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Fluoro-alkyl substituted isothianaphthene

bisimides as stable n-type semiconductors†

The development of air-stable organic semiconductors (OSCs) with high charge mobilities is of pivotal importance for practical applications of organic thin film transistors (OTFTs) in various fields. Although several OSCs with high hole mobilities have been realized, their n-type counterparts lag far behind in terms of charge mobilities and especially air-stability potentially due to the highly sensitive nature of electrons to ambient oxidants. Isothianaphthene, a kind of non-classical thiophene, possesses a unique electronic structure and isothianaphthene bismide (BTDI) has been demonstrated as an efficient core for stable n-type OSCs. In this work, we introduced two fluoric groups with different alkyl-chains at the N-position of isothianaphthene-2,3,6,7-tetracarboxylic acid diimide and obtained new air-stable n-type OSCs, BTDI-OCF₃ and BTDI-CF. The distinct length of the alkyl-chains caused different molecular packing styles of the resulting materials, as revealed by atomic force microscopy. The crystallinity of BTDI-OCF3 is found to be much better than BTDI-CF and single crystals of BTDI-OCF3 could easily be obtained by solution process. Besides, OTFTs based on these new semiconductors showed almost no hysteresis when tested in a glovebox and very small hysteresis is observed in air with a small decay of current, indicating good stability of both of these materials for their applications in n-type OTFTs.

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Introduction

As an important building block for flexible and printed organic electronics, organic thin film transistors (OTFTs) have shown extensive prospects for a variety of applications, including electronic papers (e-papers), electronic skins (e-skins), sensors, radio frequency identification (RFID) tags, active-matrix displays, and synapses, owing to their large area, low cost, mechanical flexibility, and ease of fabrication. 1-5 Since the first report on OTFTs based on polythiophene in 1986,6 intensive research efforts from both academia and industry have been dedicated to realize high performance OTFTs, leading to

remarkable progress in the past few decades. For instance, a number of organic semiconductors (OSCs) have been developed with charge carrier mobilities exceeding 10 cm² V⁻¹ s⁻¹ for holes and $1 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$ for electrons. However, the development of n-type OSCs is still far behind the p-type counterparts in terms of their carrier mobilities and especially the air-stability, 12,13 due to the high sensitivity of electrons to ambient oxidants (such as O2 and H2O), which seriously degrade the charge mobility and device stability of n-channel OTFTs. 14,15 On the other hand, it is noteworthy that a high-energy barrier between the active layer and the source/drain electrodes further hinders charge injection. 16 The imbalance between the characteristics of p-type and n-type semiconductors significantly restricts the practical applications of OTFTs in complementary circuits and p-n junctions.

In general, the ambient stability of n-type OSCs for applications in OTFTs could be improved through two different molecular design strategies. One is decreasing the lowest unoccupied molecular orbital (LUMO) level below than that of the most atmospheric trapping oxidants via introducing strong electron-withdrawing groups, such as cyano (-CN), fluoroalkyl, imide/amide and so on. It is well established that a LUMO level of ca. -4.0 eV or lower is beneficial to realize airstable carrier transport for OSCs since it could minimize most

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of the atmospheric trapping.^{17,18} On the other hand, a low LUMO level also reduces the electron injection barrier, which allows efficient and stable electron injection and transport. Until now, a series of ambient-air-stable n-type OSCs, including perylene diimdes (PDI), naphthalene diimides (NDI), thieno[3,4-c]pyrrole-4,6-dione (TPD) and isoindigo have been developed. 12,19-23 Consequently, a considerable enhancement in mobility from 10⁻⁴ to 12.6 cm² V⁻¹ s⁻¹ and excellent airstability have been realized in n-channel OTFTs. 24-26 The other approach to improve the ambient stability of n-type OSCs is promoting a more densely-packed steric barrier to the atmosphere, which could prevent moisture and oxygen penetration into the organic layer in steric.²⁷ The introduction of a fluoroalkyl chain to the N-atom of the NDI or PDI core could lead to dense self-segregation of the side fluoro-substituted chain, which impedes the movement of moisture and oxygen into the active channel.²⁸⁻³¹ More importantly, these fluoric materials not only show excellent air-stability but also exhibit high electron mobilities with considerable on/off ratios in OTFTs. 26,32-35 For instance, the highest reported electron mobility of an NDI small molecule, Cl₂-NDI, is as high as 8.6 cm² V⁻¹ s⁻¹ for single crystals with a slight mobility degradation of 13% (after 3 months) in ambient air. 34,36

