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Thiophene-based push-pull chromophores for small molecule organic solar cells (SMOSCs).

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The last decade has witnessed a rapid progress of organic photovoltaics boosted by the design and synthesis of novel π-conjugated small donor-acceptor molecules (mainly thiophene-based chromophores) and by the control and optimization of both device processing and fabrication. Although some important progress has been reached, current challenges remain to further improve their efficiency, durability and cost-effectiveness in order to compete with silicon-based solar cells. The review will provide to the scientific community both general and deep information on the structure-properties relationships related to the photocurrent efficiencies comprising a detailed I/V characteristics. It will highlight guidelines for designing new efficient and emerging alternatives to conjugated polymers on the basis of thiophenic chromophores representing, to date, the most widely used class of organic materials for such purpose) as well as important information on device processing or fabrication factors that could influence their performances.

Introduction.

Organic solar cells (OSCs) constitute the third generation solar cells technology and are nowadays of crucial interest because they can offer a good alternative to high cost solar cells based on silicon. Indeed, organic materials display several advantages compared to their inorganic counterpart, such as for instance a lighter weight, a facile chemical tailoring to fine tune their optoelectronic properties and an ease of processability, opening thus the possibilities of developing large-area and cost-effective solar cells based on ecofriendly technologies.

The first attempts to fabricate organic solar cell were described in the early 80’s by Chamberlain from thin organic layers sandwiched between two dissimilar electrodes. Such devices have shown very low solar power conversion efficiency up to 0.3% (PCE) that strongly depends on the nature of the electrodes. A major breakthrough was achieved in 1986 by Tang with the possibility of producing efficient solar cells based on organic semiconductors. A two-layer device fabricated from a copper phthalocyanine (CuPc, as donor material) and perylene tetracarboxylic derivate (PV, as acceptor), end-capped respectively by an ITO and Ag electrodes, led to a PCE value near 1%. Interestingly, in such devices, the charge generation efficiency was found to be relatively independent of the bias voltage. The donor-acceptor interface (p-n junction) provides efficient exciton dissociation in comparison to single layer solar cells. These findings stimulated new works in the field of organic and polymer-based solar cells. A new milestone was reached with the discovery of C60 as a highly efficient acceptor. Although the development of new acceptor materials is strongly requested and seems to be promising, fullerenes C60, C70 and their analogues PC61BM, PC71BM ([6,6]-phenyl-C61-butyric acid methyl ester) and [6,6]-phenyl-C71-butyric acid methyl ester respectively) are readily used and became a traditional acceptor part in OPV cells.

In spite of achieving good photovoltaic conversion efficiencies with the so-called planar heterojunction cells (PHJ), their performances are rather limited due to the small area (interface) contributing to the photovoltaic effect. To circumvent this dilemma, researchers have turned out their attention to the development of bulk heterojunction (BJH). In such structures the p-type and n-type materials are co-deposited, in a determined ratio, to ensure a greater dissociation irrespectively to the thickness. Nevertheless, despite to be a very attractive solution, numerous problematic originate and need to be solved to ensure perfect reproducibility of the devices structures.

Among them, batch-to-batch reproducible synthesis is somewhat difficult to implement, that is why since very recently scientists have paid more and more attention in the use of small organic molecules. Using this approach new OPV devices have been fabricated and fast-optimized leading to power conversion efficiency (PCE) values as high as 11%. For characterizing overall performance of solar cells devices it is appropriate to use few parameters. First of them is an open-circuit voltage, $V_{OC}$, which shows the maximum possible voltage which could be extracted from a device. In first approximation $V_{OC}$ is determined by an energy difference between LUMO level of an acceptor molecule and HOMO of donor one. Keeping in mind that in most cases fullerene derivatives are used as an acceptor, a low HOMO level of donor is responsible for high $V_{OC}$. The maximum current delivered by a device in short circuit conditions, $I_{SC}$, characterize the possibility of charge generation by the cell. In consequence, it depends from other factors, namely light absorption properties and effectiveness of exciton separation. While amelioration of optical properties usually demands a low bandgap of a donor component, the effective charge separation could not occur if LUMO of donor is less than $\approx 0.3$ eV higher than LUMO of acceptor. So, obviously, there is a competition to find an optimal donor molecule with a low enough HOMO level, a high enough LUMO and at the same time not a large bandgap between them. In OC or SC conditions photovoltaic cell do not produce any power ($W = V \times I$). The operating point is a condition with the maximum power and values of $V_{m}$ and $I_{m}$ lower then $V_{OC}$ and $I_{SC}$ respectively. A ratio between products $V_{m} \times I_{m}$ (means maximum achieved power) and $V_{OC} \times I_{SC}$ (means maximum possible power) is called a fill factor, FF, and in generally characterizes the efficacy of the cell structure. It could depend on very various factors among which the active layer morphology and charge carrier mobilities are very important. So, solubility properties of active compounds and their solid state electrical properties should also be taken in
consideration during a molecular engineering of new organic materials for photovoltaic application.

In this article, we will focus on various criteria that influence power conversion efficiencies in small organic molecules, mainly based on the thiophene skeleton, used in organic solar cells from a chemist’s viewpoint with the aim to introduce both a variety of exotic organic materials and insights that are potentially applicable to small molecule organic solar cells.

D-A type.

Despite their extensive uses as key-components in the field of Graetzel solar cells (i.e. DSSC), D-A structures have gained only recently a strong interest for OPVs devices.

Among them, triarylamine-based conjugated chromophores end-capped with acceptors, such as di- or tri-cyanovinyl groups, are some representatives’ D-A structures that have been largely tested.\(^7\)\(^8\) Elongation of the π-conjugated system between the donor and the dicyanovinyl acceptor edge substituents by the increment of thiophene units (from 3 to 6 units) does not affect significantly the \(\lambda_{\text{max}}\) on such structures. Interestingly unexpected, the highest molar extinction coefficient was attained for the tetrathiophene (4T) analogous derivative. As anticipated consecutive change of the acceptor moiety from di- to tri-cyanovinyl (DCV, TCV) leads to a significant bathochromic shift, due to an increase of the electron withdrawing strength, (from 514 to 629 nm and 524 to 643 nm for 2T and 3T derivatives, respectively)\(^9\) associated with a reduced molar extinction coefficient. Modification of the donor unit with a fluorene or heterofluorene such as carbazole constitutes an attractive way to tailor and enhance optical properties. Thus, the substitution of a benzene ring on the triphenylamine (TPA) fragment by a fluorene leads to a slight bathochromic shift with a concomitant increase of the PCE values (Scheme 1).

Scheme 1

Modification of the donor part by a dendritic-like structure based on two TPA units (3a and 3b) has a dramatic effect on electrochemical properties. It was shown that such changes reduce the HOMO energy level to 0.15-0.16 V as well the LUMO to 0.11-0.13 V compared to their linear analogs 3a relied on the donor strength. This fact results in a higher energy band gap estimated from cyclic voltammetry and correlates well with the observed optical gap. The highest PCE value (2.67 %) was obtained for the device based on the compound 1e prepared by bilayer vacuum deposition and annealed for 20 min at 100°C.

Further substitutions on the TPA part were envisioned by Ko et al.\(^10\)\(^11\) in order to fine tune the optical and electrochemical properties of the donating part (Scheme 2).

Scheme 2

It can be noticed herein that shorter π-conjugated spacer lengths provide, in that case, comparable conversion efficiencies up to 2.46 % for 5c, which can be attributed to higher \(J_{\text{SC}}\) value (8.91 mA·cm\(^{-2}\)) compared to 1a (4.92 mA·cm\(^{-2}\)) due to an optimized performance architecture in the BHJ blend with PC\(_{71}\)BM compared to the PHJ with C\(_{60}\) for 1a. Further optimization, of the phase segregation upon addition of 1-chloronaphthalene (CN) as an additive, leads to a significant enhancement of the conversion efficiency up to 3.66 %.

Additional improvement of the photophysical properties has been achieved with the fused benzothiophene 6 (Scheme 3) instead of 2-thienylbenzene as spacer.\(^12\) Therefore, compared to 5c, better optical and hole-mobility properties are attained. Subsequent device made from a blend of 6 and PC\(_{71}\)BM (1:3) with TiO\(_x\) layer affords 4% conversion efficiency.

Scheme 3

Silole bridged bithiophene (DTS) spacer has been used in order to gain further insights for the design of SMOSCs in the above TPA compounds series.\(^13\) Thus, introduction of the DTS moiety along the π-conjugated spacer (7) does not change significantly the optical properties (\(\lambda_{\text{max}} = 537 \text{ nm} \epsilon = 45 800 \text{ M}^{-1} \text{ cm}^{-1}\)) compared to 5c (\(\lambda_{\text{max}} = 536 \text{ nm} \epsilon = 32 174 \text{ M}^{-1} \text{ cm}^{-1}\)) but results in a refinement of the HOMO and LUMO energy levels which are lowered by almost 0.5 V. (Scheme 4).

Scheme 4

Fabrication of a planar-mixed heterojunction (PMHJ), by vacuum deposition, with 7 and fullerenes derivatives in the molar ratio (1:1) achieves conversion efficiency up to 2.69 % and 3.82% with C\(_{60}\) and C\(_{70}\) respectively. The higher PCE originates from the high V\(_{\text{OC}}\) potentials achieved in these devices (0.88 V and 0.83 V respectively compared to 0.774 V for 5c) as a consequence of the energy levels optimization.

Besides the common covalent bridging strategy, the design of “dwarf” chromophores (i.e. minimizing the size of the chromophore) has recently emerged to be a possible alternative for controlling the batch-to-batch reproducibility as well as to get low cost systems associated with high purity and optimized photophysical properties.\(^14\)\(^15\) Therefore, both Roncali’s group and Würthner’s group have demonstrated the validity of this concept throughout the synthesis of the smallest push-pull dyes based on thiophene rings (Schemes 5 and 6).

Scheme 5

Optimized chemical structures have been obtained by varying only the acceptor edge substituent. On the basis of this series the best performances were achieved with the compound 9c leading to PCE up to 3.0% in blended films with PC\(_{60}\)BM 55 wt %. Further devices optimizations were made, for instance replacing PC\(_{60}\)BM by PC\(_{71}\)BM, and the hole-collecting PEDOT: PSS layer by MoO\(_x\), and using a Ba/Ag metal electrode leading to attain remarkable PCE value of 4.5%.

Scheme 6

However, by changing the donor part from dialkylyamino to TPA, Roncali et al. have obtained a reasonable value of 2.53 % of conversion efficiency on un-optimized architectures devices (Scheme 6). In furtherance of their continuous efforts on the understanding of structure-properties relationships, the covalent bridging strategy was applied on the acceptor part in order to optimize the light harvesting properties (Scheme 7). Thus
substitution of the dicyanovinyl (9) by the thieno-fused-dicyanoindene (10) moiety leads to a 110 nm red shift, as expected. Although this bathochromic shift is accompanied by a two-fold decrease of the molar extinction coefficient an enhanced PCE (2.97 %) was clearly evidenced. Additional work on fine-tuning properties of the donating group has been done in order to give a better accuracy of the HOMO levels and bandgap control as well as to gain further information on the design of such chromophores (Scheme 7).\(^{13}\) Shortening the \(\pi\)-conjugated spacer results in a 55 nm hypsochromic shift (from 9 to 11) while an average shift of 27 nm was observed for compounds 12 to 13 because of the lower resonance energy of the thienyl ring. As a consequence, higher FF and \(J_{SC}\) characteristics were obtained beside a decrease of the \(V_{OC}\) leading to an increase of the photocurrent (1.60–1.70 %). Interestingly, the best performances were obtained for compound 14 evidenced by the highest \(J_{SC}\) (6.80 mA cm\(^{-2}\)) and PCE (1.92 %) values, whereas the bridged carbazole analogous 15 did not improve the photocurrent.\(^{16}\)

Scheme 7

Additionally it has been demonstrated that 16, a structural isomer of 10, exhibits a higher dihedral angle leading to a less planar \(\pi\)-conjugated system. According to the structural modification of the chromophores general behaviors can be deduced that are consistent with the optical and electrochemical experiments (Scheme 8).\(^{17}\) For instance, bridging the dicyano group leads to a positive shift of the reduction potential (from 9 to 10 and 16-19) due to a better stabilization of the radical anion upon reduction attributed to the formation of the cyclopentadienyl anion radical on those structures. Replacement (from 10 to 17-18) of the phenyl ring by thienyl or introduction of methoxy groups significantly increases the HOMO levels. Optical observations were in agreement with the cyclic voltammetry experiments, thus the narrow bandgap for 17 and 18 compared to 10 is related to an increase of the HOMO level. Evaluation of the performances of these chromophores has been achieved using a bilayer planar heterojunction with the following architecture ITO/PEDOT: PSS (40nm)/10,16,19 (35nm)/C\(_{60}\) (30nm)/Al (100nm). The highest performance give a PCE of 2.97% for 10 (\(V_{OC}= 0.97\) V, \(J_{SC}= 5.32\) mA cm\(^{-2}\) and FF= 0.52) while 16 exhibits the lowest PCE according to the optical and electrochemical data (0.57%, \(V_{OC}= 0.71\) V, \(J_{SC}= 2.07\) mA cm\(^{-2}\) and FF= 0.35).

Scheme 8

Further interesting insights were obtained from the previous D-A series by the insertion of a second acceptor along the \(\pi\)-conjugated spacer. The elegant combination of two different acceptors gives rise to chromophores with the D-A-A configuration.\(^{18,19,20}\) These D-A-A structures exhibit both narrow bandgap and lower HOMO levels compared to their D-A counterparts (Scheme 9). Thus, insertion of the benzothiadiazole (BT) unit in the D-A structures fosters the creation of quinoidal mesomeric forms over the \(\pi\)-conjugated skeleton resulting to a bathochromic shift of the maximum of absorption. Moreover, judicious choices of the substituents as well as the aromatic rings over the \(\pi\)-conjugated backbone are key parameters to control both energy levels and bandgap. For instance, substitution of the phenyl ring by a thienyl leads to a substantial red shift of 90 nm of the maximum of absorption (i.e. 22a→20a and 22b→20b).

Scheme 9

**A\(\_\)D type**

Optimization of the light harvesting properties of the chromophores is a crucial point that needs to be addressed in order to improve the conversion efficiencies. An elegant solution to tackle this problem has been developed by several groups, from 3D-structures mainly based on the TPA core. Indeed, TPA structures are of peculiar interest for this problematic, as they have already demonstrated their ability to generate efficient linear chromophores and allow the possibility to introduce various acceptor edge groups on the same skeleton. Hence, through an iterative pathway, a star-shaped series of TPA has been synthesized by Roncali et al. in which the \(\pi\)-conjugated arms offer variability in term of length, nature and acceptor edge groups. Using this strategy, disymmetric structures are therefore attainable with both a controlled acceptor load and morphology structures (Scheme 10).

