional merits with maximum absorption at 582 nm and light conversion efficiency of 8%.¹⁶ Our calculations reproduce the absorption spectrum nicely. In this work, we will focus on the effect of substituting the donor with novel ullazine groups. Results on systematic modifications of ullazine dyes with acceptors at different positions will be presented elsewhere. The oscillator strength of all the five dyes is large and increases with the length of the π spacer, and so does the LHE. The calculated LHEs are all near unity. The longest dye VB4 has the largest oscillator strength and LHE. TDDFT with traditional exchange-correlation functionals tends to underestimate the excitation energy especially for charge transfer transitions of molecules. The errors increase with the decrease in overlap between the orbitals of the ground and excited states. Hybrid functional will reduce the errors, and with long-range corrections the errors are further decreased to a level of ~0.5 eV, making the discrepancy between experiment and theory less dependent on the coupling degrees of initial and final states in optical transitions.²⁵ It should be noticed that in VB dyes the main contribution to the first absorption peak is assigned to a charge transfer (CT) transition from HOMO-1 to LUMO, which is different from most D- π -A organic dye molecules, where electron excitation from HOMO to LUMO plays a critical role. This is caused by the symmetry of the molecular orbitals: strong separation between the HOMO and LUMO levels in VB dyes makes their spatial overlap very small, resulting in a negligible oscillator strength <0.03 for HOMO \rightarrow LUMO electronic transitions.

We note that ullazine dyes have been successfully synthesized in experiment recently¹⁶ after we have carried out theoretical characterizations of VB dyes. An ullazine dye, JD32, which only slightly differ from the VB1 dye (phenylhexyloxy group in JD32 versus phenyl group in VB1 on the peripheral of ullazine donor) was reported 16 with a major absorption peak at 393 nm and a small shoulder at 540 nm. For comparison, we predicted that the dye have maximum adsorption at 376 nm (oscillator strength = 1.49) and a very small shoulder at \sim 500 nm (oscillator strength = 0.01, small shoulders are not listed in Table 1), in consistence with experiment. The small difference is attributed to structural difference in the donor as well as computational errors in DFT exchange-correlation functionals. Solar cells fabricated with JD32 (VB1) dyes present acceptable light-to-electricity efficiency of 2%. Higher efficiencies will be achieved with dyes with more red-shifted light absorbance. Experiments also reveal that dyes (JD21) with the acceptor group linked to the side position of ullazine (site 5 instead of site 6) yield additional merits with maximum absorption at 582 nm and light conversion efficiency of 8%. Our calculations reproduce the absorption spectrum of these dyes nicely. Results on systematic modifications of this group of ullazine dyes with acceptors at the different position will be presented elsewhere.

Figure 4 shows the calculated energy levels of the VB dyes. As the linker getting longer the HOMO-1 are upshifted for



Figure 4. Calculated energy levels of molecular orbitals of VB dyes. The red bars denote unoccupied orbitals, and blue ones denote occupied orbitals.

almost 1 eV and the LUMO downshift for about 0.5 eV. This is in agreement with the increase in the λ_{max} . On the other hand,

Table 1. Calculated Maximum Absorption Wavelength (λ_{max}), Oscillator Strengths (f), Light Harvesting Efficiency (LHE), and the Corresponding Electronic Transitions for VB Dyes

dye	λ_{\max} (nm)	f	LHE (λ)	assignment
VB0	315.3	1.39	0.959	H-1 \rightarrow L(67%) H \rightarrow L+1(16%) H-2 \rightarrow L(9%)
VB1	375.8	1.49	0.968	H-1 \rightarrow L(69%) H-2 \rightarrow L(13%) H \rightarrow L+2(9%)
VB2	428.6	1.83	0.985	$H-1 \rightarrow L(88\%) H-2 \rightarrow L(7\%)$
VB3	472.6	2.12	0.992	$\text{H-1} \rightarrow \text{L(87\%)}$
VB4	501.8	2.51	0.997	$H-1 \rightarrow L(85\%)$



Figure 5. Two adsorption configurations of dyes on TiO_2 (101) surface: N^O adsorption (left) and O^O adsorption (right).