The electronic structure of isothianaphthene was first studied by Joseph³⁷ and later, Fred Wudl et al. reported poly(isothianaphthene) by a polymerization method.³⁸ The conductivity of poly(isothianaphthene) could be effectively adjusted by doping various ions. Isothianaphthene was also explored as an effective unit for low band gap conjugated polymers. In our previous work, we have reported an alkyl substituted isothianaphthene derivative, N,N'-bis(n-hexyl)isothianaphthene-2,3,6,7-tetracarboxylic acid diimide (BTDI-C6), which also possesses a deep LUMO level (-4.21 eV), approximately 0.32 eV lower than that of NDI-C6 and much higher airstability of the resulting OTFT devices.³⁹ It is well known that fluorinated substituents play a significant role in enhancing the air-stability of n-type OSCs because of their dense molecular packing, which prevents the diffusion of moisture and oxygen into the device. To date, 3-(perfluorooctyl)propyl (CF) and p-(trifluoromethoxy)benzyl (BOCF₃) substituted at the N,N'-positions of NDI have shown excellent air-stability and high mobility in OTFTs. 32,33 In this study, we introduced these two fluorinated substituents to the BTDI core to investigate their effect on the molecular packing and air-stability of the resulting BTDI derivatives, i.e. BTDI-CF and BTDI-OCF₃. Similar LUMO energy levels with a slight distinction of 0.02 eV were observed for these materials. Remarkably, BTDI-CF showed reasonable solubility in chloroform ($>2 \text{ mg mL}^{-1}$) and BTDI-OCF₃ also exhibited excellent solubility in chloroform (>10 mg mL⁻¹), while NDI-OCF3 is not soluble.33 The OTFT devices based on **BTDI-CF** showed higher mobility (0.0368 cm² V⁻¹ s⁻¹) but lower air-stability than the BTDI-OCF3 based counterparts and both types of the devices demonstrated good air-stability with small hysteresis. The relationship between the material structure and the corresponding device performance is further studied systematically.

Experimental

Synthesis and thermal analysis

The synthetic routes of BTDI-OCF₃ and BTDI-CF are shown in Scheme 1. The synthetic method was similar to that reported in the literature 39,40 and the obtained products were finally purified by sublimation with satisfactory yield. The corresponding synthetic procedures, chemical structure characterization (¹H NMR, ¹³C NMR and mass spectroscopy analysis) and further experimental details are provided in the ESI.† The thermal properties of BTDI-OCF3 and BTDI-CF are characterized by thermal gravimetric analysis (TGA) and differential scanning calorimetry (DSC). BTDI-OCF₃ showed better thermal stability and a higher melting point than BTDI-CF due to the more rigid substituent. Both BTDI-OCF₃ and BTDI-CF exhibited good thermal stability with decomposition temperatures of up to 385 °C and 338 °C, respectively (see Fig. S1a, ESI†). In addition, the DSC results revealed melting temperatures of 264 °C and 214 °C for BTDI-OCF₃ and BTDI-CF, respectively, as shown in Fig. S1b (ESI†).

Electrochemical and optical properties

The electrochemical properties of BTDI-OCF₃ and BTDI-CF are studied by cyclic voltammetry (CV) in CH₂Cl₂ solution; the corresponding cyclic voltammograms are presented in Fig. 1a. As expected, both BTDI derivatives exhibited two reversible reduction peaks, indicating their typical n-type charge transfer properties. The reduction potentials of BTDI-OCF₃ and BTDI-CF estimated from the onsets of the first reduction peak are

Scheme 1 The synthetic routes of BTDI-OCF₃ and BTDI-CF.

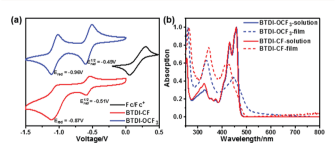


Fig. 1 (a) Cyclic voltammograms of BTDI-CF, BTDI-OCF₃ and ferrocene in DCM solution. (b) UV-vis absorption spectra of the BTDI derivatives in thin films and DCM solution