Scheme 10

As demonstrated on bulk heterojunction devices, from the structures 23 to 25, a noticeably linear improvement in power conversion efficiencies is observed ranging from 0.46 to 1.17 % respectively.\(^{22}\) Explanations of this observation originate from (i) the fact that there is a linear dependency of the intramolecular charge transfer (ICT) with the number of acceptors present over the \(\pi\)-conjugated backbone and (ii) an enhanced \(V_{OC}\) (23: 0.48 V; 24: 0.72 V; 25: 0.89 V; i.e. lower HOMO levels). Moreover increasing the number of acceptor groups promotes different spatial arrangement in the solid-state with strong intermolecular D-A interactions that can affect also the PCE. As expected, the highest \(V_{OC}\) (0.96 V) was obtained for the compound 26 containing three dicyanovinyl end-groups, however, reduced PCE (1.02 %) is achieved due to a lower filling factor compared to 25. Furthermore, optimizations of these rudimentary devices were undertaken to boost the power conversion efficiencies. For instance, addition of a thin film of LiF between the C\(_{60}\) acceptor...
layer and the Al electrode provides an impressive increase of the PCE from ~ 1% to 1.85% (with $V_{OC} = 1.15 \text{ V}$ and $J_{SC} = 4.59 \text{ mA} \text{cm}^{-2}$)\(^\text{23}\) for the device made from 26.

Taking into account these aspects the strategy is valid and worth pursuing as the observed synergistic effects lead to considerable improvement of the PCE.

With the aim to enhance the photovoltaic output signals, several structural modifications have been conducted on the TPA derivatives based on the previous obtained results. Interestingly, contrary to polymeric or polycrystalline organic structures, amorphous TPA based chromophores have the advantage not to require control of the molecular orientation. First attempts have been focused basically on the hole-transport properties by engineering TPA derivatives with linear p-type organic semiconductors acting as the donor material in PV devices. Thus, structure-like 28 (Scheme 11) has been synthesized and tested as donor material in a bilayer heterojunction with C60 as the acceptor. PCE not exceeding 0.32% ($V_{OC} = 0.67 \text{ V}; J_{SC} = 1.70 \text{ mA} \text{cm}^{-2}$)\(^\text{24}\) has been attained with such compound as mainly a consequence of a low filling factor. Although low PV efficiency has been obtained, 28 demonstrated good hole-mobility properties up $\mu_h = 1.10 \times 10^{-2} \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$ which is very promising for PV applications.

Scheme 11

Subsequent replacement of one or two arms by unit end-capped with DCV (29 and 30 respectively, Scheme 12) as acceptor has led to a drastic improvement of the PCE (1.80% and 2.02% respectively) due to a large increase of the short-circuit current density and $V_{OC}$ (29: $V_{OC} = 0.85 \text{ V}; J_{SC} = 5.30 \text{ mA} \text{cm}^{-2}$; 30: $V_{OC} = 1.07 \text{ V}; J_{SC} = 5.83 \text{ mA} \text{cm}^{-2}$) that are intrinsically correlated to the ICT.\(^\text{25}\)

Scheme 12

To achieve improved results and gain further insights into structure-properties relationships, bridged-thienyl spacers have been introduced in the TPA molecular skeleton. Covalently bridged-spacers are well-known to exalt photophysical properties compared to their un-bridged analogues.\(^\text{26}\) For instance, Li\(^\text{27}\) and co-workers used 3,6-di-hexyl-thieno[3,2-b]thiophene as electron relay between the TPA donor part and the dicyanovinyl end groups for the construction of star-shaped structures (Scheme 13). Devices made from a blend of 31 and PCBM or PC7BM in weight ratio 1:2 with a Al/Ca as cathode leads to high-performance devices under standard illumination conditions (respectively 1.63% and 2.87%) associated with exceptional high $J_{SC}$ values (4.66 to 6.80 mA cm\(^{-2}\)) and $V_{OC}$ (0.90 V and 0.96 V) that are among to the best reported in literature.

Scheme 13

To achieve readily processable active materials, Lin et al. have synthetized star-shaped structures based on the TPA scaffold using alkyl-cyanoacetate (ACA) as acceptor.\(^\text{28}\) Incorporation of ACA aims at introducing simultaneously solubilizing and acceptor groups on the molecular π-conjugated backbone and avoids tedious multistep reactions. As a result, successful synthesis of 32 has been achieved associated with a good solubility in common solvents and high molar extinction coefficient (7.3 $10^3 \text{ M}^{-1} \text{ cm}^{-1}$) (Scheme 14). Absorption maximum is red-shifted by 65 nm compared to the unsubstituted analogue 28 because of the intramolecular charge transfer. Additionally, 32 retains relatively high hole-mobility, measured by the space charge limited current (SCLC) method, up to 1.5 $10^{-3} \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$ guaranteed effective charge carrier transport and reduced photocurrent loss. Optimized BHJ solar cells made from 33 in conjunction with PC7BM (best weight ratio 1:2) as the electron acceptor gave under standard illumination condition AM1.5 a power conversion efficiency, without any post-treatments, of 3.60% ($V_{OC} = 0.89 \text{ V}, J_{SC} = 7.30 \text{ mA} \text{cm}^{-2}$).

Furthermore, the photophysical properties of the star-shaped chromophores 32 were compared with linear and 2D analogous chromophores (33–34 Scheme 15) in order to investigate the structural effects on the above properties.\(^\text{29}\)

Scheme 14

As expected, on the linear analogous derivatives, increment of the number of thiophene unit from 1 to 3 (33a to 33c) led to a 20 nm red shift of the maximum of absorption whereas increasing the number of arms around the TPA core it remains almost unchanged (32, 33a and 34 494, 498, 494 nm respectively). In addition, the energy differences between the HOMO levels of the chromophores and the LUMO level of PC7BM are sufficient to ensure the exciton dissociation and lead to a high $V_{OC}$ (from 0.87 to 0.99 V). Relative to 33a-33c, chromophores 34 and 32 exhibit the highest PCEs (at same weight ratio with PC7BM 1:2), attaining 4.03% for 34 ($J_{SC} = 9.03 \text{ mA} \text{cm}^{-2}$), due to stronger absorption and higher mobilities.

Scheme 15

Replacing the thienyl unit by an ethenic bond can be used to reduce the aromatic character of the π-conjugated arms in order to design more efficient chromophores. The ternary substituted TPA-based compound 35, end-capped with an octyl cyanoacetate (CA) group, was synthesized and evaluated by Li et al. in BHJ solar cells and compared to its analogue 32 (Scheme 16).\(^\text{30}\) Due to the less aromatic character of the arms better charge transfer occurs between the central donor and external acceptor groups. As expected, compound 35 possesses broad and strong visible absorption band which is red-shifted by 24 nm related to 32 and lower HOMO energy level. The photovoltaic performance was collected from a BHJ solar cell device with the following architecture ITO/PEDOT: PSS (30nm)/35:PC7BM (70nm)/Ca/Al (120 nm). The performances were recorded after a thermal annealing of the photosensitive layer at 80°C (10min.) at different blend weight ratios (from 1:1 to 1:3, w/w) under AM1.5 conditions. The best device, based on a blend at 1:3 ratio, exhibited a PCE of 2.1% ($V_{OC} = 0.91 \text{ V}, J_{SC} = 4.64 \text{ mA} \text{cm}^{-2}$ and FF= 0.50). The reduced photoconversion compared to 32 was ascribed to a decrease of the short-circuit current due to lower hole-mobility ($\mu_h = 2.03 \times 10^{-4} \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$ vs $\mu_h = 1.5 \times 10^{-3} \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$).

Scheme 16
Motivated by the obtained results and continuing their efforts on the structure-properties relationships of star-shaped chromophores Li et al. investigated a novel class of trisubstituted TPA. Those are based on 5-vinyl-4,4'-diethylbithiophene as π-conjugated spacer, in order to afford efficient solution-processable OSCs materials associated with good film forming properties and are end-capped with a dicyanovinyl group.\textsuperscript{31} Compared properties of 36 vs 37 show that the latter displays enhanced photophysical properties caused by the insertion of the Vinyl group onto the π-conjugated backbone. For instance 37 exhibits red shift by 40 nm of the maximum of absorption and lower HOMO level (-5.03 vs -5.22 eV for 36) likewise the bandgap (1.61 vs 1.88 eV for 36). Consequently, spin-coated films of either 36 or 37 were used to fabricate BHJ with PC$_7$BM as acceptor (with ratio 1:2 w/w respectively). Improved performances devices were obtained as anticipated for 37 exhibiting a power conversion efficiency of 3.00% ($V_{OC}$= 0.88 V, $J_{SC}$= 7.76 mA·cm$^{-2}$) which is twofold higher than 36 (Scheme 17).

\textbf{Scheme 17}

Besides processability, stability of chromophores is a crucial parameter that needs to be considered. Regarding dicyanovinyl acceptors, acidic hydrogen could be problematic for long-term stability of the active layers. In this context, Min et al.\textsuperscript{32} have designed a new star-shaped TPA 38 possessing an ethyl-dicyanovinyl group as acceptor (Scheme 18), which avoids the presence of acidic hydrogen. Additionally, 38 shows reduced torsional interactions compared to 36 because of lack of steric hindrance of the hexyl chains (leading consequently to lower HOMO level) and presents good solubility. Thus, BHJ made from 38 and PC$_7$BM has resulted in a twofold improvement of the PCE (3.10%) relative to 36 (1.40%) even without post-treatments. Moreover, as already stated, device performances could be enhanced by optimizing the processing conditions such as thermal annealing methods or solvent additives. Therefore, the previous device treated, during the processing step, with 2% of 4-bromoanisole as additive has attained a remarkable efficiency of 4.3% ($V_{OC}$= 0.79 V, $J_{SC}$= 6.41 mA·cm$^{-2}$; $V_{OC}$= 0.71 V, $J_{SC}$= 5.41 mA·cm$^{-2}$ respectively) were attained mainly due to poorer film-forming properties (large molecular size, lower solubility) and weaker ICTs.

\textbf{Scheme 18}

Additional improvements have been achieved regarding the absorption properties compared to 28 by insertion of benzothiadiazole (BT) moieties into the π-conjugated systems of the TPA starburst molecular structures. Indeed, introduction of BT lead, not only, to a broadening absorption spectrum and also to an increase of the thermal stability probably due to the S-N intramolecular bondings.\textsuperscript{33} Moreover, 39 displayed typical p-type semiconductor behavior in air associated with a hole-mobility of $\mu_h= 4.90 \times 10^{-7}$ cm$^2$ V$^{-1}$ s$^{-1}$ which is almost 50 times higher than the reported analog 28.

All these features, associated to good solubility properties, make 39 to be a donor of choice for fabricating solution-processed OSCs. Consequently, a blend of 39 and PC$_7$BM has been deposited by spin coating exhibiting a PCEs as high as 4.3% without any post-treatment (device structure: ITO/PEDOT: PSS/39: PC$_7$BM (1:2, w/w)/Ca/Al, ($V_{OC}$ = 0.87 V, $J_{SC}$ = 9.51 mA·cm$^{-2}$). The high value obtained has been correlated to the surface morphology displaying nano-aggregated domains that are suitable and beneficial to the charge separation. In order to expand the scope of this compounds family, He et al. have developed and tested in BHJ OSCs the derivatives 40a and 40b (40a or 40b: PC$_7$BM (1:3, w/w)).\textsuperscript{34} Extended π-conjugated backbone was obtained by inserting the vinylene groups and TPA end-groups in order to achieve broader absorption. Although similar optical properties to 28 were gained lower photovoltaic performances (PCEs of 1.90% and 1.23% with $V_{OC}$=0.75 V, $J_{SC}$= 6.41 mA·cm$^{-2}$; $V_{OC}$= 0.71 V, $J_{SC}$= 5.41 mA·cm$^{-2}$ respectively) were attained mainly due to poorer film-forming properties (large molecular size, lower solubility) and weaker ICTs.

\textbf{Scheme 19}

As was demonstrated for structure-like 39, thiophene or polythiophene structures can play not only a linker role but can also act as a donor. The 41a star-shaped conjugated molecule possesses a D-A structure with TPA as the donor core unit, benzo[1,2,5]thiadiazole (BT) as the acceptor unit and terthiophene (TT) donor end groups.\textsuperscript{35} Comparatively to the corresponding star-shaped molecules 41b and 41c, 41a shows a red-shifted absorption with a maximum at 538 nm and an absorption edge extending to 650 nm associated to higher hole mobility up $\mu_h= 1.1 \times 10^{-4}$ cm$^2$ V$^{-1}$ s$^{-1}$. The BHJ solar cells based on the device structure ITO/PEDOT: PSS/ 41: PC$_7$BM (70nm)/Ca/Al, with a blend 41: PC$_7$BM at a 1:3 ratio (w/w), exhibited a power conversion efficiency of 2.28% ($V_{OC}$ = 0.82V, $J_{SC}$ = 6.39 mA·cm$^{-2}$ and FF= 0.43) under standard illumination AM1.5 conditions. Compounds 41b and 41c exhibit lower efficiencies, 0.61% ($V_{OC}$ = 0.91V, $J_{SC}$ = 1.54 mA·cm$^{-2}$ and FF= 0.43, with a Ba (10nm)/Al (100nm) cathode) and 1.33% ($V_{OC}$ = 0.81V, $J_{SC}$ = 4.18 mA·cm$^{-2}$ and FF= 0.39) respectively incriminated to lower hole-mobility as well as less π-π interactions in the solid state due to the less planar TPA scaffold related to TT.

\textbf{Scheme 20}

Replacing benzothiadiazole by benzoselenadiazole (BS) into TPA star-shaped skeleton was described recently (Scheme 21).\textsuperscript{38} Selenium atom was used because of its more electron rich character compared to sulfur and induced a red shift in π-conjugated systems relative to their sulfur counterparts. Thereby, compound 42 was designed and synthesized displaying both a good solubility and a broad absorption from 300 to 650 nm similar to that of 39. BHJ OSCs based on blend films of 42 and PC$_7$BM (1:2, w/w) have given rise to photovoltaic performances up to 1.54% ($V_{OC}$ = 0.91 V, $J_{SC}$ = 4.54 mA·cm$^{-2}$; FF=0.37).

