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Small molecular push-pull donors for organic photovoltaics: effect of the heterocyclic π -spacer

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Abstract: A series of $(D-\pi-A)$ small push-pull molecules involving a triphenylamine electron-rich group (D) connected to a dicyanovinyl electron-deficient unit (A) through different chalcogenophene type π -connectors has been synthesized. Optical and electrochemical results reveal that the replacement of furan by thiophene and selenophene leads to a progressive decrease of the optical band gap of the material and to a parallel increase of hole mobility and power conversion efficiency (*PCE*). Thus, a *PCE* of 3.33% has been obtained for a simple air-processed solar cell involving the selenophene-based donor and the [6,6]-phenyl-C₆₁-butyric acid methyl ester (PC₆₁BM) as the acceptor.

1. Introduction

Over the past decades, organic solar cells (OSCs) have generated a considerable research effort due to the lightness, flexibility, low environmental impact and potential low cost of organic materials.^{1, 2} Recent progress in device fabrication and material engineering have led to remarkable progress with power conversion efficiencies (*PCE*) now exceeding the symbolic value of 10%.³⁻⁵

However, beyond striving for improved *PCEs*, the development of active materials by simple, clean, cost-effective, efficient and scalable syntheses is a mandatory condition for a future industrial production.⁶⁻⁸ Owing to their well-defined chemical structure, molecular active materials present the advantages of a better reproducibility of synthesis and purification and a

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more precise analysis of the structure-property relationships compared to polydisperse polymers.⁹⁻¹¹ Among them, Donor-Connector-Acceptor (D- π -A) type push-pull molecules based on arylamines donor blocks represent interesting molecular donors that combine synthetic accessibility, simplicity and efficiency.^{6, 12-15} Indeed, **TPA-T-DCV** (Figure 1), prepared in good yields in a few synthetic steps, has been over the past few years an interesting working structure to investigate the effects of various structural variations such as (*i*) the covalent bridging of the dicyanovinyl (DCV) group,¹⁶ (*ii*) its replacement by other electron-withdrawing units,¹⁷ (*iii*) the substitution of one phenyl ring of TPA by fused ring systems or aliphatic chains,^{18, 19} or (*iv*) the drastic size reduction to assess the limit of simplification.²⁰

As a further step of this systematic analysis of structure-properties relationships, we report here on the effect of the nature of the heterocyclic π -connector by the synthesis, characterization and photovoltaic evaluation of the furan (**TPA-F-DCV**) and selenophene (**TPA-S-DCV**) analogues of **TPA-T-DCV**. (Figure 1).



Figure 1. Chemical structure of the D- π -A target molecules

Indeed, while several articles deal with the effect of the replacement of the thiophene ring by other chalcogenophene in conjugated polymers for OSCs,²¹⁻²⁵ only few examples of molecular donors, especially incorporating the selenophene ring, have been reported.²⁶⁻²⁹

2. Results and discussions

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The synthesis of the three molecules is depicted in Scheme 1.

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Scheme 1. Synthetic route to TPA-F-DCV, TPA-T-DCV and TPA-S-DCV

5-Formylfuran-2-boronic acid was engaged in a Suzuki cross-coupling reaction with the commercially available 4-bromotriphenylamine **1** affording the carbonyl compound **4**.³⁰ In parallel, the thiophene and selenophene analogues **5** and **6** were prepared by a two-step procedure involving a Stille cross-coupling of 4-bromotriphenylamine **1** and the appropriate trimethyl stannylated chalcogenophene, followed by a Vilsmeier-Haack formylation with POCl₃/DMF.¹⁶ Finally, a Knoevenagel condensation between malononitrile and the carbaldehydes **4**, **5** or **6** led to the target products **TPA-F-DCV**, **TPA-T-DCV** and **TPA-S-DCV** respectively.

The compounds show good solubility in common organic solvents and their thermal stabilities were assessed by thermogravimetric analysis (Figure 2).



Figure 2. Thermogravimetric analysis of TPA-F-DCV (black), TPA-T-DCV (red) and TPA-S-DCV (blue) recorded at 5° C/min under N₂.

The furan and thiophene derivatives exhibit a comparable decomposition temperature (Td) of ca 255°C. However, introducing a selenium atom slightly increases Td since a 5 % weight loss was recorded at ca 280 °C for **TPA-S-DCV**.

UV-Vis absorption spectra were recorded on chloroform solutions and on films spin-cast on glass.



Figure 3. UV-Vis absorption spectra of **TPA-F-DCV** (black squares), **TPA-T-DCV** (red circles) and **TPA-S-DCV** (blue triangles) in chloroform (a) and as thin film on glass (b).

