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dve-sensitized solar cells (DSSCs).



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Efficiency difference between furan- and thiophene-based $D-\pi-A$ dyes in DSSCs explained by theoretical calculations

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The performance of two donor- π -bridge-acceptor type phenothiazine dyes bearing different π -bridges (furan and thiophene) was investigated by density functional theory and time-dependent density functional theory to explore the reasons for the differences in DSSC efficiency. It was revealed that **dye1** with furan showed higher short-circuit photocurrent density due to its larger driving force and better light harvesting efficiency compared with **dye2**. Moreover, a larger number of photo-injected electrons into TiO_2 for **dye1** leads to higher open-circuit photovoltage. Our results indicate that furan could be used as a promising π -bridge to improve the efficiency of PTZ dyes. We hope that our work can provide a theoretical basis and view for designing efficient dyes in

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Introduction

Dye-sensitized solar cells (DSSCs) have garnered considerable attention due to their great advantages, such as light weight, lowcost, and easy processing, as well as colorful and transparent features. 1-3 As one of the crucial parts in dye-sensitized solar cells, the photosensitizers including ruthenium complexes, 4-6 porphyrins,7 and metal-free organic dyes8-13 have been designed and applied to DSSCs in the past decades. Although an overall powerconversion efficiency above 11% has been established with ruthenium-based sensitizers,14 the limited resources and the environmental issues related to ruthenium use make it necessary to seek alternative dyes. Organic dyes are one of the most widely studied sensitizers for DSSCs because of their high extinction coefficients and environmental effects.15 However, the low conversion efficiency limits their practical application in our daily life. In recent years, much effort has been devoted to overcome this bottleneck.16-19 Most of the reported organic dyes with high efficiencies depended on thiophene or thiophene-based heterocycles as the π -bridge, for example, coumarin-thiophene, ²⁰ carbazoleoligothiophene,21 and benzothiadizole-thiophene.22 These dyes show good stability because of the fact that holes in such dyes are located on the thiophene moieties. And furan, thiophene's oxygen analog, which has a high oxidation potential, would be more efficient for the holes location and reinforce the stability of the dye-

Theoretical background

The overall conversion efficiency η of solar cells is determined by the short-circuit photocurrent density J_{sc} , the open-circuit photovoltage V_{oc} , the fill factor FF, and the intensity of the solar power P_{in} :

$$\eta = FF \frac{V_{\rm oc} J_{\rm sc}}{P_{\rm in}} \tag{1}$$

In DSSCs, the $J_{\rm sc}$ is determined by the following equation:²⁶

$$J_{\rm sc} = \int LHE(\lambda) \Phi_{\rm inj}(\lambda) \eta_{\rm coll}(\lambda) d\lambda$$
 (2)

• LHE(λ), the light harvesting efficiency (LHE) at the given wave-length λ ;

sensitizers.^{21,23} However, furan has received much less attention compared to thiophene unit. Furan unit was incorporated into phenothiazine (PTZ) dye as an alternative to thiophene from 2011, showing better conversion efficiency than that containing thiophene unit.²⁴ The similar experimental phenomena had also been reported by Hua *et al.* but again without deep understanding why the efficiency was improved.²⁵ In current contribution, density functional theory (DFT) and time-dependent density functional theory (TD-DFT) were employed to explore the different performances of the dyes with respective furan and thiophene π -bridge.²⁴ We hope the current contribution would provide theoretical guidance on optimizing the efficiency of existing donor– π bridge–acceptor (D– π –A) dyes or even designing new D– π –A dyes with better conversion efficiency.

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• $\Phi_{\rm inj}(\lambda)$, the electron injection efficiency at the given wavelength;

• $\eta_{coll}(\lambda)$, electron collection efficiency on the interface between dye and the semiconductor at the given wave-length.

In the similar DSSCs systems with only one difference part in the π -bridge fragment of the sensitizer, $\eta_{\rm coll}(\lambda)$ could be assumed to be a constant due to the similar structure of the dye-semiconductor interface. As a result, for the enhancement of $J_{\rm sc}$ the strategy of research should focus on improving LHE and $\Phi_{\rm ini}$. The light harvesting efficiency of the dye in the whole UV-vis range could be evaluated by calculating the area of the absorption bands. Obviously, the larger area of the absorption bands, the better for the short-circuit photocurrent density. In principle, to make the electron transfer energetically, on the one hand, the redox potential of the dye in the excited state, E^{dye*}, should be higher than conduction band of the semiconductor; on the other hand, the energetics of the excited states should be high enough to provide a thermodynamic driving force (D_f) for charge injection, assuming injection takes place at or near the Frank-Condon region. Furthermore, we made the approximation that the energetics of the dye are not different whether it is in CH₂Cl₂ solution or attached to TiO2 film, which is supported by previous studies performed by Kuciauskas. The driving force, $D_{\rm f}$, can be expressed as^{29,30}

$$D_{\rm f} = E^{\rm dye^*} - E_{\rm CB} = E_{\rm dye} + \lambda_{\rm max} - E_{\rm CB} \tag{3}$$