\textbf{Scheme 21}

Due to their intrinsic attractive properties, diketopyrrolopyrrole (DPP) have been proven promising materials for the design of efficient OSCs. Thus changing BT or BS acceptors by a DPP moiety in TPA based-materials has led to the design of novel materials that can be used as non-fullerene acceptors for solution-processed OSCs. To this end, Zhan et al.\textsuperscript{39} have synthesized a
soluble chromophore 43 and have therefore demonstrated its potential application in OSCs (Scheme 22). A device structure ITO/PEDOT: PSS/P3HT: 43 (1:1, w/w)/Cu/Al leads to an impressive increase of the VOC of 1.18 V (JSC = 2.68 mA.cm⁻²) associated to a conversion efficiency as high as 1.20%.

**Scheme 22**

Shi and co-workers have recently optimized the photophysical properties of the above DPP architectures by end-capping with substituted or unsubstituted benzene rings (compounds 44a–44c – Scheme 23).⁴³ The latter were investigated as π-conjugated donors in blend films with PC₆₀BM as the acceptor at ratio 1:1 (w/w) (for 44a and 44b) or 1:2 (for 44c). Best performances were obtained after improving the films morphologies by adding 2% of DIO (1,8-diiodooctane) during the solution-process step. Generally, the synthesized chromophores exhibit roughly similar optical and electrochemical behaviors with slight differences relative to the endgroup that influences mainly the π-π stacking in the solid state. The device structures demonstrated that the highest PCE (2.98%) was obtained for 44a and was correlated to its most efficient charge transport properties and higher hole-mobility among the others (44a: μh = 8.63 10⁻⁵ cm² V⁻¹ s⁻¹, VOC = 0.72 V, JSC = 7.94 mA.cm⁻² vs 44b: μh = 2.42 10⁻⁵ cm² V⁻¹ s⁻¹, VOC = 0.76 V, JSC = 5.14 mA.cm⁻² and 44c: μh = 4.43 10⁻⁵ cm² V⁻¹ s⁻¹, VOC = 0.80 V, JSC = 5.83 mA.cm⁻²) as well as a smoother film morphology and an higher exciton dissociation efficiency.

**Scheme 23**

Expanding the π-conjugated system of 39 by embedding an additional thienyl ring has demonstrated to be benefit to improve the light harvesting properties on the TPA-DPP based structures. Indeed, improvement of the photophysical properties were ascribed to an increase of the molecular ordering that inter alia impacts on the absorption properties and on the charge mobilities (Scheme 24).⁴⁴ Consequently, a dramatic enhancement of the photovoltaic performances, more than two fold magnitude, was obtained in favor to 45 (2.95%, VOC = 0.81 V, JSC = 10.02 mA.cm⁻²) compared to 39 (1.38%, VOC = 0.77 V, JSC = 5.08 mA.cm⁻²) in conventional device structures with PC₆₀BM as acceptor (at ratio 1:1 and 1:2 (w/w) respectively). Similar results were obtained when PC₆₀BM was used as acceptor (1.81%, VOC = 0.83 V, JSC = 5.60 mA.cm⁻² vs 1.13%, VOC = 0.84 V, JSC = 4.19 mA.cm⁻²). Moreover, the HOMO (-5.13 eV) and LUMO (-3.32 eV) energy levels suggest that 45 could be used as an acceptor in blended films with P3HT yielding a PCE of 0.63% with an impressive VOC = 1.14 V, (JSC = 1.33 mA.cm⁻²).

**Scheme 24**

Original star-shaped chromophores based on non-TPA cores, with a C₃ᵥ symmetry, are of interest due to the possibility to fine-tune the properties of such structures by modifying for instance the planarity and controlling the solid-state interactions. Based on these considerations, Xiao et al.⁴² and Bo et al.⁴³ have envisioned the use of planar structures such as benzothiophene (BTT) and benzotrisindole (BTI) respectively to develop new star-shaped structures (Schemes 25-26).

Fine-tuning the properties of molecules 46a–46c by a selective increase of the thiophenic units leads to a control of the HOMO level while the LUMO level remains almost unaffected. Compounds 46a–46c exhibit broad optical absorption with similar behaviors; the main difference came from the highest energy band due to the increase of the number of thienyl units. BHJ solar cells were fabricated with the device structure ITO/PEDOT: PSS (45nm)/46: PC₆₀BM or 46: PC₆₀BM /Ca (20nm)/Al (80 nm). Increasing the number of thienyl units led to a significant enhancement of the PCE from 0.16% for 46a (VOC = 0.54 V, JSC = 0.95 mA.cm⁻² and FF = 0.31) to 0.74% for 46c (VOC = 0.69 V, JSC = 2.93 mA.cm⁻² and FF = 0.7) at a 46: PC₆₀BM 1:1 ratio. Changing the blend to 46c: PC₆₀BM (1:2 ratio) improves the conversion efficiency up to 1.36% (VOC = 0.72 V, JSC = 4.13 mA.cm⁻² and FF = 0.46) due to a better intimate donor acceptor mixing and smoother surface.

**Scheme 25**

BTI core has been used for the synthesis of chromophores 47-48 and due to presence of the N-atom it should display a higher donor character relative to BTI. The films made from 47 and 48 display a broad absorption in the visible associated to an optical bandgap of 1.82eV and 2.07eV respectively. From the photoelectrochemical experiments the HOMO and LUMO energy levels have been calculated to be -5.12eV and -3.30eV for 47 and -5.07eV and -3.00eV for 48 attesting their ability to be used as donors in BHJ in conjunction with PC₆₀BM or PC₇₀BM as acceptors. Hence, solar cells devices with the architecture ITO/PEDOT: PSS (40nm)/47-48: PC₇₀BM /LiF (0.5 nm)/Al (100 nm) were prepared and evaluated. The photovoltaic performances were found to be dependent to the ratio of donor to acceptor. The best results were obtained for a ratio of 1:3 with a thickness of the photoactive layer of about 105 nm. Under these conditions, PCEs of 1.95% for 47 (VOC = 0.82 V, JSC = 5.31 mA.cm⁻² and FF = 0.45) and 1.73% for 48 (VOC = 0.65 V, JSC = 8.13 mA.cm⁻² and FF = 0.33) were achieved. However, a post-deposition thermal annealing at 130°C for 5 min. for 48 increases the conversion efficiency up to 2.05% (VOC = 0.72 V, JSC = 6.41 mA.cm⁻² and FF = 0.44) while an improvement for 47 is obtained when 0.5% of CN is used as solvent additive leading to a PCE of 2.29% (VOC = 0.82 V, JSC = 6.00 mA.cm⁻² and FF = 0.47). The enhancement has been incriminated to marked changes of the film morphologies when the post-treatments were performed as attested by the surface and morphology analysis. The results obtained from these structures make them very promising candidates for the elaboration of SMOSCs and need to be further explored.

**Scheme 26**

Recently Wu et al. have envisioned the synthesis of cruciform-like structures originating from the link of two linear chromophores (Scheme 27).⁴⁴ Thus, two new structures 49 and 50 have been designed and synthesized in order to be investigated as potential new efficient donor for organic solar cells. They differ one to each other by the length of the arms. As expected, 50 exhibiting the longest π-conjugated system led to the highest maximum absorption (λmax = 525 nm). Interestingly, from solution (λmax = 495 nm) to thin film (λmax = 499 nm) 49 shown similar
spectra suggesting weak intermolecular contacts in the solid-state while 50 displays a 13 nm red shift due to better \( \pi \)-\( \pi \) interactions. From the solid-state spectra the optical band gap can be deduced to be 2.07 eV and 1.82 eV respectively to 49 and 50. The lower lying HOMO energy levels of 49 and 50 are appropriate to generate large \( V_{OC} \) in the devices.

Conventional structure of ITO/PEDOT: PSS (30 nm)/49 or 50: PC\(_{70}\)BM (1:3, w/w, 90 nm)/Ca (15 nm)/Al (100 nm) have been used for the fabrication of BHJ solar cells. Photovoltaic performances were determined under standard illumination conditions AM1.5. Conversion efficiencies of 0.74% for 49 (\( V_{OC} = 1.09 \) V, \( J_{SC} = 2.47 \) mA cm\(^{-2} \) and FF = 0.27) and 3.14% for 50 (\( V_{OC} = 1.08 \) V, \( J_{SC} = 6.75 \) mA cm\(^{-2} \) and FF = 0.43) were obtained without any treatments. The significant differences of the PCEs in favor to 50 came from the improved short-circuit current and fill factor related to a better phase separation and broader absorption spectrum.

Scheme 27

A – D – A type

Besides the recently developed TPA-based chromophores most efforts have been devoted to the linear oligomeric thiophene \( \pi \)-conjugated systems. On such derivatives the thiophene \( \pi \)-conjugated backbone, is routinely combined with selected acceptors in order to fine-tune their photophysical properties. From a given donor (D) and acceptor (A) different combinations are possible leading to several defined architectures namely D-A, A-D-A, D-A-D etc… Symmetric and disymmetric structures can be easily obtained using iterative synthetic pathways and offer most advantages than polymers because of their higher solubility, monodisperse character and the possibility to better fine tune their properties.

In addition, as evidenced on TPA star-shaped structures, combination of a donor with a given strength acceptor broadens the absorption spectrum and optimizes the light-harvesting properties. Consequently a reduction of \( E_g \) was produced concomitantly with an increase of the \( V_{OC} \) due to a decrease of the HOMO level. All these phenomena are in favor of optimizing the photovoltaic performances.

In this context, association of a sepi thiophene with two different acceptors namely the dicyanovinyl (DCN) and the diethylthiobarbituric (DETA) acid affords two symmetrical (A3D3A, D3A3D etc… Symmetric and dissymmetric structures are possible leading to several defined architectures namely D3A, A – D – A type conjugated backbone, is routinely combined with selected acceptors. Nonetheless, the nature of the terminal acceptors will affect not only the optical properties but also dramatically the solid-state packing as well as on the solubility. That might impact on the final properties (54a, 54b and 54c) respectively. The observed variations can be incriminated to the formation of different grain-sized domains organizations in the blend as evidenced by AFM. For instance, microphotographs of 54b, exhibit well separated domains (assigned to the acceptor-rich and donor-rich parts) associated to a higher roughness (4.05 nm compared to 2.51 nm for 54a and 54c). Comparison of the volume occupied by the side chains (linear or hyperbranched alkyl chains) is of importance on the molecular design of linear systems.

Continuing their studies on the structure-property relationships modulation of the properties have been achieved by varying the nature of the acceptor on symmetrical structures (Scheme 28).

Scheme 28

As expected, the replacement of one DCN by DETA (51 to 53) leads to a 31 nm red shift and subsequent exchange of the second DCN by DETA (51 to 52) leads to another 21 nm red shift of the \( \lambda_{max} \). Surprisingly, it can be noticed that from 51 to 52 a reduction of the band gap is attained while a concomitant diminution of the power conversion efficiency is obtained as the number of DETA increase in a PHJ configuration with a spin-coated donor layer and a vacuum deposited Ca layer (1.64% 51, 1.21% 53 and 0.36% 52 respectively).

From these observations it appears that the molecule desymmetrization affects both the optical properties as well as the packing arrangement in the solid state, which has the dramatic effect on the PCE. Further investigations in BHJ solar cells based on a blend of 51 and PC\(_{70}\)BM (1:1.4 (w/w) ratio) led to an improvement of the PCE value up to 2.45% and 3.7%.

Chen’s group further studied an analogous series by changing the nature of the acceptor. Interestingly, it was found that the alkyl-cyanoacetate (ACA) end-group acts as a quite effective acceptor group with the benefit to improve the film morphologies qualities by changing the nature of the alkyl chain (Scheme 29).

Scheme 29

Thus, compounds 54a, 54b and 54c display similar optical properties in CHCl\(_3\) (absorption maxima ca. 492 nm and absorption coefficients are 7.6×10\(^{4}\) M\(^{-1}\) cm\(^{-1}\), respectively). PCE were evaluated on a BHJ devices, made from a blend of each of these donors and PC\(_{70}\)BM (1:0.5 w/w), leading to high efficiency, up to 4.46%, 5.08% and 4.52% for 54a, 54b and 54c respectively. The observed variations can be incriminated to the formation of different grain-sized domains organizations in the blend as evidenced by AFM. For instance, microphotographs of 54b, exhibit well separated domains (assigned to the acceptor-rich and donor-rich parts) associated to a higher roughness (4.05 nm compared to 2.51 nm for 54a and 54c). Comparison of the volume occupied by the side chains (linear or hyperbranched alkyl chains) is of importance on the final properties (54b and 54c) and has to be considered in the molecular design of linear systems.

Continuing their studies on the structure-property relationships modulation of the properties have been achieved by varying the nature of the acceptor on symmetrical structures (Scheme 30).

Scheme 30

As expected the energy levels and bandgap can be easily fine-tuned on these architectures by changing the strength of the acceptors. Nonetheless, the nature of the terminal acceptors will affect not only the optical properties but also dramatically the solid-state packing as well as on the solubility. That might impact both on the hole and electron mobilities and on the grain size domains. Hence, hole-mobilities were measured by the space charge limited current method (SCLC) on films made from the compounds 55 to 58 giving the following mobilities \( \mu_h = 0.47·10^{-4} \) cm\(^2\) V\(^{-1}\) s\(^{-1}\), \( \mu_b = 3.26·10^{-4} \) cm\(^2\) V\(^{-1}\) s\(^{-1}\), \( \mu_h = 1.50·10^{-4} \) cm\(^2\) V\(^{-1}\) s\(^{-1}\) and \( \mu_b = 0.24·10^{-4} \) cm\(^2\) V\(^{-1}\) s\(^{-1}\) respectively. Bulk heterojunction devices fabricated with a blend of each donors and PC\(_{70}\)BM in the ratio 1:0.5 (w/w) (except for 55: 1:0.8 (w/w)) led moderate to high PCE as high as 4.05%, 5.08%, 6.10% and 2.46% for 55 – 58 respectively. The enhancement of PCE for 57 compared to 56 is attributed to the higher \( J_{SC} \) value (\( J_{SC} = 13.98 \) mA cm\(^{-2}\) versus \( J_{SC} = 10.74 \) mA cm\(^{-2}\)) due to a stronger red shift of the maximum of absorption. Changing the previous acceptors by the 1,3-indanedione analogous in compound 61 leads to a reduced band gap and
should improve their photovoltaic conversion efficiencies (Scheme 31). Nevertheless, the poor packing in the solid state (as revealed by XRD analysis) and its low LUMO level (-3.86 eV) provide a relatively low PCE for 60 that do not exceed 0.66%. The rather low PCE could be assigned to the lack of solubility for those chromophores, which turns to be prohibitive for fabrication solution processes (i.e. for 61). However, considering the better solubility of 59, the 59-based structure gives rise to an efficient device with a PCE of 4.93% and I/V characteristics close to that of 56 and 57.51

Scheme 31

Further modulations were achieved from these linear systems by inserting into the π-conjugated systems various unsubstituted or substituted rigid blocks (Schemes 32-33) that can help to adjust the photovoltaic characteristics as well as their intrinsic optoelectronic properties. For instance, the dithienosilole derivative 62 demonstrated greater light harvesting efficiency. BHJ devices based on this chromophore showed a PCE of 5.84% along with a noticeably high fill factor of 0.64.52 As an archetype structure benzothioiophene (BTD) 63 was first investigated and used for the preparation of an efficient BHJ cell (PCE = 5.44%) with a high Voc of 0.93 V.53 Encouraged by these results further optimization were achieved by changing the nature of the acceptors (to have greater light absorption) and also by increasing their solubility by the introduction of alkyl side chains (64-65).54 Hence, an optimized BHJ device made from a blend of 65, PC71BM and PDMS as additive in the active layer led to one of the highest PCE value (7.38%) with an average 7.18% obtained from a hundred devices.