the HOMO level does not shift much. This could be understood since the HOMO is strongly localized on the donor part and does not change its characteristics as the π spacer elongates. Some may argue that after photoexcitation from the HOMO-1 to LUMO level the presence of higher lying HOMO may deteriorate DSSC performance. We note that the energy level of HOMO orbitals, about -5.0 eV, is sufficiently lower than the redox potential of I^-/I_3^- electrolyte at -4.6 eV.²⁶ In addition, this character could provide an advantage to separately optimize the optical absorption and the regeneration potential of dye sensitizers. Ru-based high efficiency dyes, for instance C101 and C106 with efficiency larger than 11%, have similar HOMO levels (-5.34 and -5.05 eV, respectively). Long-term stability under moderate thermal stress and visiblelight soaking has been proved in experiments.²⁷ Some metalfree dyes also show a HOMO level in the range of -5.1 to -5.2eV and present high efficiencies up to 9%.28

Dye Adsorption onto TiO₂ Surfaces. On the basis of the knowledge of isolated dyes, we extend to study the energy alignment, electronic coupling, and interface dipole moment of these dyes when adsorbed onto TiO₂ substrate to analyze other factors affecting the energy conversion efficiency. TiO₂ anatase (101) surface is modeled using a slab with a thickness of six atomic layers. Periodic boundary conditions are employed, with a surface supercell of 10.24×22.70 Å² and a vacuum layer of at least 10 Å to prevent the interactions between adjacent slabs. Two adsorption configurations are considered (see Figure 5). The N^AO adsorption configuration has the C-N and one of the two C–O groups in cyanoacrylic carboxylic acid binding to the surface Ti atoms. The H in carboxylic acid forms a hydrogen bond with the surface O atom. The O[^]O configuration denotes the configuration that both O atoms in the carboxylic acid group form a Ti-O bond with surface Ti atoms. In the latter adsorption mode, the dye is deprotonated to have H from the carboxylic acid covalently bound to a surface O atom during geometry relaxation. The N[^]O adsorption is more stable for all dyes considered here (see adsorption energies listed in Table 2), consistent with our previous findings. This configuration is then adopted for further

Table 2. Computed Adsorption Energies (in eV)

adsorption	VB0	VB2	VB4	L2	M2
N^O	1.39	1.28	1.25	1.30	1.33
O^O	0.84	0.87	0.91	0.90	0.93

discussion. Representative VB0, VB2, and VB4 dyes are taken as examples, since other dyes show qualitatively the same trend. Following eq 6, we calculated driving force ΔG_{iniect} listed in

Table 3, to estimate the injection efficiency Φ_{inject} . From VB0,

Table 3. Calculated Electronic Properties of VB0, VB2, and VB4 Adsorbed onto TiO₂ Anatase (101)

dye	$E^{\rm dye}$ (eV)	E_{0-0} (eV)	$E^{\rm dye}*({ m eV})$	$\begin{array}{c} E_{\mathrm{CB}} \\ \mathrm{(eV)} \end{array}$	${\Delta G_{ m inject} \over m (eV)}$	$\mu_{ m normal} \ (D)$	n _c
VB0	-6.38	3.93	-2.45	-4.17	1.72	8.67	0.72
VB2	-6.03	2.89	-3.14	-4.05	0.91	11.66	0.12
VB4	-5.71	2.47	-3.24	-3.95	0.71	16.38	0.04

VB2, to VB4, the reduction potential of the conduction band $E_{\rm CB}$ increases as the dipole moment $\mu_{\rm normal}$ pointing out of the surface gets larger. At the same time, the $E^{\rm dye_{\#}}$ decreases so $\Delta G_{\rm inject}$ and thus $\Phi_{\rm inject}$ decrease (see Figure 6). Considering that the LHE increases with the length of the linker group, we need to find a balance between competing $\Phi_{\rm inject}$ and LHE to reach a large $J_{\rm SC}$.

We could also estimate the $V_{\rm OC}$ through eq 2. It is reasonable to assume $N_{\rm CB}$ and $E_{\rm redox}$ to be constant for different dyes as they are determined by the properties of the substrate and electrolyte, respectively. The conduction band edge of semiconductor could be extracted from projected density of states calculations ($E_{\rm CB}$ in Table 3). The temperature of 300 K and typical $N_{\rm CB}$ density of 7 × 10²⁰ cm⁻³ are adopted.⁸ The dye loading density⁶ is on the scale of 100 mmol cm⁻³. Then kT $\ln(n_c/N_{CB})$ is calculated to be 0.11, 0.06, and 0.03 eV for VB0, VB2, and VB4, respectively. However, the effect of electron occupation in conduction band is relatively small compared to the influence of surface dipole moment. As a result, the $V_{\rm OC}$ for VB2 and VB4 are 0.07 and 0.14 V larger, respectively, than VB0. In real devices, dark current is another influencing factor that must be considered. Experiments show that longer dye may cause larger dark current.⁶ Possible reasons include the loss of surface protection for long dye and flexible donor groups binding to the surface. Taking all above considerations, we expect that VB2 will show a proper V_{OC} in devices. Other factors affecting energy conversion efficiency including dye loading concentration, dye aggregation, and desorption are out of the scope of this paper and will be investigated in future work.