- E^{dye*}, the redox potential of the excited dye;
- \bullet E_{dye} , the redox potential of dye in ground state;
- λ_{max} , the largest vertical transition energy;
- \bullet $E_{\rm CB}$, the conduction band edge of the semiconductor, which is sensitive to the conditions, and the experimental value -4.00 eV (vs. vacuum) is used widely.31

The DSSCs open-circuit photovoltage V_{oc} is the difference of electronic potential between the quasi-Fermi level of the semiconductor under illumination and the redox potential of the electrolyte. We can calculate the $V_{\rm oc}$ as:³²

$$V_{\rm oc} = \frac{E_{\rm C} + \Delta {\rm CB}}{q} + \frac{kT}{q} \ln \left(\frac{n_{\rm c}}{N_{\rm CB}} \right) - \frac{E_{\rm readox}}{q} \tag{4}$$

where q is the unit charge, E_c is the conduction band edge (CBE) of the semiconductor substrate, T is the absolute temperature, kis the Boltzmann constant, n_c is the number of electrons in the conduction band, $N_{\rm CB}$ is the density of accessible states in the conduction band, $E_{\rm readox}$ is the reduction-oxidation potential of electrolyte, Δ CB is the shift of the E_c when the dyes are adsorbed on substrate and can be expressed as:33

$$\Delta CB = \frac{q\mu_{\text{normal}}\gamma}{\varepsilon_0 \varepsilon} \tag{5}$$

here q is electron charge, γ is the dye's surface concentration, μ_{normal} is the dipole moment of individual dye molecules perpendicular to the surface of the semiconductor substrate, ε_0 and ε represent the vacuum permittivity and the dielectric permittivity, respectively. According to eqn (4) and (5), it is obvious that n_c and μ_{normal} exert crucial influences on V_{oc} .

Computational details

The ground state of the isolated dyes was optimized at B3LYP/ 6-31G(d)³⁴ level of theory in gas phase. Frequency calculations were performed at the same level of theory as the geometry optimizations to ensure that the ground state geometry corresponds to a minimum point on the potential energy surface. For TD-DFT calculations of isolated organic dyes in CH₂Cl₂, we evaluated several functional, including B3LYP,³⁴ CAM-B3LYP,³⁵ PBE0,^{36,37} BHandHLYP,³⁸ and M062X,³⁹ to screen out a proper exchange-correlation functional for current study. The conductor-like polarizable continuum model (C-PCM) was considered to simulate the solvent effects of dichloromethane. The results in Table 1 given by TD-M062X/6-31G(d) were in good agreement with the experimental values in terms of transition energies.24 The absorption spectra were afterwards simulated by using Gaussian function with full-width at half-maximum (FWHM) of 0.3 eV. The oxidation potential of the ground state was calculated by B3LYP/UB3LYP/6-31G(d) level of theory according to the Born-Haber cycle.28 For geometry optimization of TiO2 cluster and dye/TiO2 complexes, plane-wave DFT calculations were performed using projector augmented wave (PAW) pseudopotentials as implemented in the VASP package.39,40 These calculations were carried by using generalized gradient approximation (GGA) of Perdew-Burke-Ernzerhof41 to account for the exchange-correlation effects. After optimization, the adsorption energies (E_{ads}) of dyes on TiO_2 surface were calculated with the following equation:

$$E_{\text{ads}} = E_{\text{dve}} + E_{\text{TiO}_2} - E_{\text{dve+TiO}_2} \tag{6}$$