Scheme 32

Scheme 33

With the aim to enlarge the scope of this series, chromophore 65 was further optimized into 2D structures by expanding laterally the π-conjugated system with different thiényl substituents (66a-66c, Scheme 33).55 Solution-processed BHJ devices were fabricated from these chromophores and PC71BM as electron acceptor. Similarly to 65, the optimal weight ratio D:A was selected to be 1:0.8, and the device performances were improved by addition of PDMS as an additive (0.2 mg mL-1) (PDMS). Additionally, when other additives such as 1,8-diodooctane and 1-chloronaphthalene were used no significant improvement was attained. For instance, optimized device performance based on 66a showed an exceptional PCE value of 8.12% with a Voc= 0.93V, Jsc= 13.17 mA.cm-2 and a fill factor of 0.66.

Cui et al.55 prepared the analogous 2D molecule 67 of derivative 64 with an A-D-A framework (Scheme 34). Although optical properties were not modified significantly compared to 64, the BHJ OSC device based on 67: PC71BM (1:0.5, w/w) leads to an increase of the PCE 4.0% (which almost twofold magnitude higher than for 64) associated to a notable fill factor of 0.63 without any specific treatments.

Scheme 34

The use of benzothiophene (BTT) as central core instead of BDT was investigated by Patra et al. (Scheme 35).56 The first attempts to make solar cells devices from 68b and PC71BM (in 1:0.75 ratio (w/w)) led to a respectably power conversion efficiency of 3.61% with Voc= 0.88V, Jsc= 7.39 mA.cm-2 and a fill factor of 0.57. Improvements during the fabrication process by adding 0.25 vol% of 1-chloronaphthalene, as an additive, during the formation of the blend film of 68b: PC71BM, increased significantly the PCE to 5.05% with Voc= 0.86V, Jsc= 9.94 mA.cm-2 and a fill factor of 0.59. In such devices, the enhancement was attributed to the suppression of the nanophase molecular aggregation.

Scheme 35

Pursuing their efforts on the optimization of the molecular structures led Shen et al. to develop shorter derivatives of compounds 63-67 that are depicted in scheme 36.57 Thus, four new solution-processable A-D-A chromophores (70-72) were designed and synthetized with a BDT as central core end-capped with indenedione (ID) as acceptor and connected by thiophene or bithiophene as π-conjugated spacers. The photovoltaic conversion efficiencies were determined for each compound in presence of PC71BM (1.5:1 ratio (w/w)). The results indicate that in such structures the π-conjugated spacer is of importance, as well as the side chain on BDT, because the highest efficiencies were obtained for the longest spacer and correlate perfectly with the results for the series 63-67. Indeed, the PCE are respectively 5.67% for 69, 6.75% for 70, 4.15% for 71 and 5.11% for 72.

Scheme 36

Since recently new interesting acceptors have been commonly used for the synthesis of advanced chromophores in optoelectronic applications namely the diketopyrrolopyrrole core (DPP). Thus, Yao et al.58 end-capped the previous DBT core with DPP and fabricate an efficient device with PC71BM as acceptor (Scheme 37). Indeed, when compound 73 was deposited as a blend by spin-casting from an α-dichlorobenzene solution in presence of an optimized percentage of DIO (0.70%), used as additive, a continuous interpenetrating network film was formed associated to a photovoltaic efficiency up to 5.29% and gathering the following I/V characteristics Voc= 0.72V, Jsc= 11.86 mA.cm-2 and a fill factor of 0.62. The lowest value obtained for 73 compared to 69 was incriminated to a reduction of the Voc. Slight enhancements have been achieved, from the work of Zhian et al.,59 by changing the nature of the acceptor, from PC71BM to PC61BM, and applying a thermal annealing at 110°C during 10 min. Consequently a PCE of 5.79% was attained.

Scheme 37

Besides this example, the DPP unit has been used in conjunction with several others π-conjugated spacers in order to investigate the influence of the spacer on the electropical properties of the synthesized chromophores. Thereby, Jo and Choi60 have studied and compared the influence of electro-donating character and planarity of different spacers on the
properties (Scheme 38).

**Scheme 38**

From this study it appears that the insertion of weak \( \pi \)-donating spacers, such as biphenyl (76) and naphthalene (77), contribute to higher \( V_{OC} \) inversely to stronger systems like dithienyl (74), or fused-biophene (75) due to a lower HOMO energy level. In addition, when more rigid and planar \( \pi \)-conjugated systems were used it was observed an increase of the short-circuit current density \( J_{SC} \) (75 and 77) owing to a higher hole-mobility. Therefore the use of naphthalene in this series corresponds to the best compromise owing its enhanced \( V_{OC} \) and \( J_{SC} \) characteristics leading to a device with a power conversion efficiency of 4.40%.

Y. Li et al. have done further investigation by replacing the central core by other rigid and non-rigid nitrogen aromatic \( \pi \)-conjugated structures (Scheme 39). Owing their light harvesting properties and appropriate energy levels the new compounds have been tested in a device having the following structure: ITO/PEDOT: PSS:78: PC_{61}BM/LiF/Al. Nevertheless among the examined structures only 78a gave satisfactory performance with a PCE of 1.50%, along the I/V characteristics \( V_{OC}=0.66\text{V}, J_{SC}=4.12\text{mAcm}^{-2} \) and a FF of 0.44.

**Scheme 39**

Extension of the carbazole \( \pi \)-conjugated spacer of the compound 78a, based on the two regioisomeric indolo[3,2-b]carbazoles cores (Compounds 79-80, scheme 40), has been carried out by Yagai et al. Interestingly, depending on the regiochemistry of the backbone linkages (3,9- or 2,8-) two different solid-state behaviors are found (crystalline vs amorphous). As a result, significant dissimilarities in phase segregation are obtained from 79 and 80 blended films with PC_{61}BM that affect dramatically the photophysical properties. For instance, compound 79 presents a linear structure (crystalline) whereas 80 displays a S-shape structure (amorphous). These differences imply that thermal annealing would decrease or increase respectively the degree of phase separation and would modify their molecular packing. Consequently, although the compounds display the same optical and electrochemical properties their PCE are quite different upon thermal annealing due to a difference on the degree of phase separation. The best results (PCE = 1.80%) were obtained for 79 after thermal annealing at 130°C (\( V_{OC}=0.77\text{V}, J_{SC}=5.91\text{mAcm}^{-2} \) and a FF of 0.40) while 80 presents under the same condition a PCE of 1.40% (\( V_{OC}=0.72\text{V}, J_{SC}=3.92\text{mAcm}^{-2} \) and a FF of 0.50).

**Scheme 40**

Thereafter, a novel series of bis-DPP derivatives bridged with various \( \pi \)-conjugated spacers were also investigated by Nguyen and co-workers in order to elucidate and understand more deeply the molecular features that can be helpful for the construction of novel effective molecular structures (Scheme 41). Additionally, their studies have been correlated with DFT calculations that have been proven to be very effective as an aide prediction tools. A rapid screening of compounds 81-88 has revealed that the structure 86d constitutes one of the most promising structures and was chosen for further optimization. Therefore, the best performances (PCE up to 4.20% with \( V_{OC}=0.85\text{V}, J_{SC}=9.90\text{mAcm}^{-2} \) and FF= 0.49) for 86d have been achieved for a blend of 86d:PC_{70}BM (1:1 w/w) annealed at 80°C in presence of chloronaphthlene (0.50%) as solvent additive.

**Scheme 41**

In a similarly work, R. A. J. Janssen et al. have studied four novel tetraphiophene derivatives flanked with DKKP end-capped either with benzothiophene or benzofuran moieties (Scheme 42). Interestingly, all structures (89a-89d) possess comparable electro-optical properties but display quite different solubility behaviors. During the device fabrication, careful adaptation of the processing conditions (concentration, solvent, temperature, additive) for each compounds in presence of PC_{71}BM as acceptor have given cells performances ranking from 3.6 to 4.6%. This experiments point out that small structural changes will strongly affect the solubility properties and consequently the photovoltaic performances.

**Scheme 42**

An analogous derivative 90 (Scheme 43) has been recently described by Zhu et al. showing good optical properties and optimum energy levels for OPV applications. Solution processed BHJ solar cells have been fabricated from 90 and PC_{70}BM (1:1.5, w/w) leading to devices with PCE of 2.82% after thermal annealing. The lowered values obtained from this device compared to the previous, based on tetraphiophene core, might be relied on a decrease of the molecular packing. More interestingly, compound 90 has been successfully used as an acceptor in a BHJ device in a blend film with P3HT (1:1.2, w/w respectively) yielding a PCE of 0.83% with a \( V_{OC}=1.17\text{V} \) after thermal annealing. This evidences an ambipolar character for 90 with the possibilities to use such compounds either as donor or acceptor in the construction of solar cells.

**Scheme 43**

As already mentioned the possibilities to boost the photo-absorption properties as well as the electron and hole-mobilities can be achieved both by chemical design and engineering, but drive to more and more complicated structures. In addition, the aim to use solution-processed technologies requires the use of large solubilizing units that can affect at the meantime the solid-state arrangement. To circumvent such difficulties Bäuerle et al. studied, using vacuum deposition process, some oligothiophenic structures substituted with shorter alkyl-chains, based on the \( \alpha \)-quinquethiophene core, and end-capped with dicyanovinyl groups (Scheme 44) in OPV devices.

**Scheme 44**

Thus, optimized planar heterojunctions made from compounds 91 and C_{60} demonstrated PCE of 3.4% with \( V_{OC}=0.98\text{V} \) and \( J_{SC}=10.60\text{mAcm}^{-2} \). Based on those promising results a series of unsubstituted oligothiophene 92 (n = 1-6) end-capped with DCV was further investigated (Scheme 45).
For instance, 92a showed a well-structured absorption band centered at 406 nm. Increasing the number of thiophene units leads to a bathochromic shift of the maximum of absorption from 406 nm (92a) to 532 nm for 92f associated to a hyperchromic effect (92a: ε = 39 000 L mol⁻¹ cm⁻¹ to 92e: ε = 73 300 L mol⁻¹ cm⁻¹). Devices made by vacuum deposition of 15 nm of a C₆₀ layer and 6 nm of 92d-92f, showed a high values of PCE = 1.20%, 2.60% and 2.80% respectively. Beside, BHJ cells based on 92e and C₆₀ were also prepared leading to PCE as high as 5.2% after device fabrication optimizations. Solid-state packing was also studied on the methyl substituted electron relay between the acceptors. Replacement of thiophene units by analogous selenophene or selenium containing oligomers by a more number of selenophene decreases also the photovoltaic effect (Scheme 46). Using vacuum deposition technique PHJ solar cells were also investigated (Scheme 47). These modified oligomers (93-95) were incorporated by vacuum deposition process in BHJ solar cells exhibiting PCEs ranking from 4.80 to 6.10%. Subsequent optimization of the deposition process and changes of the fabrication conditions have led to an enhanced PCE up to 6.90%. This result suggested the existence of strong specific intermolecular interactions within the organic layer associated to effective π-π overlap and multidirectional electronic coupling evidenced by single-crystal X-ray structure analysis of 94.

**Scheme 45**

**Scheme 46**

Replacement of thiophene units by analogous selenophene or ethylenedioxythiophene (EDOT) core were also investigated (Scheme 47).

**Scheme 47**

Nevertheless, despite a broader and more intense absorption properties selenium-containing oligomers (96-98) have shown lower performances compared to 92e with PCEs ranging from 2.50 to 3.10%. The lowest obtained values might be attributed to lowered solid-state packing due to the longer alky chains used along the π-conjugated backbone. In addition increasing the number of selenophene decreases also the photovoltaic characteristics. Later on Palomares et al. have changed the thiophene or selenium containing oligomers by a more electron rich unit such as EDOT (99). A BHJ device structure based on 99:PC₇₁BM gave rise to an enhancement of the PCE up to 3.75% associated to good annealing properties and high open-circuit voltage and fill-factor (V_OC= 1.01V, J_SC= 6.00 mA.cm⁻² and FF= 0.63). Benzothiadiazole derivatives have been also examined as terminal acceptors in oligothiophene. Thus compounds 100 and 101 have been synthetized (Scheme 48) using a terthiophene as electron relay between the acceptors.

**Scheme 48**

Combination of benzo[c][1,2,5]thiadiazole with 2,7-carbazole end-capped with oligothiophene have been explored by Li et al. in order to readjust HOMO levels compared to molecules 100 or 101 and optimized the band gap (Scheme 49). Thus, solution processed organic solar cells based on a blend 102a/PC₇₁BM exhibit a PCE of 2.26% associated with a high open-circuit voltage close to that of 101 (V_OC= 0.98-1.05V). Expanding the terminal π-conjugated system by an oligothiophene moiety (102b and 102c) leads to a slight decrease of the V_OC (V_OC= 0.92 and 0.86V respectively) on devices having the same configuration accompanied with higher PCE values up to 3.44% and 3.90% respectively without any post-treatment due to an enhancement of the J_SC and FF.

**Scheme 49**

Different type of s-tetrazine derivatives have been investigated and tested as end-groups in a oligothiophene series synthesized by Tu et al. Thus, active solar cells obtained from the series of the compound 103 (Scheme 50) show, by AFM and TEM experiments, that on blended films of 103: PC₆₀BM (1:1 w/w ratio) well-ordered packing structure of 103 in the crystalline state leads to higher photovoltaic performances. In addition, replacement of PC₆₀BM by PC₇₁BM in presence of 1% of DIO as additive affords a device with a power conversion efficiency reaching 3.24% (J_SC= 7.87 mA.cm⁻²).