-0.49 eV and -0.51 eV, from which nearly-similar LUMO energy levels of -4.13 and -4.11 eV versus ferrocene were deduced. The photophysical properties of BTDI-OCF₃ and BTDI-CF were investigated in solution and thin-film form (Fig. 1b). Similar ultraviolet (UV) absorption bands were observed for both of these compounds in dichloromethane (DCM) solution. However, the absorption spectrum of the BTDI-OCF3 thin-film showed a slight bathochromic shift with respect to BTDI-CF, which could be attributed to the difference in the molecular packing geometry of these materials. These results suggest that BTDI-OCF₃ may exhibit stronger intermolecular interactions in the solidstate thin-film. The optical bandgap obtained from the absorption onset of thin-film spectra is estimated to be 2.43 eV for BTDI-OCF₃ and 2.53 eV for BTDI-CF. On the other hand, density functional theory (DFT) calculations were performed at the B3LYP/6-13G(d) level by using the Gaussian 09 program package, which predicted nearly-similar LUMO levels of -4.25 eV and -4.28 eV for BTDI-OCF₃ and BTDI-CF, respectively. Both the experimental and theoretical results revealed comparable LUMO levels of these compounds, indicating that the introduction of different fluorinated substituents at the N_1N' -positions may not greatly affect the LUMO energy level of the resulting material. The corresponding electrochemical and optical properties of BTDI-OCF₃ and BTDI-CF are summarized in Table 1.

Device fabrication and characterization

OTFTs were fabricated by thermal deposition at various substrate temperatures (T_{sub}) to optimize the device performance since T_{sub} could influence the molecular packing, grain size, grain boundaries and defects in the thin film. The device performance of OTFTs based on BTDI-OCF3 and BTDI-CF was tested in a glovebox; the corresponding parameters are summarized in Table 2, and the data for devices tested in air are provided in the ESI.† Both of these materials showed unique electron transfer characteristics and the BTDI-CF based device exhibited higher mobility than BTDI-OCF₃.

We observed that when T_{sub} is increased, the mobility of both materials is enhanced; the mobility of BTDI-OCF₃ showed a slight dependence on the T_{sub} , while that of BTDI-CF increased 2-fold at $T_{\rm sub}$ = 80 °C. This will be further explained by the thin film morphological study. Interestingly, the threshold voltage (V_{th}) of the devices based on these materials showed a contrast shift trend with an increase in the T_{sub} , as can be observed from the transfer curves (Fig. 2). The $I_d^{1/2}$ - V_g curves of the fabricated devices are close to linear, indicating that the mobility is independent of the gate voltage when the influence of water and oxygen is eliminated, which is ideal for logic circuits. The $I_{\rm on}/I_{\rm off}$ ratios of both devices are improved since $I_{\rm off}$

Table 2 Device performance summary of the OTFTs tested in a glovebox

	$T_{\rm sub}/^{\circ}{\rm C}$	$\mu_{\rm e}/{\rm cm}^2~{\rm V}^{-1}~{\rm s}^{-1}$	$I_{\rm on}/I_{\rm off}$	$V_{\rm th}/{ m V}$
BTDI-OCF ₃	RT^b	$6.47 \times 10^{-3a} (7.05 \times 10^{-3})^c$	10^{3}	-35
BTDI-OCF ₃	60	$8.94 \times 10^{-3} (9.77 \times 10^{-3})$	10^4	-43
BTDI-OCF ₃	90	$9.21 \times 10^{-3} (1.11 \times 10^{-2})$	10^{4}	-50
BTDI-CF	RT	$1.50 \times 10^{-2} (1.58 \times 10^{-2})$	10^{5}	-50
BTDI-CF	50	$3.58 \times 10^{-2} (3.72 \times 10^{-2})$	10^{6}	-33
BTDI-CF	80	$3.68 \times 10^{-2} (3.84 \times 10^{-2})$	10^{7}	-15

^a Average mobility. ^b Room temperature. ^c Highest mobility.

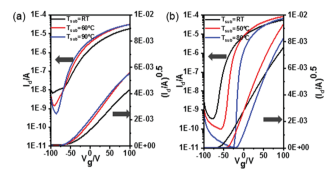


Fig. 2 Transfer characteristics of (a) BTDI-BOCF₃ and (b) BTDI-CF based OTFTs at different T_{sub}

decreased with an increase in the T_{sub} . Overall, the device performance is observed to be enhanced by elevating the T_{sub} .