Here E_{dye} denotes the total energy of isolated dye, E_{TiO_2} represents the total energy of TiO_2 cluster, and $E_{dve+TiO_3}$ denotes the total energy of dye/TiO2 complex. The frontier molecular orbitals (FMO) analysis of the dye/TiO2 were performed under PBE0 level, and the natural population analysis (NPA) was carried out at M062X for ground state and TD-M062X for excited state to evaluate the number of photoinjected electrons (Δq). The dipole moments of free dyes were obtained at the geometry of the dyes bound to TiO₂ cluster under the C-PCM-B3LYP level. The 6-31G(d) for nonmetal atoms and LanL2DZ basis set for Ti atoms have been selected for all the calculations for dye/TiO₂ complexes. The calculations of the geometry optimizations of dye/TiO2 complexes and TiO₂ cluster were in VASP, and the others were performed with Gaussian 09 program package.42

Table 1 Absorption peak comparison between experimental value λ_{max} and calculated λ_{max} under TD-DFT/6-31G* with PCM(CH₂Cl₂) level of theory

Functional	B3LYP	CAM-B3LYP	PBE0	BHandHLYP	M062X	Exp ²⁴
$\lambda_{\text{max}}/(\text{nm})$	534 0.569		503 0.685	414 1.296	420 1.222	442

Results and discussion

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The effect of different π -bridge on isolated dyes

To the dye molecules used in DSSCs, it is desirable to ensure that there exists a strong charge transport property and the electrons can be successively transferred to TiO_2 along D- π -A direction. From the perspective of molecular structure, the dye molecules need to be a flat structure and have small distances between adjacent parts. The optimized ground-state geometries of the **dye1** and **dye2** are displayed in Fig. 1, and the important geometrical parameters are collected in Table 2. The dyes are constructed by phenothiazine donor moiety (D), furan or thiophene as π -bridge (π) and cyanoacrylic acid acceptor part (A).

In the D- π -A dyes, the π -bridge is expected to function as the bridge of intramolecular charge transfer (ICT). 43,44 Therefore, the bridge bond between D and π -bridge or π -bridge and A has important effects on ICT. Herein, the bridge bonds between D and π -bridge, and π -bridge and A were marked as D- π and π -A, respectively. The shorter length of bridge bonds is more beneficial for ICT in the D- π -A dyes. As shown in Table 2, the values of D- π bond lengths of dye1 and dye2 are 1.451 and 1.461 Å, respectively, which implies the intensity of interaction between PTZ and π -bridge is in the order of furan > thiophene. On the other hand, the π -A bond lengths of **dye1** and **dye2** are 1.416 and 1.426 Å, which indicates that interaction of furan-A is stronger than that of thiophene-A. For **dye2**, the dihedral angle between donor and π -bridge was calculated to be 22.3°, and it becomes 0.2° in **dve1**. For these two dves, the acceptor groups (A) were nearly coplanar with the π -bridge groups (π) as π -A dihedral angles were calculated to be 0.1° and 0.9° for dye1 and dye2, respectively. Considering above, dye1 may be more beneficial for ICT due to its shorter bridge bond lengths and flatter conjugated structure.

One of the important requirements for improving the efficiencies of dyes is to extend the cover of the absorption spectra. The calculated absorption spectra of both dyes are shown in

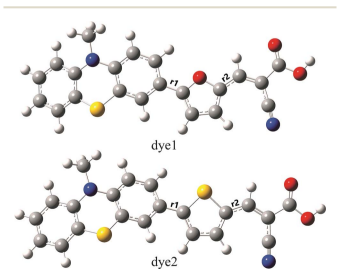


Fig. 1 Optimized ground-state geometries with tagged bond lengths of dye1 and dye2 (C atom in grey; H atom in white; O atom in red; S atom in yellow; N atom in blue).

Table 2 The selected bond lengths (r_1 and r_2 are marked out in Fig. 1) in angstroms and dihedral angles (Φ) in degrees of **dye1** and **dye2** (D: donor; π : π -bridge; A: acceptor)

	D-π		π-Α		
Molecule	r_1	Φ	r_2	Φ	
dye1	1.451	0.2	1.416	0.1	
dye1 dye2	1.461	22.3	1.426	0.9	

Fig. 2, and the corresponding data is summarized in Table 3. The results are in agreement with experimental measurement and this method is best for simulating the absorption spectra in our calculations. Both of dyes evidently exhibit major absorption bands in the 420 nm region. Markedly, the absorption intensity of **dye1** is higher a bit than the **dye2**. All the absorption peaks are contributed from the HOMO to LUMO or HOMO -1 to LUMO transitions with the D- π -A direction. The integral area of the absorption bands for **dye1** and **dye2** are 128 and 117, respectively, revealing that **dye1** has better LHE to **dye2** over the almost same absorption region. Therefore, it can illustrate that the modification of π -bridge, by replacing thiophene with furan, has gradually enhancing effect in light harvesting efficiency (LHE).