**Scheme 50**

The strategy has been applied toward the synthesis of extended A-D-A-D-A chromophores incorporating two different acceptors at central and terminal positions along the π-conjugated backbone namely the benzothiadiazole and trifluoroacetyl. Using vacuum deposition technique PHJ solar cells were prepared from the compounds 104-106. The presence of these acceptors lower the HOMO level, providing an unusually high V_OC (V_OC= 1.17V (105) and 1.10V (106)) associated to moderate PCE of 1.56% and 1.45% respectively.

**Scheme 51**

Similar studies have been conducted by Y. Chen et al. on the solution processable chromophore 107 incorporated in BHJ solar cells (Scheme 52) through a spin coating process. An optimized donor/acceptor weight ratio was found to be 1:0.8 for 107:PC₆₀BM affording a PCE of 2.43% without any post-treatment (V_OC= 0.76V, J_SC= 5.20 mA.cm⁻² and FF= 0.61%) and using LiF/Al as cathode. Thermal annealing at 150 °C for 10 min improved the PCE up to 3.07% (V_OC= 0.78V, J_SC= 7.10 mA.cm⁻² and FF= 0.55), suggesting that the enhancement can be attributed to a better molecular packing at the solid state associated with an improvement of the absorption intensity and hence the shunt current circuit density from 5.20 to 7.10 mA.cm⁻².

**Scheme 52**

Thienothiadiazole and thienopyrazine have been also examined as central acceptors in oligothiophene series end-capped with DCV or trifluoroacetyl groups (Scheme 53). Insertion of such central acceptors leads to a red shift of the maximum of absorption compared to their analogous parent compounds with...
benzothiadiazole (707 nm, 675 nm, 620 nm for 108, 109 and 110 respectively). Nevertheless, the different attempts to optimize the solar cells during the process of fabrication have not permitted to enhance efficiently the photovoltaic characteristics. Indeed, only moderate PCE values were attained in the range of 0.8% to 1.3% ascribed to the lack of transport of the photo-generated electrons and holes ($J_{sc}$ are not exceeding 3.20 mA.cm$^{-2}$), the low molar absorptivity and the low-lying LUMO energy levels of the donor.\textsuperscript{[75]}

Scheme 53

With the aim to enhance the electron withdrawing efficiency properties of DCV and to reduce the optical band gap, a nitrobenzene ring can be used in place of one of the cyano groups. Mikroyannidis et al. have successfully applied this strategy during the synthesis of the phenylenevinylene-based chromophore 111 (Scheme 54).\textsuperscript{[76]} Replacement of one of the cyano groups leads to a reduction of the optical band gap, with a maximum of absorption around 640 nm, compared to the unsubstituted DCV compound. BHJ solar cells device were fabricated using 111: PCBM as active material sandwiched between ITO/Al electrodes. The low PCE, 1.36%, is mainly attributed to the low $J_{sc}$ ($V_{oc}$= 0.73 V, $J_{sc}$= 0.056 mA.cm$^{-2}$ and FF= 0.31) due to the small exciton diffusion lengths as well as low mobility of hole. Thermal annealing of the blend affords an improvement of the PCE up to 2.33% induced by an increase of the packing density facilitating both the exciton dissociation and charge transportation.

Scheme 54

In addition, changing the PC$_{60}$BM acceptor by perylene diimide PDI in the previous device leads to a remarkable PCE of 1.87%. Further improvement, notably by insertion of ZnO in between the active layer and the Al-cathode followed by a thermal annealing (5 min., 100°C) gives rise to an exceptional PCE of 3.17%.\textsuperscript{[77]}

With the aim to enhance the light-harvesting abilities and optimize the optical bandgap compared to A-D-A counterparts, Wong et al. have envisioned the combination of two strong electron-withdrawing groups leading to the A-A-D-A-A structures.\textsuperscript{[78]} Hence, in the following examples 112-113 (Scheme 55) the dithienosilole (DTS) core has been selected as $p$-type building block in respect with its better packing abilities and higher hole-mobilities compared to their carbon-based homologues.

Scheme 55

Elaboration of planar mixed heterojunction devices based on 112 and fullerene acceptors (C$_{60}$ or C$_{70}$) leads to decent power conversion efficiencies from 2.3% to 3.7% respectively ($V_{oc}$= 1.05 V, $J_{sc}$= 5.80 mA.cm$^{-2}$ and FF= 0.38, $V_{oc}$= 1.01 V, $J_{sc}$= 9.79 mA.cm$^{-2}$ and FF= 0.38) after tuning the donor: acceptor ratio (from 1:1 to 1:1.5) without any post-treatments such as thermal of solvent annealing processes.

Further investigations have been done with the DTS core incorporating solubilizing chains (Scheme 56). The roles of such chains are to improve the solubility of chromophores in organic solvents for solution casting processes as well as their packing structure and their solid-state miscibility with fullerenes. A careful attention on the molecular design, chain position, and electron density has to be paid to ensure the best properties. In this context, Wang et al. described in a recent work the impact of such molecular modification on the photoconversion properties.\textsuperscript{[79]}

Scheme 56

The best results were obtained for 116 and PC$_{60}$BM (1:0.8 ratio) with a PCE up to 3.81% ($V_{oc}$= 0.92 V, $J_{sc}$= 8.73 mA.cm$^{-2}$ and FF= 0.49). AFM microphotographs of spin-coated films from chloroform solutions show better interpenetrating network for 116 compared to 114-115 associated to higher roughness and better optical absorption properties. These results highlight again the importance to control the size, nature, length and position of the alkyl chains that strongly impact the optical absorption, energy levels, packing and film morphology.

To gain further insights, Kim et al. have introduced on the terminal alkyl substituents an amide function (compounds 118) that would favor intermolecular interactions via a hydrogen bonding network\textsuperscript{[80]} compared to 117 structures (Scheme 57). This hydrogen bonding will affect dramatically the interfacial interactions, optical properties, the film morphology and therefore the photovoltaic properties. Thus, it has been demonstrated in this work that the introduction of intermolecular interaction induces well-ordered molecular packing that is not suited for the optimization of the photovoltaic performance. Indeed, the best result was obtained for 117b with a PCE exceeding 4.30% ($V_{oc}$= 0.82 V, $J_{sc}$= 9.79 mA.cm$^{-2}$ and FF= 0.54) while its counterpart 118b leads only a PCE of 3.22% under the same conditions ($V_{oc}$= 0.87 V, $J_{sc}$= 7.94 mA.cm$^{-2}$ and FF= 0.47).

Scheme 57

An analogous strategy was adopted by Lam and co-workers but using a terminal acid group (CO$_{2}$H) (Scheme 58).\textsuperscript{[81]} Interestingly, in this case an improvement of the photovoltaic efficiency was observed for the derivative 119b compared to its ester analogue (119a). The enhanced PCE value was mainly attributed to a better hydrogen-bonding network that induces strong $\pi-\pi$ interactions and highly ordered films with a vertical alignment. This behavior leads to higher hole-mobilities in favor of 119b as measured by the SCLC method ($\mu_h$ = 2.83 $10^{-5}$ cm$^2$ V$^{-1}$ s$^{-1}$ (119b) $\mu_h$ = 6.91 $10^{-6}$ cm$^2$ V$^{-1}$ s$^{-1}$ (119a)) that directly impact on the photovoltaic conversion efficiency (0.91% ($V_{oc}$= 0.71 V, $J_{sc}$= 2.86 mA.cm$^{-2}$, FF= 0.45) and 0.66% ($V_{oc}$= 0.53 V, $J_{sc}$= 0.45 mA.cm$^{-2}$, FF= 0.26) respectively). In addition, it is noted that the formation of nanofibers as-cast films prevents the use of post-treatments or solvent additives.

Scheme 58

D – A – D

Besides the A-D-A structures, several groups have developed in parallel D-A-D architectures with the same aim. For instance, one can cite the work of Li\textsuperscript{[82]} who has widely used the
benzo[1,2,5]thiadiazole core as central acceptor part flanked with TPA donors and connected together by thiophenic linkers (Scheme 59).

Scheme 59

Preliminary studies based on the derivative 120 have been performed with different devices architectures made from pristine 120, a PHJ structure (120 and C60) and a BHJ from a blend of 120 and PC60BM. In a typical device, the active layer was sandwiched between a ITO/PEDOT: PSS as anode and LiF/Al as cathode. From the obtained results it has been demonstrated herein that the configuration BHJ shows the highest photovoltaic performances with a PCE 0.19%. Further optimization of cathode material by using Ba/Al instead of LiF/Al gave rise to an enhancement of the PCE up to 0.26%.52

In order to increase the photovoltaic performances of such derivatives, multi-arms architectures were built up as shown in scheme 60 and compared with their linear D-A-D counterparts.83

Scheme 60

All structures are well soluble in dichlorobenzene, which makes them suitable to elaborate organic solar cells by spin-coating processes. Thus, in devices with structure of ITO/PEDOT: PSS/active layer/LiF/Al and optimized conditions (thickness (70-90 nm), donor: acceptor (PC71BM) ratio 1:3 (w/w)) the highest PCE was obtained for the four-arms derivatives 122c (1.80%, Voc = 0.92V, Jsc = 4.90 mA.cm⁻² and FF= 0.41). Nevertheless for all compounds tested the PCE values are comprised between 1.10 to 1.80% demonstrating that other parameters have to be controlled and taken into account to optimize the devices.

Further investigations were done on compounds 123 and 124 (Scheme 61). To ensure sufficient solubility alkyl chains have been introduced along the π-conjugated backbone (123) or at the terminal position (124). The derivative 123 is crystalline while its counterpart 124 is amorphous, this difference will certainly affect the performances of the devices. As expected, 124 exhibits a bathochromic shift (λmax=456 nm and λmax=353 respectively) and a higher HOMO level compared to 125 due to its extended π-conjugated system. Organic solar cells were fabricated and the device based on 123:PC60BM (1:3 ratio (w/w) exhibits higher Voc and PCE than that for 124:PC60BM (0.56% (Voc= 0.80, Jsc= 1.80 mA.cm⁻², FF= 0.39) and 0.42% (Voc= 0.62V, Jsc= 1.89 mA.cm⁻², FF= 0.36) respectively) attributed to the lower HOMO energy level of 123. Changing the nature of the acceptor (123: PC71BM) yielded to a higher PCE up to 1.23% (Voc= 0.86, Jsc= 3.50 mA.cm⁻², FF= 0.41), which is not improved by thermal annealing.84

Scheme 61

In the following example Zhen et al. have investigated the star-shaped bis-(tri-substituted)TPA 125 end-capped at two of its extremities by an hexyl-thiophene and connected through a benzo[1,2,5]thiadiazole core (Scheme 62). Organic solar cells based on 125: PC71BM (donor: acceptor ratio 1:3) yielded, without any solvent additives or thermal post-treatments, a PCE of 3.18% (Voc= 0.96V, Jsc= 6.83 mA.cm⁻², FF= 0.48). This example demonstrates that the connection pathway of the different building blocks along the π-conjugated system is of crucial importance to control the photoconversion efficiencies.

Scheme 62

The nature of the π-conjugated linker plays also an essential role in the achieved properties. In this context, Li and coll. have structurally modified the archetype chromophore 120 in order to improve its efficiency in BHJ solar cells (Scheme 63).66 Thus the insertion of hexyl side chains (126) helps in getting better blend morphology. As a consequence a PCE up to 1.44% was attained. (126: PC71BM (1:3 ratio) Voc= 0.79V, Jsc= 4.84 mA.cm⁻², FF= 0.37). Extending the π-conjugated system with thieno[3,2-b]thiophene (127) instead of thiophene allows significantly to broaden the absorption spectra. However, even if an increase of the shunt current circuit density is noticed (from Jsc= 4.84 mA.cm⁻² to Jsc= 5.71 mA.cm⁻²) the PCE for 126 still the same (1.44%) as the Voc has decreased (from Voc= 0.79V to Voc= 0.74V) due to a higher HOMO energy level. The introduction of bulky alkyl chains at the periphery (128) reduced significantly the PCE from 1.44% to 0.75% related to poorer miscibility with PC71BM as evidenced by the AFM microphotographs and the lower hole-mobilities (from µh = 1.48 10⁻⁴ cm² V⁻¹ s⁻¹ (127) to µh = 2.78 10⁻⁵ cm² V⁻¹ s⁻¹ (128)) determined by the SCLC (space-charge-limited-current) method.

Scheme 63

The results demonstrate that broadening the absorption spectra with remaining the lower HOMO energy level have to be considered in the synthesis and design of high performance organic chromophores.

Pursuing their efforts in the understanding and in the design/synthesis of efficient chromophores for photovoltaic applications, Li et al. have replaced the benzo[1,2,5]thiazole core by a (dicyanomethylene)-pyrrole unit in chromophores 129 to 131 (Scheme 64). Organic solar cells based on 128: PC71BM (1:3 ratio w/w) exhibits a PCE of 1.40% (Voc= 0.71V, Jsc= 5.07 mA.cm⁻², FF= 0.38) which corresponds to almost twice the value obtained from that based on 129: PC60BM (PCE= 0.79%) due to the stronger absorption properties of PC71BM in the wavelength range 350-550nm. Modifying the linker from benzene (129) to thiophene (130) alters the photophysical properties. For instance a red shift of the maximum of absorption was observed as well as a lower bandgap. Therefore, under the same conditions an improvement up to 2.06% of the photovoltaic efficiency was attained with 129: PC71BM (Voc= 0.79V, Jsc= 5.94 mA.cm⁻², FF= 0.44). Introduction of solubilizing chains along the π-conjugated system (131) leads to a better miscibility with PC71BM, without altering the optical properties, associated to an enhancement of the PCE to 2.10% (Voc= 0.78V, Jsc= 6.74 mA.cm⁻², FF= 0.39).87

Scheme 64

Further work has been done Zhan and coll. with other central acceptors such as thieno[3,4-c]pyrrole-4,6-dione (132a-c)88 and bisthiazole (133a-c)89 that are widely used to synthesize low-
bandgap polymers due to their strong electron withdrawing abilities (Scheme 65).