The spectrum of all compounds shows an intense absorption band with a maximum (λ_{max}) around 500 nm assigned to an internal charge transfer (ICT) from the TPA moiety to the electron-deficient DCV group (Fig. 3). Replacement of the furan spacer by thiophene and selenophene leads to a progressive bathochromic shift of λ_{max} from 499 to 520 nm with a parallel increase of the molar extinction coefficient (ε) from 33000 to 36800 M⁻¹ cm⁻¹ (Table 1), in agreement with previous observations (Figure 3).^{23, 31, 32} Broader absorption bands and red-shifted absorption edges with onsets at *ca* 595 nm, 606 nm and 630 nm corresponding to optical band gaps (E_g^{opt}) of 2.08 eV, 2.04 eV and 1.96 eV for **TPA-F-DCV**, **TPA-T-DCV** and **TPA-S-DCV** respectively were observed on thin films (Figure 3, b).

Tał	ole	1.	U	V-'	Vis	absor	ption	data.
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Compound	$\lambda_{\max}(nm)$	3	$\lambda_{\mathrm{onset}}(\mathrm{nm})$	$\lambda_{\max}(nm)$	λ_{onset} (nm)	E_g^{opt} (eV)
Compound	solution	$(M^{-1} cm^{-1})$	solution	film	film	film
TPA-F-DCV	499	33000	560	517	595	2.08
TPA-T-DCV	505	35000	572	523	606	2.04
TPA-S-DCV	520	36800	587	535	630	1.96

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The electrochemical properties of the three molecules have been analyzed by cyclic voltammetry in dichloromethane solution in the presence of Bu_4NPF_6 as the supporting electrolyte. Data are summarized in Table 2.

Table 2. Electrochemical data for the target compounds (0.5 mM in 0.10 M Bu_4NPF_6/CH_2Cl_2 , scan rate 100 mV s⁻¹, Pt working and counter electrodes, ref. SCE)

Compound	E _{pa} ¹ (V)	E _{Ox/onset} (V)	$\frac{E_{\rm pc}^{-1}}{\rm (V)}$	E _{Red/onset} (V)	HOMO ^{a)} (eV)	LUMO ^{b)} (eV)	∠E ^{elec} (eV)
TPA-F-DCV	0.98	0.86	-1.24	-1.07	-5.85	-3.92	1.93
TPA-T-DCV	1.01	0.87	-1.20	-1.05	-5.86	-3.94	1.92
TPA-S-DCV	1.01	0.87	-1.20	-1.01	-5.86	-3.98	1.88

a) HOMO $(eV) = -(E_{Ox/onset}(V) + 4.99)$, b) LUMO $(eV) = -(E_{Red/onset}(V) + 4.99)^{33}$

The cyclic voltammogram (CV) of all compounds shows a reversible one-electron oxidation wave, assigned to the formation of a stable radical cation (Figure 4). While the CVs of **TPA-T-DCV** and **TPA-S-DCV** show comparable anodic peak potentials at $E_{pa} = 1.01$ V, the furan derivative presents a slight negative shift of E_{pa} to 0.98 V indicative of an increase of the HOMO level. In the negative potentials region, the CV of all donors exhibits an irreversible reduction process with cathodic potential peak at $E_{pc} = -1.24$ V for **TPA-F-DCV** and -1.20 V for both the thiophene and selenophene compounds. HOMO and LUMO levels were calculated respectively from the onsets of oxidation and reduction waves (Table 2).



Figure 4. Cyclic voltammograms of **TPA-F-DCV** (black), **TPA-T-DCV** (red) and **TPA-S-DCV** (blue) 0.5 mM in 0.1 M Bu₄NPF₆/CH₂Cl₂, scan rate 100 mV s⁻¹, Pt working and counter electrodes, ref SCE.

A slight reduction of the electrochemical gap (ΔE^{elec}) is observed from the furan derivative (1.93 eV) to the selenophene one (1.88 eV), in agreement with optical data. Electrochemical data also show that this decrease of ΔE^{elec} is mainly associated to the lowering of the LUMO energy levels when varying the heteroatom of the chalcogenophene from oxygen to selenium whereas the HOMO energy levels remains relatively unaffected.

To gain further insight into the electronic properties, the frontier orbitals and energy levels of the compounds have been investigated by theoretical calculations performed with Gaussian 09 program using Becke's three-parameter gradient-corrected functional (B3LYP) with the 6-31G(d,p) basis set (Figure 5).

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Figure 5. HOMO and LUMO energy levels and their representations for **TPA-F-DCV** (left), **TPA-T-DCV** (middle) and **TPA-S-DCV** (right) after optimization with Gaussian 09 at the B3LYP/ 6-31G(d,p) level of theory.