Rapid and efficient electron injection after photo-excitation is also very important for improving the efficiency of the cell. Thereupon, the driving force (D_f) was also studied in detail. The calculated D_f as well as E_{dye} and E^{dye*} for both PTZ dyes are listed in Table 4. It can be found that electron injection efficiency $(\Phi_{\rm ini})$ increases along with higher $E^{\rm dye^*}$ and larger $D_{\rm f}$ in dye1 in contrast to dye2, thus indicating that the injection of the electron from excited state of dye1 to the TiO2 is much easier than dye2. Notably, the change of thiophene to furan improved the LHE but also shifts the $D_{\rm f}$ by 0.20 eV. Hence, considering the improved LHE and $\Phi_{\rm inj}$, it can be expected that the DSSCs based on **dye1** should have minute intrinsic enhanced J_{sc} than **dye2**, and that is also reflected in the experimental results (12.18 vs. 12.05 mA cm⁻² in I_{sc} measurement). These tiny changes have also recently been captured by experimental scientists and used in the design of more new D- π -A dye molecules, ^{45,46} and now we have a clear theoretical cognition.

The effect of different π -bridge on dye-TiO₂ complexes

The anatase (101) surface is the most active surface since it has unpassivated Ti and O atoms, which has been widely used to

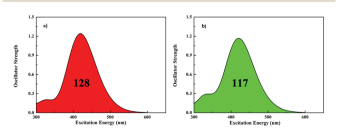


Fig. 2 Simulated absorption spectra for dye1 (a) and dye2 (b) in CH_2Cl_2 solution (128 and 117 represent the area of absorption bands for dye1 and dye2 in the whole UV-vis region, respectively).

Table 3 The excitation energies (λ_{max} /nm), oscillator strengths (f), and compositions of the lowest state for **dye1** and **dye2** (H: HOMO; L: LUMO)

	State	Composition (%)	λ_{max}	f	$\lambda_{\rm max}$ (exp with molar absorption coefficient) ²⁴
dye1	$S_0 \to S_1 S_0 \to S_1$	$H \rightarrow L (82.5), H - 1 \rightarrow L (11.8)$	420	1.222	442 (15 100)
dye2		$H \rightarrow L (81.6), H - 1 \rightarrow L (11.7)$	421	1.117	448 (16 200)

Table 4 Calculated redox potential, excitation energy λ_{max} and D_{f} (in eV), dipole moment μ_{normal} (in Debye) and number of electron injection Δq (in e) of **dye1** and **dye2**

	$E_{ m dye}$	λ_{max}	$E^{\mathrm{dye}*}$	$D_{ m f}$	$\mu_{ m normal}$	Δq
dye1 dye2	-5.07 -5.27	2.95 2.95	-2.21 -2.32	1.88 1.68	2.68 4.71	0.129 0.107

adsorb dyes for the interface investigation. To model the TiO₂ nanoparticle surface, we considered a (TiO₂)₃₈ cluster, obtained by appropriately "cutting" from anatase slab exposing the majority (1 0 1) surface.47 The considered model represents a reasonable tradeoff between accuracy and computational convenience, and nicely reproduces the main electronic characteristics of TiO2 nanoparticles. We obtained HOMO-LUMO gap of 4.13 eV, while the TD-DFT lowest excitation energy of 3.63 eV is in agreement with the experimental spectrometric measurement. For the studied (TiO₂)₃₈ (101) model, the occupied MO can be attributed the contribution of 2p orbitals character of O atoms. While, the unoccupied MO can be attributed from the contribution of 3d orbitals of Ti atoms. These characters are in agreement with that of the understanding in TiO2 semiconductor material. Though HOMO-LUMO gap for this model is extended than the experimental band gap of crystal (3.2-3.3 eV),31 it corresponds well with previously calculated HOMO-LUMO gaps for other (TiO2)38 clusters.48-50 We used the plane-wave DFT to optimize the ground-state geometry of the (TiO₂)₃₈ cluster and dye/(TiO₂)₃₈ complex. The electronic structure and related results of the (TiO₂)₃₈ (101) model is reasonable and reliable for describing the conduction band properties of TiO2 semiconductor based on the both of accuracy and computational cost consideration.