Scheme 65

5 Photovoltaic cells were fabricated with a structure of ITO/PEDOT: PSS/132-a-c: PC71BM/Al. Interestingly the use of thieno[3,4-c]pyrrole-4,6-dione in 132 as acceptor appears to be a promising candidate as its photovoltaic efficiency was higher than its counterparts 121-a. Indeed, under identical conditions 121-a and 123 have a PCE of 1.30% (VSC = 0.71V, JSC = 4.80 mA.cm-2, FF = 0.38) and 1.70% (VSC = 0.87V, JSC = 4.80 mA.cm-2, FF = 0.40) respectively while 132 exhibits values of PCE ranking from 1.91% to 2.41% (VSC = 0.85V, JSC = 5.52 mA.cm-2, FF = 0.41 and VSC = 0.86V, JSC = 5.11 mA.cm-2, FF = 0.43). By changing the donor:acceptor ratio from 1:3 (w/w) to 1:4 (w/w) enhanced the PCE up to 2.87% (VSC = 0.94V, JSC = 6.94 mA.cm-2, FF = 0.44) and 2.90% (VSC = 0.88V, JSC = 7.73 mA.cm-2, FF = 0.43) respectively. Finally, thermal annealing of the blend 132:PC71BM (1:4 w/w) led to a PCE of 3.31% (VSC = 0.91V, JSC = 7.70 mA.cm-2, FF = 0.47). Substitution of thieno[3,4-c]pyrrole-4,6-dione by bithiophene leads to the chromophores 133-a-c. Increasing the π-conjugated system from 133a to 133c provides better intermolecular interactions in the solid-sate and generates strong tendency to absorb the low energy photons as evidenced by the UV-Vis spectroscopy. The chromophores show typical p-type semi-conductor behavior in air with hole-mobilities as high as µh = 3.60 x 10⁻⁸ cm² V⁻¹ s⁻¹ (133c). BHJ solar cells based on 133a-c: PC71BM (1:4; w: w) blend films showed increasing PCE (up to 2.61%) due to broader and stronger absorption as well as higher hole-mobility as expanding the π-conjugated system (spin-cast solutions: 133a: VSC = 0.71V, JSC = 3.42 mA.cm-2, FF = 0.30, 133b: VSC = 0.81V, JSC = 4.69 mA.cm-2, FF = 0.33 and 133c: VSC = 0.84V, JSC = 7.72 mA.cm-2, FF = 0.40).

A narrow band gap chromophore 134 based on triphenylamine and isoindigo was further investigated by Yang and coll. (Scheme 66) due to its efficiency previously demonstrated in DSSCs devices. AFM microphotographs evidenced a good miscibility between 134 and PC61BM with interpenetrating structures, which would facilitate charge separation in the solid state, associated to a broad absorption spectrum from 300 to 750 nm. BHJ OSCs devices with a layered structure of ITO/PEDOT: PSS/134: PC61BM (1:1 ratio w/w)(80nm)/Ca(10nm)/Al(100nm) were fabricated and led to a PCE of 0.61% (VOC = 1.00V, JSC = 2.35 mA.cm², FF = 0.26). VOC is among the highest values in all-small molecules or polymers highlighting the importance of such acceptor in future development of SMOSCs due to the high VOC attained.

Scheme 66

A α-heptathiophene series of D-A-D-type oligomers 135, 136, and 137 based on TPD (thieno[3,4-c]pyrrole-4,6-dione), TPN (thienopyrrole naphthalene) and TIN (thienoimidazoyle naphthalene) as central acceptor core respectively have been developed by H.-Y. Wang and coll. (Scheme 67). Modification of the diimide unit of 135 by perylene or imidazole analogues in 136 and 137 has the benefits to enhance the coplanarity and to tune the energy levels due to a better intermolecular stacking in the solid-state. The photovoltaic performances were determined by fabricating BHJ devices (ITO/PEDOT: PSS/135-137: PC71BM/LiF/Al) with a donor: acceptor ratio (2:1, w/w). The device 135: PC71BM blends showed the best PCE of 1.87% (VOC = 0.87V, JSC = 4.93 mA.cm², FF = 0.44) while 136: PC71BM and 137: PC71BM blends in the same conditions led only to PCE of 0.77% and 0.11% respectively (VOC = 0.76V, JSC = 2.22 mA.cm², FF = 0.45, VOC = 0.75V, JSC = 0.58 mA.cm², FF = 0.25). However, for 136 and 137 efficiencies can be improved in blends device structures with a weight ratio of 1:1 (w/w) respectively to 1.24% and 0.34% (VOC = 0.77V, JSC = 2.75 mA.cm², FF = 0.59, VOC = 0.69V, JSC = 1.91 mA.cm², FF = 0.26). The lower PCE are mainly attributed to the different morphology of the blend films and the presence of high crystalline domains.

Scheme 67

By comparison of the work done by Zhan (Scheme 65), Suh et al. have developed a new series 138a-b based on the bithiophene imide (BTI) as an electron deficient unit. (Scheme 68) The BTI is unit of choice due to its strong electron-withdrawing character, planar architecture, good solubility, and close π-π stacking in the solid state. In addition, BTI presents the advantage to decrease the steric hindrance between the alkyl chains of the electron rich unit and the imide group and changes the shape of the oligomers (as banana-like structure) compared to the 132 analogues. J-V characteristics obtained from devices with the geometry ITO/PEDOT: PSS (40 nm)/138: PC71BM (80 nm)/Al (100 nm) led to the best performances in favour of 138a with a PCE of 1.36% (1:3 ratio, VOC = 0.91V, JSC = 3.93 mA.cm², FF = 0.38) after a thermal annealing a 130°C. Extension of the π-conjugated system for 138b didn’t induced any improvement of the photovoltaic characteristics, as the best PCE attained is only of 1.08% (1:4 ratio, thermal annealing at 120°C, VOC = 0.80V, JSC = 3.79 mA.cm², FF = 0.36) even after addition of solvent additives such as 1,8-octanedithiol (ODT) or 1,8-diiodooctane (DIO).

Scheme 68

Cao et al. have investigated the influence of the extension of the π-conjugated system of the chromophores 121, using fused fluorenyl TPA-based units, on the photovoltaic properties (Scheme 69). BHJ solar cells were fabricated with the device configuration ITO/PEDOT: PSS/139: PC61BM or 140: PC61BM/Ba/Al. Several weight ratios were studied to improve the efficiencies (1: 1, 1: 2 and 1: 4). As demonstrated improvement of the device characteristics were obtained by increasing the PC61BM content in the blend films. In both examples the best PCE achieved were 0.75% and 0.22% (1:4 ratio, VOC = 0.75V, JSC = 1.95 mA.cm², FF = 0.34, 1:4 ratio, VOC = 0.70V, JSC = 1.59 mA.cm², FF = 0.22) respectively. Despite an extended photosensitivity response to near-infrared region for those chromophores and good film forming properties, the PCE was still low. The low PCE was incriminated to a less efficient charge separation due to LUMO levels close to the LUMO level of PC61BM.
Similarly to 139, Hyun et al. have evaluated the photovoltaic properties of thiadiazoloquinoloxaline-based units99 by varying the nature of the π-conjugated backbone as well as the substituents on the quinoxaline moiety (for instance isobutyl and thiophene) (Scheme 70). The nature of the substituents on the quinoxaline part has a dramatic influence on the final optical properties; from isobutyl to thiophene the absorption onset was shifted from 900 to 1100 nm in the solid state. Device structures with the configuration ITO/PEDOT: PSS/PC71BM (1:3 w/w)/TiOx/Al have been fabricated and the performances evaluated under the AM1.5G conditions. 141:PC71BM blend films showed better PCE (0.24%, $V_{OC}=0.42$V, $J_{SC}=2.42$ mA.cm$^{-2}$, FF=0.24) than 142: PC71BM (0.14%, $V_{OC}=0.38$V, $J_{SC}=1.40$ mA.cm$^{-2}$, FF=0.36) even if 142 possesses a broader absorption up to 1100 nm. The lowest PCE for 142 was attributed to a reduced charge separation and a relatively rougher surface. Compared to its counterpart 139 the lower PCE attained was also due to the weakness of the donor used.

**Scheme 70**

Cyan substituents have been introduced along the π-conjugated systems in order to take advantage of additional intramoelcular charge transfer to reduce the bandgap and to extend the absorption spectrum of the chromophores (Scheme 71). Hence, Li et al. have conducted preliminary investigations on compounds 143 bearing two different acceptors namely BX (benzoxadiazole 143a) and BT (benzothiadiazole 143b).99 BX was used to lower the HOMO level of molecular donors, compared to BT, and thus to increase the open-circuit voltage. Devices fabricated with the configuration ITO/PEDOT: PSS/143: PC60BM/Al showed PCE of 0.33% for 143a ($V_{OC}=0.58$V, $J_{SC}=2.39$ mA.cm$^{-2}$, FF=0.24) while 143b exhibited a PCE of 0.50% ($V_{OC}=0.60$V, $J_{SC}=2.86$ mA.cm$^{-2}$, FF=0.29). Interestingly, the introduction of alkoxy side chains on the BT core has a dramatic effect on the photovoltaic properties (144-145). The photovoltaic performances were evaluated on devices with the structure ITO/PEDOT: PSS/144-145: PC60BM (1:2 ratio, w/w)/Al. The BHJ fabricated with 145: PC60BM as a blend film provided an impressive PCE of 3.85% ($V_{OC}=1.04$V, $J_{SC}=11.20$ mA.cm$^{-2}$, FF=0.33) while 144: PC60BM did not exceed 2% ($V_{OC}=0.94$V, $J_{SC}=6.33$ mA.cm$^{-2}$, FF=0.33).97 The highest PCE was consistent with a deeper low-lying HOMO level induced by the cyan groups and better film morphology compared to 143b.

To get further insights on molecular design of efficient chromophores Ko et al. have studied the unsymmetrical D-A-D 146 chromophores based on a benzo[1,2,5]thiadiazole central core (Scheme 72).98

**Scheme 72**

Their photovoltaic performances were assessed with the devices configuration ITO/PEDOT: PSS (35nm)/146: PC70BM (55-65nm) (1:5 ratio, w/w)/Al (100nm). Importantly, it was observed that fluorination increased the short-circuit current $J_{SC}$ and the $V_{OC}$ induced by improved hole-mobility and reduced HOMO levels, respectively. The best performance was obtained for 146c affording a PCE up to 3.24% ($V_{OC}=0.84$V, $J_{SC}=9.45$ mA.cm$^{-2}$, FF=0.41) while 146b and 146a afforded 3.05% ($V_{OC}=0.81$V, $J_{SC}=9.37$ mA.cm$^{-2}$, FF=0.40) and 2.70% ($V_{OC}=0.80$V, $J_{SC}=9.29$ mA.cm$^{-2}$, FF=0.38) respectively. In addition, insertion of a thin layer of TiOx as an optical spacer enhanced the photovoltaic properties by reducing the interfacial resistance between the active layer and the Al metal electrode. Performances were enhanced in all cases reaching a PCE of 4.24% for 146c ($V_{OC}=0.88$V, $J_{SC}=10.76$ mA.cm$^{-2}$, FF=0.44), 3.73% for 146b ($V_{OC}=0.85$V, $J_{SC}=10.55$ mA.cm$^{-2}$, FF=0.44) and 3.54% for 146a ($V_{OC}=0.82$V, $J_{SC}=10.52$ mA.cm$^{-2}$, FF=0.42).

Fused bisthiazole, namely thiazolothiazole, has served to build up D-A-D architectures because it possesses a rigid coplanar structure and the presence of the imine bonds confers to it an electron-accepting character. Thus, linear chromophores 147 flanked with oligothiophenic units have been investigated by Shin et al.99 The terminal hexyl side chains ensure their solubility in common organic solvents and induce liquid crystalline properties that can benefit for the supramolecular arrangement in the solid state (Scheme 73). Solution processed BHJ have been fabricated with the structure ITO/PEDOT: PSS (50nm)/147: PC60BM (100nm, 1:1 ratio, w/w)/Al (80-100nm) and evaluated. Reasonably high PCE of 1.57% were achieved ($V_{OC}=0.65$V, $J_{SC}=7.85$ mA.cm$^{-2}$, FF=0.31). Furthermore, increasing the donor: acceptor ratio to 1:2 has not led to an improvement of the efficiency and the device has exhibited a PCE of 1.20% ($V_{OC}=0.42$V, $J_{SC}=7.59$ mA.cm$^{-2}$, FF=0.37).

**Scheme 73**

To improve the efficiencies of thiazolothiazole-based chromophores, Zhan et al. have used of triphenylamine as end-capped unit associated to different length of the π-conjugated bridge (Scheme 74).100 As expected, linear extension of the π-conjugated system, from 148 to 150, induced a bathochromic shift of the maximum of absorption and an enhancement of the molar absorptivity.

**Scheme 74**

Organic solar cells were fabricated with the configuration ITO/PEDOT: PSS (35nm)/148-150: PC70BM (70-80 nm, 1:2, 1:3 or 1:4 ratio, w/w)/Ca (15nm)/Al (60nm) and tested. The best performances were obtained for a donor: acceptor ratio of 1:4 and thermal annealing at 110°C (148, 149) or 120°C (150) affording a PCE of 2.17% ($V_{OC}=0.86$V, $J_{SC}=6.08$ mA.cm$^{-2}$, FF=0.415) for 148, 3.61% for 149 ($V_{OC}=0.90$V, $J_{SC}=9.28$ mA.cm$^{-2}$,FF=0.428) and 3.89% for 149 ($V_{OC}=0.85$V, $J_{SC}=9.74$ mA.cm$^{-2}$, FF=0.47) for 150. Particularly, the thermal annealing has affected the electron mobility and hole-transport contributing into their balance (for instance for 150 $\mu_{h}=1.70 \times 10^{-6}$ cm$^{2}$V$^{-1}$s$^{-1}$ and $\mu_{e}=1.13 \times 10^{-4}$ cm$^{2}$V$^{-1}$s$^{-1}$ ($\mu_{h}/\mu_{e}=66.47$) and after the thermal annealing $\mu_{h}=1.01 \times 10^{-3}$ cm$^{2}$V$^{-1}$s$^{-1}$ and $\mu_{e}=1.89 \times 10^{-3}$ cm$^{2}$V$^{-1}$s$^{-1}$ ($\mu_{h}/\mu_{e}=1.89$) and increased the miscibility of PC70BM facilitating the phase separation.

Analogously to 147 Nguyen et al. have designed and synthesized the chromophore 151 having a diketopyrrolopyrrole (DPP) as central electron-withdrawing core (Scheme 75).101
Replacement of the thiazolothiazole core by DPP has led to a strong red shift of 169 nm in chloroform solution (447 nm to 616 nm). In thin film, 151 exhibited two absorption bands centered at 660 nm and 742 nm shifting the absorption band edge from 1.72eV to 1.51eV. This significant change was ascribed to an improved molecular ordering as evidenced by AFM microphotographs and hole-mobility measurement (μ_h = 3.00 10^{-6} cm^2 V^{-1} s^{-1}). BHJ solar cells were fabricated with the structure ITO/PEDOT: PSS (40nm)/PC_71BM (100 nm, blend ratios 30/70, 50/50, 70/30)/Al (100nm). Photovoltaic performances were evaluated for each donor: acceptor ratios. The best PCE of 2.33% (V_{OC}= 0.67V, J_{SC}= 8.42 mA.cm^{-2}, FF= 0.45) was obtained for a 70:30 ratio without any thermal post-treatment to prevent the BOC group elimination.