The optimized geometries and computed electronic structures at the observed minima reveal that the LUMO is strongly localized on the five-membered ring-DCV block whereas the HOMO is mainly distributed on the TPA-spacer. In addition, the geometry of **TPA-F-DCV** presents the smallest dihedral angle between the phenyl ring and the heterocycle (1.4° *vs* 19.6° and 17.6° for **TPA-T-DCV** and **TPA-S-DCV** respectively) ensuring a better planarity of the structure. The calculated values of the HOMO and LUMO levels are consistent with optical and electrochemical results.

The photovoltaic properties of the three donors have been evaluated in bulk heterojunction solar cells of 0.27 cm² of configuration: ITO/PEDOT: PSS (*ca* 40 nm)/ **Donor**: PC₆₁BM blend/ LiF (1 nm)/ Al (120 nm). Except for LiF and aluminum depositions, the cells were fabricated in ambient atmosphere. Table 3 gathers the photovoltaic parameters of these devices and Figure 6a shows the best current density-voltage (*J-V*) characteristics measured under AM. 1.5 simulated solar illumination (80 mW cm⁻²).

Donor	V _{oc} (V)	J_{sc} (mAcm ⁻²)	FF (%)	PCE _{max/ave} ^{a)} (%)	$\mu_h ({\rm cm}^2 {\rm V}^{-1} {\rm s}^{-1})$
 TPA-F-DCV	0.97	5.28	38.9	2.50/2.23	8.1 x 10 ⁻⁶
TPA-T-DCV	0.97	6.53	37.9	3.00/2.86	1.2 x 10 ⁻⁵
TPA-S-DCV	0.95	7.15	39.2	3.33/3.17	1.7 x 10 ⁻⁵

Table 3. Photovoltaic properties of **TPA-F-DCV**, **TPA-T-DCV** and **TPA-S-DCV** blended with $PC_{61}BM$ (1:2 w/w).

a) Average value recorded over 18 devices.

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The high open-circuit voltages ($V_{oc} \sim 1.0$ V) are consistent with the low-lying HOMO level of the donors resulting in a large offset with the LUMO level of PC₆₁BM. Although comparable V_{oc} and fill factors (*FF*) values are obtained for all cells, the short-circuit current-density (J_{sc}) progressively increases from **TPA-F-DCV** to **TPA-S-DCV** leading to a parallel increase of *PCE* from 2.50 to 3.33% (Table 3). This trend is confirmed by external quantum efficiency (*EQE*) measurements performed under monochromatic irradiation (Figure 6b).



Figure 6. J –V characteristics and EQE curves of **TPA-F-DCV** (black squares), **TPA-T-DCV** (red circles) and **TPA-S-DCV** (blue triangles) based OSCs.

Thus, varying the heteroatom from oxygen to selenium results in a slight extension of the contribution of the donor in the long wavelengths region with a maximum photon-to-current conversion of 52 % at 504 nm, 54% at 523 nm and 55% at 540 nm for **TPA-F-DCV**, **TPA-T-DCV** and **TPA-S-DCV** respectively.

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The hole-mobility of the materials (μ_h) was determined using the space-charge limited current (SCLC) method on hole-only devices (Figure 7).



Figure 7. J–V characteristics of hole only devices ITO/PEDOT:PSS/**TPA-F-DCV** (black squares), **TPA-T-DCV** (red circles) or **TPA-S-DCV** (blue triangles)/Au.

As shown in Table 3, the furan-containing molecule exhibits the lowest mobility (of *ca* 8.1×10^{-6} cm² V⁻¹ s⁻¹) and the selenophene derivative the highest $(1.7 \times 10^{-5} \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1})$. These results, in agreement with the J_{sc} trend, can be correlated to the improved orbitals overlap of heavier chalcogen heteroatoms resulting in enhanced intermolecular electronic interactions.

3. Conclusion

Triphenylamine based D- π -A type small molecular donors with three different fivemembered heterocycles as π -conjugating connector have been synthesized. Replacement of furan by thiophene and selenophene produces a small reduction of the band gap and an improvement of the hole-mobility and photovoltaic conversion efficiency of the donor. In addition to the synthetic accessibility (*ie*, three steps from commercially available materials), promising power conversion efficiencies above 3.0% were obtained with simple BHJ solar cells processed in ambient atmosphere with PC₆₁BM as acceptor. Further device optimization (morphology, additives, and interfaces) can be expected to significantly improve these preliminary results.

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5. References

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6. Graphical Abstract