The stability of the fabricated devices is investigated under different conditions. The electrical stability of the OTFTs is evaluated by a successive scan process in a glovebox. As can be seen in Fig. 3a and c, the drain-source current of BTDI-OCF₃ slightly increased during the scan process, while V_{th} gradually shifted in the negative direction. This improvement in I_d could be attributed to the reduced electron trap density during constant scanning. For the BTDI-CF based device, after 100 cycles of scanning, the I_d - V_g curve almost remained unchanged, which showed superior stability of the device performance under N2. This certified the remarkable operational stability of the BTDI-CF based device in N2 and also indicated that the device would potentially work stably for a long time upon encapsulating it effectively. Moreover, the air-stability is also investigated by measuring the device in an ambient environment. The performance of the OTFTs tested in air initially decreased a little compared with those tested in a glovebox, illustrating that the fluoro-alkyl could protect the device against oxygen and moisture to a certain extent. However, the device performance degraded gradually with time when stored in air since oxygen and moisture could slowly diffuse into the active layer. Vth of both of the devices gradually shifted in the positive direction when stored in air,

Table 1 The electrochemical and optical properties of BTDI-OCF₃ and BTDI-CF

			Experimental		Calculated					
Compound	$E_{\rm red}^{-1/2}/{\rm V}$	$\lambda_{\rm on}/nm$	$E_{\text{HOMO}}/\text{eV}$	$E_{\mathrm{LUMO}}/\mathrm{eV}$	Gap/eV	$E_{\text{HOMO}}/\text{eV}$	$E_{\mathrm{LUMO}}/\mathrm{eV}$	Gap/eV	λ_e/eV	VEA/eV
BTDI-OCF ₃ BTDI-CF	-0.49 -0.51	510.3 489.5	$-6.56 \\ -6.64$	$-4.13 \\ -4.11$	2.43 2.53	-7.03 -7.30	$-4.25 \\ -4.28$	2.78 3.02	0.397 0.330	2.806 2.809

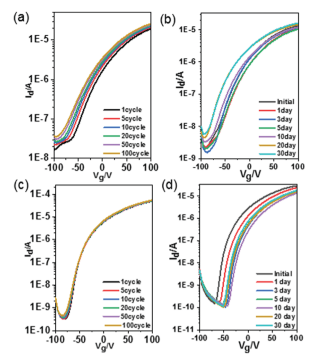


Fig. 3 Operational stability of (a) BTDI-OCF₃ and (c) BTDI-CF based OTFTs under N2. Transfer curves of (b) BTDI-OCF3 and (d) BTDI-CF based OTFTs stored and tested in air

which was induced by the reaction of oxygen with residual -OH in the dielectric layer. 41 Moreover, oxygen and water caused extra traps for electron transfer, which led to a drop in the mobility. The devices could still maintain n-type characteristics comparable to its initial behavior for a long time exposure to air. The scope of degradation of BTDI-CF was a little larger since the three methane groups (-CH₂-) negatively influenced the dense packing of the perfluorooctyl, but the device performance finally stabilized after ten days, as displayed in Table S1 (ESI†). Furthermore, OTFTs based on both BTDI-CF and BTDI-OCF₃ exhibited almost no hysteresis when tested in N₂ and small hysteresis in air; however, the BTDI-C6 based device showed hysteresis in N2 and much larger hysteresis in air with a higher decay of $I_{\rm on}$ (Fig. 3). In general, the hysteresis is mainly caused by water and oxygen. To our knowledge, the fluroalkyl substituted BTDIs could exhibit densely packed molecular structures, which prevent water and oxygen from penetrating into the active layer, thereby reducing the hysteresis in the corresponding devices. This significant improvement in the device performance compared with our previous work³⁹ demonstrates that -C₃H₆C₈F₁₇ and p-(trifluoromethoxy)benzyl can kinetically protect the device performance (Fig. 4).

Structural and morphological analysis of thin films

It is widely accepted that the morphology and the microstructure of thin-films are the key factors affecting the charge mobility. To understand the mobility difference between **BTDI-CF** and **BTDI-OCF**₃ at various T_{sub} , we analyzed the morphologies of the corresponding thin films by atomic force microscopy (AFM) and X-ray diffraction (XRD). The out-of-plane

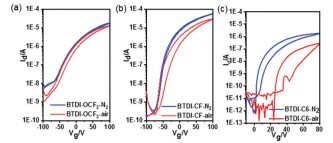


Fig. 4 Transfer characteristics of (a) BTDI-OCF₃, (b) BTDI-CF and (c) BTDI-C6 based OTFTs under different testing conditions

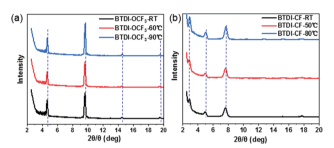


Fig. 5 XRD patterns of (a) BTDI-OCF₃ and (b) BTDI-CF thin films deposited on OTS-treated SiO_2/Si at different T_{sub} .