Adashi et al. also have investigated mesomorphic DPP-based chromophores, in order to achieve optimized photovoltaic efficiencies taking into account their intrinsic self-organizing properties (Scheme 76). Effect of the nature of the alkyl chains, with the appropriate ratio, on the photovoltaic performances was studied. As expected, compounds 152 exhibit liquid-crystalline properties and form well-developed nanostructured bulk heterojunction (BHJ) architectures with a fullerene derivative (PC_71BM). The organic solar cells have been fabricated with the standard device structure ITO/PEDOT: PSS (40nm)/PC_71BM blend (90-120nm, 1:1, 1:5, 1:21 ratios, w/w)/LiF (1nm)/Al (100nm) and evaluated under AM 1.5G illumination. The best performances were achieved with a 1:1 ratio after a thermal annealing at 140°C for 152a. As evidenced on UV-Vis spectra, upon thermal post-treatment major structural changes were obtained in the solid state for the blend 152a:PC_71BM highlighting the formation of H-aggregates at the expense of J-aggregates that are predominant for 152b even after heating. This thermo-responsive organization was fully in agreement with the LC state of 152a occurring from 117°C upon heating. The self-assembling into ordered domains strongly affects the stacking in the solid state, the charge carriers mobilities and the solar cells performances. As anticipated 152a exhibited the highest PCE of 4.30% (V_{OC}= 0.93V, J_{SC}= 8.37 mA.cm^{-2}, FF= 0.55) after thermal annealing (at 140°C) without any additives while 152b gave only a PCE of 1.20% (V_{OC}= 0.93V, J_{SC}= 3.90 mA.cm^{-2}, FF= 0.36) (with a thermal post treatment at 120°C).

To favor the π-π arrangement and the intermolecular contacts in the solid state DPP 153 end-capped with benzofuran units (Scheme 77) has been synthetized and tested by Nguyen et al. Thin film properties have been investigated from pure 153 to blends films of 153: PC_71BM at various ratios (30:70, 50:50, 60:40, 30:70) and temperatures (from room temperature to 150°C). From these experiments, it has been shown that changing the annealing temperature one can control and adjust the surface domains sizes for a given blend. These changes in morphologies are fully in agreement with the absorption and XRD experiments and correlated with the charge carrier mobility measured. Hence, from the understanding thin-films properties, the best conditions are obtained for a given donor: acceptor ratio of 60:40 annealed at 110°C. Using these criterions a PCE as high as 4.40% was achieved (V_{OC}= 0.92V, J_{SC}= 10.00 mA.cm^{-2}, FF= 0.48) in BHJ solar cells.

Pursuing their efforts in structure-properties relationships in the DPP chromophores family, T.-Q. Nguyen et al. have studied the effects of stereoisomerism on 153 on the crystallization properties. Typically, 2-ethylhexyl chains are widely used to improve the solution processability of the synthesized chromophore but possess the disadvantage to present an asymmetric carbon (R or S configuration) affording a mixture of stereoisomers that induce different solid-state properties. Thus, to elucidate the influence of the nature of the stereocenters on the optoelectrical properties the three stereoisomers, namely 153-RR, 153-SS and 153-RS (mesomer), were synthesized and purified by a preparative chiral HPLC. As expected, from crystallographic data 153-RR and 153-SS exhibit the same parameters (as they are enantiomers) while the mesomer 153-RS exhibit two additional structures. In addition, due to its centrosymmetry 153-RS adopt a coplanar structure with a π-π stacking of 3.28Å unlike 153-RR or 153-SS exhibit a twisted structure associated to inferior intermolecular stacking (3.47Å). Among these stereoisomers 153-RS possesses the highest crystallinity and carrier mobility (μ_h = 2.80 10^{-3} cm^2 V^{-1} s^{-1} (after annealing at 100°C for 10 min.), for 153-RR μ_h = 8.60 10^{-4} cm^2 V^{-1} s^{-1} and 153-SS μ_h = 9.10 10^{-4} cm^2 V^{-1} s^{-1}). Unlike to 153, 153-RS exhibits larger crystalline domain size and better intermolecular contacts demonstrating the significant effect of the isomeric purity on the material’s optoelectrical properties that could be benefit to improve the photovoltaic performances.

As seen in the previous sections, the most reported BHJ solar cells combine a small molecule (p-type) and exclusively a fullerene derivative (n-type). However, there still need to develop other n-type materials that can be used as alternatives of fullerene derivatives. For this purpose, Nguyen et al. envisioned to investigate a novel heterojunction based on their DPP chromophores, such as 153 and/or 154 and the dicyanoimidazole 155 (vinazene) acceptor (Scheme 78).

BHJ solar cells were fabricated with the architecture ITO/PEDOT: PSS/153 or 154:155a or 155b/Al. Current-density characteristics were determined with different blend ratios and at different annealing temperatures and compared with devices made from PC_60BM as acceptor. As expected, the nature of the alkyl chains on vinazene derivatives 155 influences the optoelectrical properties: the less hindered and bulky hexyl chain led to more favorable morphology than its 2-ethylhexyl counterpart. Upon thermal annealing marginal improvement was observed. Nevertheless the best photovoltaic performances were obtained with a blend 154:155a (1:1 ratio, 80°C) 0.80% (V_{OC}= 1.23V, J_{SC}= 2.30 mA.cm^{-2}, FF= 0.28) and 154:155b (1:1 ratio, 110°C) 1.10% (V_{OC}= 1.08V, J_{SC}= 2.76 mA.cm^{-2}, FF= 0.35).

Higher PCE values were achieved when PC_60BM was used as acceptor up to 3.72% in the case of the blend 154: PC_60BM (1:1, 110°C, V_{OC}= 0.84V, J_{SC}= 9.26 mA.cm^{-2}, FF= 0.48) but with lower V_{OC}.
A more deeply structure-properties relationship on 153 analogues have been conducted by Roncali et al. using a combination of benzothiophene or benzofuran with thielenyl or furanyl-DPP-based D-A-D molecules (153, 156) as well the replacement of the carbonyl groups of DPP by thioketo (157) (Scheme 79).107 Interestingly, within the series 156 the composition of the side chains has no effect on the HOMOLUMO levels as evidenced by the absorption spectroscopy and the cyclic voltammetry. Furthermore and surprisingly using a planar heterojunction device structure ITO/PEDOT: PSS (40nm)/153-156 (20nm)/C60(50nm)/Al (100nm), only molecule 153 gave rise to a reasonable PCE of 2.50% (VOC = 0.72V, JSC = 7.90 mA.cm−2, FF= 0.39) while for the others structures the PCE did not exceed 0.80%. Hence, minor structural modifications may cause dramatic changes on the optoelectrical properties. Additionally, although the replacement of the carbonyl of DPP by thioketo groups produces a significant reduction of the bandgap extension of the π-conjugated system, by the insertion of an acetylene linkage between the pyrene end-group and the DPP (163), and its consequences on PCE was investigated by Park et al. (Scheme 80).108 The insertion of the linkage favors a more planar geometry of the π-conjugated backbone for derivatives 163, as supported by DFT calculation. This behavior should be beneficial to increase the PCE relative to 161. BHJ with the structure ITO/PEDOT: PSS (30-40nm)/161 or 163: PC71BM (7:3 ratio, 130nm)/Al (100nm) were prepared according to reported procedures. The performances of the solar cells were recorded under standard illumination conditions AM1.5 giving a PCE of 2.95% (VOC = 0.85V, JSC = 8.89 mA.cm−2, FF= 0.42) that is almost 6.5 times greater than 161 measured in the same conditions (0.46% (VOC = 0.79V, JSC = 2.38 mA.cm−2, FF= 0.27). The improved performances were unambiguously assigned to the acetylene linkage that induced a more planar backbone facilitating the intermolecular contacts and increased absorption.

Scheme 81

Extension of the π-conjugated system, by the insertion of an acetylene linkage between the pyrene end-group and the DPP (163), and its consequences on PCE was investigated by Park et al. (Scheme 82).108 The insertion of the linkage favors a more planar geometry of the π-conjugated backbone for derivatives 163, as supported by DFT calculation. This behavior should be beneficial to increase the PCE relative to 161. BHJ with the structure ITO/PEDOT: PSS (30-40nm)/161 or 163: PC71BM (7:3 ratio, 130nm)/Al (100nm) were prepared according to reported procedures. The performances of the solar cells were recorded under standard illumination conditions AM1.5 giving a PCE of 2.95% (VOC = 0.85V, JSC = 8.89 mA.cm−2, FF= 0.42) that is almost 6.5 times greater than 161 measured in the same conditions (0.46% (VOC = 0.79V, JSC = 2.38 mA.cm−2, FF= 0.27). The improved performances were unambiguously assigned to the acetylene linkage that induced a more planar backbone facilitating the intermolecular contacts and increased absorption.

Scheme 82

Controlling the morphology of the donor/acceptor domains in the active layer is of crucial importance to optimize the performance of the solar cells. Stupp et al. proposed recently an elegant strategy by combining synergistically different weak interactions within the active layer (namely H-bonds and π-π interactions) to control the supramolecular organization of each component upon proper and optimized conditions. Thus, they have combined a trans-1,2-diaminocyclohexene building block, that possesses the capability to engender H-bonding, with the DPP chromophore 164 affording the hairpin-shaped chromophore 165 (Scheme 83).111 Upon a careful stepwise cooling process associated to a minimal solution stirring led to the formation of a supramolecular nanowire driven by H-bonds and π–π stacking. This supramolecular arrangement should be helpful to improve the electrooptical properties. Additionally, the shaped of 165 is also favorable to create cavities allowing better cofacial interfaces between donor and acceptor. Solar cells were fabricated using 165 as donor and PC71BM as acceptor under different conditions (systematically donor: acceptor ratio, temperature and solvent). Hence, a blend of disrupted assembly of 165: PC71BM affords only a PCE of 1.40 10−3% while a 400-fold greater PCE (0.53%, VOC = 0.66V, JSC = 2.98 mA.cm−2, FF= 0.27) was achieved with devices fabricated with assembled 165 solutions. If 164 is used under the same conditions a PCE of 0.24% (VOC = 0.57V, JSC = 1.43 mA.cm−2, FF= 0.29) was obtained, nevertheless in this case the concentration of the active component is higher attesting de facto that the self-assembling contributes significantly to enhance the device performances.

To get further insights various terminal modifications have been explored by Frechet et al. in order to promote their solid-state self-assembly via end-to-end π-π interactions (Scheme 81).109 End-groups were selected for their tendency to induce strong π-stacks like pyrene (derivatives 161-162) and compared with non-planar analogues (159) and planar ones bearing non-coplanar substituents (160). Thin-film BHJ solar cells were fabricated with the following architecture ITO/PEDOT: PSS (30-40nm)/159-162: PC71BM/Al (100nm). Devices prepared from 159 to 161 where optimized at a 1:4 donor: PC71BM ratio while a blend ratio of 2:1 was used for 162. Increasing the donor part in the devices comprising compounds 159-161 leads to poorer conversion efficiencies even though an enhancement of the hole-mobilities was gained. In contrast in devices made from 162 higher concentration of donor up to the optimized 2:1 ratio enhanced both the fill factor and the hole-mobilities suggesting that pyrene promotes the molecular packing and film morphology in the solid-state that are favorable to an increase of the PCE. Indeed, the highest PCE of 3.73% (VOC = 0.76V, JSC = 8.30 mA.cm−2, FF= 0.58) was achieved for 162b after thermal annealing.

Scheme 80

To fine tune the device performances Li et al. have investigated and synthesized highly polarizable DPP-based chromophores 158 using thiophene (158a), selenophene (157b) and/or fused thienothiophene (158c) as π-conjugated side-chains (Scheme 80).109 Electrooptical properties and films properties were analyzed revealing that blends of 158b and PC61BM exhibited the best film-forming characteristics and 158c the poorer packing order. ITO/PEDOT: PSS/158: PC61BM/Ca/Al devices were fabricated and evaluated. Increasing the donor: acceptor ratio (1:1, 2:1, 3:1) leads to an improvement of the photovoltaic performances in all cases. As expected from the solid-state properties analysis the best performance was achieved for 158b with a PCE of 2.30% (3:1, VOC = 0.86V, JSC = 5.81 mA.cm−2, FF= 0.46) while 160c exhibits the lowest PCE (1.25% (3:1, VOC = 0.79V, JSC = 4.51 mA.cm−2, FF= 0.36).
Besides DPP, other interesting acceptors could be used for the synthesis of novel D-A-D structures. Among them, fluorenone has been widely used in this context. To study the role of fluorenone on photovoltaic performances, symmetrically chromophores bearing different π-conjugated side chains have been synthesized by Demadrille et al. (Scheme 84). Solar cells were fabricated with the following device structure ITO/PEDOT: PSS/166-169: PC61BM (1:2 or 1:1 ratio, 100 ±20nm)/LiF (0.80nm)/Al. As expected the molecules possessing the strongest π-conjugated systems have exhibited the lowest conversion efficiencies. The highest PCE reported for these compounds reaches 1.19%, for 169 (VOC= 0.82V, JSC= 3.61 mA.cm^2, FF= 0.40) after post-thermal treatment at 70°C at a 1:1 ratio. For compounds 166-168 PCEs are ranged from 0.14% to 0.58% (VOC= 0.72V, JSC= 0.73 mA.cm^2, FF= 0.27 (166), (VOC= 0.52V, JSC= 0.91 mA.cm^2, FF= 0.29, (167), (VOC= 0.68V, JSC= 2.67 mA.cm^2, FF= 0.32 (168))) under the same conditions and without thermal annealing. Furthermore, for 166 and 167 increasing the ratio from 1:1 to 1:2 gives to a substantial improvement of the PCE up to 0.37% (VOC= 0.67V, JSC= 1.85 mA.cm^2, FF= 0.30) and 0.42% (VOC= 0.75V, JSC= 1.86 mA.cm^2, FF= 0.30) respectively. Further enhancement can be achieved by thermal annealing at 80°C.

Encouraged by the results reported on DPP oligothiophenes by Nguyen, isoindigo-based chromophores in conjunction with thiophene as π-conjugated donor were developed and evaluated by Reynolds et al. (Scheme 85). Compounds 170 and 171 present remarkable broad spectra in the solid-state with estimated optical band-gaps of 1.67eV and 1.72eV, respectively. BHJ solar cells were made from spin-coated chlorobenzene solution of 170-171: PC61BM blends onto ITO/PEDOT: PSS glass substrate. Various parameters have been optimized (comprising the solution concentrations, blends ratio, annealing temperature) in order to achieve the highest PCEs. For instance, optimized blend ratios were found to be 1:1 for 170 and 1:5:1 for 171. After thermal annealing at 100 °C, the device fabricated from 170 showed significant better photovoltaic performances (PCE=1.76%) than the solar cell made from 171 (0.55%) (VOC= 0.74V, JSC= 6.30 mA.cm^2, FF= 0.38 and 0.66V, JSC= 2.40 mA.cm^2, FF= 0.36, respectively). The higher PCE for 170 was assigned to a better solid-state ordering, greater film morphology and lower energy gap.