Comparison to Dyes with a Different Donor. Compared to L and M series of dyes, VB dyes have an obviously larger donor group. In general, a large donor unit



Figure 6. Projected density of states (PDOS) of VB0, VB2, and VB4 adsorbed on TiO_2 anatase (101) surface. The unoccupied states of the dye have been upshifted according to the vertical excitation energy from TDDFT calculations.

could prevent electrolyte from contact with the semiconductor substrate, effectively blocking charge recombination between TiO_2 and the electrolyte and reducing the unfavorable aggregation of dyes at the semiconductor surface.²⁹ Therefore, VB dyes would be more favorable in DSSC applications over L and M series of dyes. To make a more accurate evaluation of VB dye characteristics and the potential utilizations in DSSCs, we compare the distinct properties among VB2, L2, and M2 dyes, which are different only in the donor unit (see Figure 1).

As shown in Table 4, although VB2 has maximum absorption at $\lambda_{max} = 429$ nm, slightly smaller than M2 (452 nm) and L2 (462 nm), VB2 presents the largest oscillator strength, leading to the largest LHE in turn. When VB2 adsorbed on TiO₂ semiconductor substrate, the ΔG_{inject} is slightly smaller than that of L2 and M2. However, an energy difference of 0.91 eV between dye LUMO and TiO₂ CBE is large enough to guarantee efficient electron injection. On the other hand, too large ΔG_{inject} may introduce energy redundancy, resulting a smaller V_{OC} and large thermalization losses.²⁹ In the latter process the interactions between excited charge carrier and lattice phonons would have the carrier cooled down to the bandgap edge, losing its initial high potential energy. More importantly, the number of transferred electrons in TiO₂ conduction band upon VB2 adsorption is almost twice that for L2 and M2. This implies a stronger surface polarization and electronic coupling between VB2 and the TiO₂ substrate. Also, a large electron transfer will favor a higher $V_{\rm OC}$. As the conduction band edge of TiO₂ $E_{\rm CB}$ is located at -4.05 and -4.18 V respectively upon VB2 and L2 adsorption, the $V_{\rm OC}$ will be 0.13 V larger for VB2 dye than for L2. The enhanced LHE and higher $V_{\rm OC}$ suggest that VB2 is more suitable for highly efficient DSSCs.

In conclusion, we report a new type of donor group, ullazine donor, to be used for organic dye sensitized solar cells. By lengthening the π bridge, we successfully increase the dye's light harvesting efficiency and dipole moment, and we find VB2 is the best among all the five VB dyes because it performs nicely on the four key parameters (λ_{max} , LHE, ΔG_{iniect} , V_{OC}) and achieves a good balance among these factors. Furthermore, compared with widely adopted existing donor groups, diphenylaminophenyl and dimethylaminophenyl donors, application of the new ullazine donor shows an enhanced extinction coefficient and LHE, an increased number of electrons transferred to conduction bands of TiO₂, and a high V_{OC} . In addition, the photoabsorption property and regeneration potential of VB dyes could be separately optimized. All these characteristics suggest VB dyes a competitive candidate for allorganic, highly efficient, near-red absorption DSSCs.

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Author Contributions

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Notes

The authors declare no competing financial interest.

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Table 4. Calculated Optical and Electronic Nature of VB2, L2, and M2 Dyes^a

dy	e λ_{\max} (nm)	f	LHE	assignment	$\Delta G_{ m inject}$ (eV)	$\mu_{\rm normal}$ (D)	n _c
VE	2 428.6	1.83	0.985	$H-1 \rightarrow L(88\%)$	0.91	11.66	0.12
				$H-2 \rightarrow L(7\%)$			
L2	462.3	1.71	0.980	$H \rightarrow L(83\%)$	1.01	14.30	0.07
				$H-1 \rightarrow L(11\%)$			
M	452.2	1.51	0.969	$H \rightarrow L(91\%)$	0.98	16.86	0.06

^aThe three groups are only different in their donor groups.

The Journal of Physical Chemistry C

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