In dye/TiO2 adsorption model, organic dyes adsorbed on the TiO₂ surface through the carboxylic group with several adsorption configurations owing to molecular or dissociative adsorption processes. The molecular adsorption is formed via hydrogen bonding between TiO2 surface and the dye. When the dissociative adsorption happens (H atom of carboxylic acid dissociates), the bond is formed between carboxylic oxygen atoms and the surface titanium of TiO2, which can be monodentate ester, bidentate chelating, and bidentate bridging types. 51,52 However, Fourier-transform infrared spectroscopy (FTIR) results for organic systems in many experimental studies indicate that carboxylic acid usually adsorbs on TiO2 surface through bidentate bridging structure, which was most studied in the literature and was thought to be the most stable.⁵²⁻⁵⁴ Therefore, only bidentate bridging adsorption was considered in current study. This adopted TiO2 model well described the interface charge transfer and the most important thing is that the adsorbed dye on TiO_2 *via* its carboxylate group in a bidentate bridging manner, with one proton transferred to a nearby surface oxygen has been proved to be the most stable adsorption mode for this anchoring group.

The optimized structures of dye/TiO₂ complexes are shown in Fig. 3, and the important bond lengths and adsorption energies are listed in Table 3. The adsorption energy of **dye1**, calculated to be 22.9 kcal mol⁻¹, is larger than that of **dye2** (21.4 kcal mol⁻¹) and both of them are large enough to allow the dyes to be adsorbed on the cluster in the chemical way. The bond lengths between Ti atoms and the carboxylic oxygen atoms were calculated to be 2.061 and 2.079 Å for **dye1**, 2.035 and 2.108 Å for **dye2**, respectively. The average Ti–O bond length of **dye1**/TiO₂ is slightly shorter than that of **dye2**/TiO₂, indicating somewhat stronger binding of **dye1** with TiO₂ cluster. Intuitively, one can indeed expect a stronger electronic coupling in the **dye1**/TiO₂ system rather than **dye2**/TiO₂ system (Table 5).

In the most of important excitations (in Table 3), the LUMO shows considerable involvement and has largest contribution at largest adsorption energy $\lambda_{\rm max}$. Hence, the electronic coupling between the dye's LUMO and TiO₂ conduction band is important for electron injection from the former to the latter. To gain insight into the electronic coupling between the dye's LUMO and TiO₂ conduction band and the electron-transfer features during photo excitation, the frontier molecular orbitals (FMO) of dye/TiO₂ system were analyzed. HOMO, LUMO, and the interacting orbitals for **dye1**/TiO₂ are shown in Fig. 4. The HOMO is localized on dye part of the complex as can be seen from Fig. 4(a), which is characteristically similar to that of the isolated dye. While, the LUMO is entirely localized on TiO₂ part

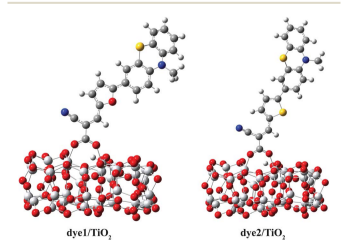


Fig. 3 Optimized structures of dye1 and dye2 adsorbed on TiO_2 surface.

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Table 5 Optimized bond lengths (in angstrom) and adsorption energy (in kcal mol^{-1}) of $dye1/TiO_2$ and $dye2/TiO_2$

	O-Ti (1)	O-Ti (2)	$E_{ m ads}$
$\begin{array}{l} \textbf{dye1/TiO}_2 \\ \textbf{dye2/TiO}_2 \end{array}$	2.061	2.079	22.9
	2.035	2.108	21.4

of the complex (Fig. 4(b)). The interacting LUMOs that have non-negligible contributions from the isolated free dye's LUMO are situated higher up in the conduction band (LUMO + 49). For dye2/TiO2 system, the FMOs features are similar to those of the dye1/TiO2 system. In DSSCs, the efficient electrons are transferred from donor through π -bridge to cyanoacrylate. Based on this, the spatial orientation of the FMOs for the isolated dye (shown in Fig. 5) and the dye/TiO2 interface here (shown in Fig. 4) are ideal conditions for DSSCs. Fig. 6 shows the density of states for dye1 and dye2 before and after adsorbing on TiO2 surface. From which we find that the HOMO of both dyes are almost unaffected by interaction with TiO₂, showing the alignment of the energy levels as in the no-interacting case. The obvious difference between the isolated and interacting systems is in the shape and distribution of the LUMOs. Both of LUMOs of dyes adsorbed on TiO2 surface decrease due to the interaction with TiO2 semiconductor, and this is in agreement with previous reports of the perylene dye and TiO₂ cluster.⁵⁵ For dye1, the LUMO is 0.25 eV lower after absorbing on the TiO₂ surface. While for dye2, the decrease of LUMO is calculated to be 0.23 eV, indicating slightly weaker electronic coupling to TiO₂ surface than dye1. In case of similar molecular geometry when anchored to TiO2, the large stabilization to the LUMO will enhance the electronic coupling between the dye and the titanium 3d orbitals and favor the electron injection. The charge transfer states with direct charge injection character are performed by TD-DFT, and the results indicate that the absorption transitions of both adsorbed systems are bathochromic shifted near 500 nm, and the dye1/TiO2 is increasing 40 nm towards longer wavelength with higher intensity (oscillator strength 0.98 vs. 0.82) in contrast with dye2/TiO2. These results are also in concert with previous adsorption energy and binding strength values.