XRD patterns of BTDI thin films deposited at different T_{sub} are shown in Fig. 5. For BTDI-OCF₃, diffraction peaks up to the 4th order were observed at $2\theta = 4.6^{\circ}$, which correspond to (001) reflections according to the single-crystal data. The interlayer distance (d-spacing) estimated from the primary peak at $2\theta = 4.6^{\circ}$ is 1.93 nm, which is close to the unit cell vector c-axis of 1.97 nm obtained from single crystals. The molecular length of BTDI-OCF₃ (Fig. S4, ESI†) is 2.2 nm, which is similar to the step-height of the AFM image (Fig. S5, ESI†). These results indicate that BTDI-OCF3 molecules are oriented vertically with respect to the OTS-treated SiO2/Si substrate. Generally, this kind of molecular orientation in thin films on the substrate is beneficial for carrier transport. Besides, the intensity of the diffraction peaks exhibited a slight enhancement as the T_{sub} was increased from RT to 90 °C, indicating improved thin-film quality. This is consistent with the experimentally observed mobility of the BTDI-OCF3 based OTFTs (see Table 2). In contrast, diffraction peaks of the BTDI-CF thin films up to the 3rd order are clearly visible and the d-spacing obtained from the primary peak at $2\theta = 2.86^{\circ}$ is 3.07 nm, which corresponds to its longer molecular length, as evident in Fig. S4 (ESI†). Obviously, the d-spacing of BTDI-CF is larger than that of BTDI-OCF3, which is attributed to its long fluoroalkyl chain. As expected, a slight increase in the peak intensity of the BTDI-CF thin films is observed after increasing the T_{sub} , which is in good agreement with the mobility observations of the BTDI-CF based OTFTs.

In addition, the temperature-dependent thin film morphologies of BTDI-OCF₃ and BTDI-CF thin films were studied by AFM, as shown in Fig. 6. Among them, BTDI-OCF₃ showed a welldefined terrace structure in thin-films with good crystallinity, demonstrating a remarkable improvement in the morphology

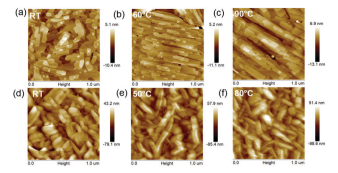


Fig. 6 AFM images of (a-c) BTDI-OCF₃ and (d-f) BTDI-CF thin-films deposited at different T_{sub}

upon increasing the deposition temperature. As displayed in Fig. 6a-c, the step-heights of the terraces were measured to be approximately 2.2 nm, which is in accord with the corresponding molecular length (Fig. S4, ESI†). These observations strongly support the aforementioned XRD analysis of the molecular arrangement on the substrate. Moreover, the grain size was significantly increased with increasing the T_{sub} . In general, well-defined thin-films with large grain size and good crystallinity are beneficial for efficient charge transport, suggesting that the mobility of BTDI-OCF₃ based OTFTs should increase with an increase in the T_{sub} . The thin-film morphology of BTDI-CF was observed to be considerably different from that of BTDI-OCF3. The BTDI-CF thin-film exhibited a wormlike structure with a grain size of 100-200 nm in length, as shown in Fig. 6d. In this case, T_{sub} only exhibited a slight effect on the thinfilm morphology. As depicted in Fig. 6e, the grain size was slightly increased after increasing the $T_{\rm sub}$ from RT to 50 °C. However, the film morphology hardly changed after continuously raising T_{sub} (Fig. 6f), indicating that the device performance would remain unchanged at higher deposition temperatures. As expected, the above investigations are consistent with the observed mobility tendency of BTDI-OCF₃ and BTDI-CF based OTFTs, as summarized in Table 2.

Conclusions

In summary, we have successfully synthesized two kinds of fluoro-alkyl substituted isothianaphthene materials, BTDI-OCF₃ and BTDI-CF. Both of these BTDI derivatives have similar LUMO energy levels and optical properties since the substituent on the nitrogen atom does not greatly affect the LUMO energy. However, the molecular packing and thin film morphologies are quite different due to the significant difference of the alkyl-chain lengths. This further influenced the device performance; the **BTDI-CF** based OTFT showed higher mobility $(0.0368 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1})$ with excellent operational stability in a glovebox, while the BTDI-OCF₃ based device demonstrated highly ordered molecular packing, which resulted in enhanced air-stability. Notably, the hysteresis phenomenon and current decay tested in air compared with those under N2 of the OTFTs based on both of these materials exhibited great improvement as compared to the BTDI-C6 reported in our previous work, indicating better stability of these newly developed fluoric BTDI derivatives.

Moreover, the solubility of these compounds is much higher than the NDI analogues with the same substituent. Further investigation of solution processing of OTFTs based on these materials is underway in our lab.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

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