Through eqn (5), the different dipole is expected to substantially influence the $E_{\rm CB}$ of ${\rm TiO_2}$, which will lead to a change in open-circuit photovoltage $V_{\rm oc}$. Grätzel and

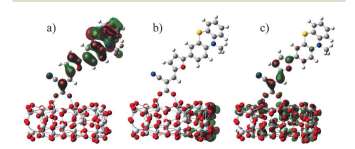


Fig. 4 Some selected MO profiles for the HOMO (a), LUMO (b) and interaction orbital (c) of $dye1/TiO_2$.

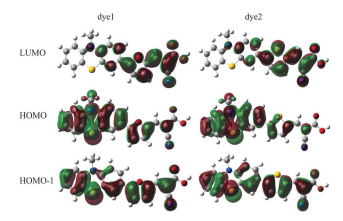


Fig. 5 Selected Frontier molecular orbitals at ground state of **dye1** and **dye2**.

coworkers have confirmed theoretically that the larger the dipole moment μ_{normal} of the adsorbed molecules pointing outward from the semiconductor surface induced the larger $V_{\rm oc}$. Accordingly, $\mu_{\rm normal}$ of both dyes were calculated under B3LYP/6-31G(d) level including solvent effects at the geometry of dyes adsorbed on TiO2 and the results are listed in Table 4. It turns out that **dye2** has much larger μ_{normal} than **dye1**, thus it can be inferred that dye2 should have larger V_{oc} than dye1. But this is not in line with the experiment results (734.6 and 706.6 mV for dye1 and dye2, respectively).24 According to eqn (4), we know that V_{oc} also depends on the charge distribution in the semiconductor. As a result, we calculated the number of charge injected into the TiO_2 conduction band Δq by means of natural population analysis (NPA) and the results are listed in Table 4. **Dye1** has a larger Δq than **dye2** (0.129e > 0.107e), which could shift the quasi-Fermi level of TiO2 accordingly, then lead to larger $V_{\rm oc}$. Therefore, we believe that the larger experimental $V_{\rm oc}$ of **dye1** is mostly ascribed to the more electrons transferred from **dye1** to TiO_2 . Although the difference in V_{oc} for **dye1** and dye2 has been demonstrated, it needs to be pointed out that the $V_{\rm oc}$ is also determined by other factors, such as electron lifetime. The relevant study is underway in our following work.

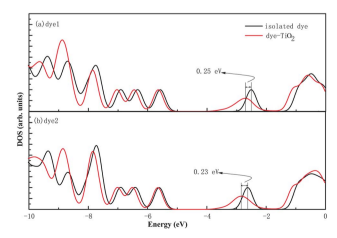


Fig. 6 The density of states of dye1 and dye2 before and after adsorbed on the TiO_2 surface.

Conclusion

In summary, the geometries, electronic structures, absorption spectrum of two PTZ dyes with different π -bridges were investigated to gain insights into the effects of the π -bridge group of furan and thiophene on the performance of the dyes. The electrons in furan-containing dye could be more easily transferred from PTZ-donor to acceptor than that in thiophenecontaining one due to the shorter bridge bonds and the flatter structure. After adsorption to the TiO₂ cluster, as compared with dye2, dye1 shows a stronger coupling with the TiO₂. dye1 with furan shows intrinsic enhanced short-circuit photocurrent density J_{sc} than **dye2** with thiophene because it has larger driving force and comparable light harvesting efficiency, and the more electrons transferred from dye1 to TiO2 surface lead to larger open-circuit photovoltage $V_{\rm oc}$. Our results indicate that furan could be used as a promising π -bridge to improve the efficiency of PTZ dyes. We also hope our work will be helpful for designing more efficient PTZ dyes in DSSCs with reduced economical cost and synthesis effort.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

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