Recently, tetracyanobuta-1,3-diene (TCBD) has been developed as an efficient electron-withdrawing group for the synthesis of NLOphores with improved nonlinear properties and thermal stabilities. Thanks to its intrinsic properties, Roncali et al. have envisioned to use such TCBD for the design of active materials in photovoltaic devices. Thus centrosymmetric and non-centrosymmetric D-A-D structures have been designed and synthesized (Scheme 86). Compounds exhibit non-planar structures (restricting π -electron delocalization in the TCBD core) associated to good solubility and reduced aggregation in solution, high oxidation potentials and strong absorption properties. Bilayer heterojunction solar cells have been fabricated with the following structure ITO/PEDOT: PSS (40nm)/172-174 (25nm)/C60 (20nm)/Al (100nm) and PCEs were determined under AM1.5 conditions at 90 mW/cm². As spin-casted from chloroform solution at 10 mM, chromophore 172 led to a PCE of 1.08% (VOC= 0.97V, JSC= 3.06 mA.cm^2, FF= 0.33) while this PCE is reduced down to 0.56% when the chromophore is deposited by vacuum as a film of 13 nm. These encouraging results demonstrate the interest to use such acceptor for developing BHJ solar cells but improvements have to be pursued.

D – A – D – A – D

Further improvements can be achieved by inserting additionally D-A π-conjugated chromophores to D-A-D architectures leading to D-A-D-A-D structures that should exhibit higher performances due to the extended π-conjugated backbone. Nevertheless, careful disposition and selection of the D and A have to be taken into account in order to optimize the optoelectrical properties.

Recently, Lee et al. have studied and compared the properties of D-A-D having the bithiazole (175) and the fused bithiazole (176, analogous of 149) (Scheme 87). BHJ solar cells have been fabricated with the following structure ITO/PEDOT: PSS/175-176: PC71BM (65-75nm)/Al and evaluated. The best performances, after device optimizations, were found to be 1.30% (1:3 ratio, VOC= 0.97V, JSC= 4.63 mA.cm², FF= 0.29) and 2.39% (1:3 ratio, VOC= 0.94V, JSC= 6.49 mA.cm², FF= 0.39). These performances might be improved by further optimization of the device morphology using processing additives, annealing, and so on. Additionally, it was speculated that the design of new molecular structures with these thiazole derivatives might also provide better tunings for seeking desirable band gap and energy levels and thereby enable better performance.

Hence, from these thiazole-based chromophores were designed and synthesized the extended π-conjugated systems 177-179 depicted in the Scheme 87. Incorporation of fused π-conjugated aromatic units into small molecules have the benefit to stabilize the quinoidal structure and to increase both the rigidity and coplanarity of the molecular backbone. In turn, improved charge transport characteristics, reduction of the bandgap and enhancement of the absorption properties are expected that might all contribute to greater device performances.

BHJ organic solar cells were prepared as follow glass/ITO/PEDOT: PSS (40nm)/177-179: PC71BM (60-70nm)/LiF (0.50nm)/Al (120nm). Compounds 177 and 178 exhibited similar absorption profiles in solution, however in the solid state 177 showed a better well-defined vibronic splitting than 178 indicating a greater efficient π-stacking and strong intermolecular interaction in the film confirmed by thin-film XRD. Furthermore, their electrochemical properties are consistent with their optical properties and molecular structures.
The device performances were evaluated under AM1.5 conditions at different weight ratios (1:1 to 1:4) and using chloroform (CF) or chlorobenzene (CB) as solvents. The best performances resulting from the optimization of the conditions led to a PCE of 1.09% for 177 (1:3 ratio in CF, $V_{OC} = 0.81$ V, $J_{SC} = 3.35$ mA cm$^{-2}$, $FF = 0.41$) and 1.62% for 178 (1:1 ratio in CF, $V_{OC} = 0.76$ V, $J_{SC} = 4.47$ mA cm$^{-2}$, $FF = 0.48$). Despite to better solid-state properties of 178, the greater PCE attained for 179 than 178, even when similar conditions were used (PCE is only 0.76% for 178 when 1:1 ratio is used), is ascribed to the finest film morphology for 178. Analogously to 175 and 176, changing the acceptor from 177 to 179 should lead to enhanced performances in favor of 179. However, the device characteristics gave for the best BHJ solar cell a PCE of 1.44% (1:2 ratio in CB, $V_{OC} = 0.75$ V, $J_{SC} = 5.10$ mA cm$^{-2}$, $FF = 0.38$) that is inferior to what is expected and obtained for 178. Nevertheless under the same conditions, i.e. 1:2 ratio in CB, 178 led to a PCE of 0.65% ($V_{OC} = 0.73$ V, $J_{SC} = 3.01$ mA cm$^{-2}$, $FF = 0.29$) confirming that fused bithiazole is a better acceptor than bithiazole.

Following the same strategy Marks et al have studied the influence of the shape of the naphthodithiophene core, i.e. linear (NDT) vs zig-zag (zNDT), on the photovoltaic performances of D-A-D-A-D comprising a DPP group as acceptor (Scheme 88).

Interestingly, by changing the shape of the central core from NDT to zNDT, both 180 and 181 exhibit the same optical properties and the same optical bandgap (1.70 eV). However 181 exhibits a predictable reduced oxidation potential compared to 180 that should affect in turn the $V_{OC}$. BHJ solar cells were fabricated with the structure ITO/PEDOT: PSS/P380-181/PC$_{61}$BM/LiF/Al and evaluated. The device performances after optimizing the donor: acceptor ratio and the thermal annealing (1.51, 110°C for 180 and 1.21, 140°C for 181) led to a PCE of 4.00% ($V_{OC} = 0.84$ V, $J_{SC} = 11.20$ mA cm$^{-2}$, $FF = 0.43$) and 4.40% ($V_{OC} = 0.75$ V, $J_{SC} = 11.70$ mA cm$^{-2}$, $FF = 0.50$) respectively. The results are consistent with the greater hole-mobility for 181 ($\mu_h = 1.00 \times 10^{-4}$ cm$^2$ V$^{-1}$s$^{-1}$) than 180 ($\mu_h = 2.50 \times 10^{-2}$ cm$^2$ V$^{-1}$s$^{-1}$) leading to a substantial enhancement of the short-circuit current in favor of 181 due to a more efficient charge separation and reduced recombination process.

Scheme 88

Recently, the indacenodithiophene (IDT) has sparked great interest as donor molecular building block due to its intrinsic properties, for instance a planar rigid structure, beneficial for the fabrication of SMOSCs. Thus, Hou et al have designed and synthesized small IDT-based chromophores with the structure 182 depicted on scheme 89. The synthesized chromophore presents optoelectronic properties that make it suitable for the fabrication of organic solar cells ($E_g = 1.80$ eV, broad absorption in the visible region, low-lying HOMO energy, high thermal stability).

Hence, BHJ solar cells based on the blend 182:PC$_{61}$BM have been fabricated with the following architecture ITO/PEDOT: PSS (30nm)/182:PC$_{61}$BM (80-100nm)/Ca/Al. Several conditions have been tested by varying the donor: acceptor ratio (from 1:1 to 1:4) leading to the highest PCE of 4.25% ($V_{OC} = 0.93$ V, $J_{SC} = 9.42$ mA cm$^{-2}$, $FF = 0.48$) for a 1:3 ratio. No further improvements have been achieved by the addition of DIO as solvent additive or even by thermal annealing. The results indicated that chromophores based on IDT core are promising materials for the fabrication of SMOSCs.

Scheme 89

Alongside the work of Marks based on NDT derivatives, Lee et al. have design new molecular D-A-D-A-D architectures based on the BDT unit, analogously to those described in scheme 32. Chromophore 183 has been synthesized following known procedures and evaluated as donor in BHJ solar cells (Scheme 90). The solar cells based on the blend 183:PC$_{71}$BM have fabricated at different weight ratios, from 1:0.5 to 1:4, with the structure ITO/PEDOT: PSS (40nm)/183:PC$_{71}$BM (50-55nm)/LiF (0.5nm)/Al (120nm). The best performance achieved without any post-treatment gave a PCE of 1.18% ($V_{OC} = 0.83$ V, $J_{SC} = 4.80$ mA cm$^{-2}$, $FF = 0.29$) at a 1:3 weight ratio. Impressive enhancement of the PCE up to 2.83% ($V_{OC} = 0.85$ V, $J_{SC} = 7.81$ mA cm$^{-2}$, $FF = 0.39$) was produced upon thermal annealing at 180°C due to a higher π-stacking order and increased hole-mobility up to $\mu_h = 2.70 \times 10^{-4}$ cm$^2$ V$^{-1}$s$^{-1}$ (untreated film $\mu_h = 1.40 \times 10^{-6}$ cm$^2$ V$^{-1}$s$^{-1}$) corroborating the solid-state analysis. In addition the alkoxy side-chains are helpful to ensure good film morphology and phase separation.

Scheme 90

DTS core has also demonstrated to be an important and useful molecular scaffold for the construction of active component for SMOSCs (see for example scheme 32). To get further insights on the molecular design of new active donors based on the DTS unit, Bazan et al. have synthesized new D-A-D-A-D structures in association with thiadiazolopyridine (PT) as acceptor and deeply studied the parameters affecting their photovoltaic performances (Scheme 91). From the solution to the solid-state the chromophore showed a broadening and a red shift of the optical absorption along with the emergence of a vibronic splitting coherent with a more rigid and ordered structure in the film ($E_g = 1.51$ eV). Such features lead to greater π-electron delocalization across the molecular backbone and enhanced inter-chromophore interactions. Upon thermal annealing up to and below 110°C the lowest energy absorption peak is enhanced consistent with a better π-stacking. In agreement with the opto-electrochemical properties of 184, SM-BHJ photovoltaic devices were made using a standard device architecture of ITO/Plexcore® OC AQ/184: PC$_{71}$BM (85-105nm)/Al. The active layers were optimized by varying the 184:PC$_{71}$BM weight ratios at 3 different thickness layers and thermal annealing 2 min. at 110°C. From this set of experiments the best performances was achieved for a 60:40 ratio with a blend thickness of 85nm leading to a PCE of 3.20% ($V_{OC} = 0.70$ V, $J_{SC} = 10.90$ mA cm$^{-2}$, $FF = 0.42$). Determination of the hole and electrons mobilities ($\mu_h = 1.50 \times 10^{-6}$ cm$^2$ V$^{-1}$s$^{-1}$, $\mu_e = 5.30 \times 10^{-3}$ cm$^2$ V$^{-1}$s$^{-1}$, respectively) leads to unbalanced charge carrier transport and may be responsible for the lower fill factor.

Scheme 91

In close collaboration with A.J. Heeger, mechanistic studies have been performed on the effect of solvent additives processed
active layers during the film morphology formation and their impact on the photovoltaic efficiencies.\(^{123, 124}\) During this study it has been clearly demonstrated that the use of DIO improves, beyond doubts, the interfacial charge transfer between donor and acceptor domains. This effect was also investigated theoretically. It was also demonstrated that Si-based chromophores are more efficient than their carbon counterparts.\(^{125}\) Carbon analogues of \(184\) showed only a PCE of 0.2% (as cast, \(V_{OC} = 0.66\) V, \(J_{SC} = 1.00\) mA cm\(^{-2}\), \(FF = 0.26\)) and 0.09% (annealed, \(V_{OC} = 0.50\) V, \(J_{SC} = 0.60\) mA cm\(^{-2}\), \(FF = 0.28\)) vs a PCE of 0.9% (as cast, \(V_{OC} = 0.67\) V, \(J_{SC} = 3.60\) mA cm\(^{-2}\), \(FF = 0.29\)) and 3.3% (annealed, \(V_{OC} = 0.70\) V, \(J_{SC} = 9.90\) mA cm\(^{-2}\), \(FF = 0.47\)) respectively for the blend \(185:\)PC\(_3\)BM without additives. The lower efficiency for the carbon analogue is assigned to the presence of larger crystalline domains.

Heeger et al. have conducted additional structural modification on chromophore \(184\). Thus, chromophore \(185\) corresponding to an isomer of \(184\) has been designed and synthesized (Scheme \(92\)).\(^{126}\) The difference came only from the position of the N atom on the PT scaffold. Different weight ratios were evaluated and it was found that at a 7:3 ratio a remarkable PCE of 4.52% \((V_{OC} = 1.40\) V, \(J_{SC} = 12.50\) mA cm\(^{-2}\), \(FF = 0.59\)) was achieved for the blend \(185:\)PC\(_3\)BM with a device structure ITO/MoO\(_3\)/BM/Al.\(^{127}\) Further improvement has been achieved by the introduction of 0.25% v/v DIO as solvent additive an increase of the PCE up to 7.00% \((V_{OC} = 0.81\) V, \(J_{SC} = 12.80\) mA cm\(^{-2}\), \(FF = 0.68\)). Encouraged by these results further investigations in order to improve the device performances were pursued notably by using solvent additives. Progressive increase of DIO up to 0.4 v/v% followed by a thermal treatment at 70°C led to an impressive PCE of 7.60% \((V_{OC} = 0.81\) V, \(J_{SC} = 12.80\) mA cm\(^{-2}\), \(FF = 0.68\)). Interestingly, a non-fullererene acceptor, namely the perylene diimide PDI, has been used for the fabrication of BHJ solar cells in combination with \(188\) as donor.\(^{128}\) PDI exhibits complementary absorption spectra to that of \(187\) leading to a broad coverage of the visible spectrum. Screenings of the parameters that can influence the performances such as blend ratios, solvent additives and thermal annealing have been examined. Optimized devices were obtained for a 1:1 blend ratio dissolved in CB (30 mg mL\(^{-1}\)) with 0.4% v/v of DIO. BHJ solar cells were fabricated with the standard architecture of ITO/PEDOT: PSS (45nm)/P3HT:PC\(_{70}\)BM/Ca/Al and evaluated.\(^{129}\) As a cast film without any post-treatment the device delivers a moderate photocurrent with a PCE of 1.80% \((V_{OC} = 0.78\) V, \(J_{SC} = 6.60\) mA cm\(^{-2}\), \(FF = 0.36\)). To improve the above efficiency, post-deposition thermal annealing were investigated and it was shown that heating 10 min. at 130°C gives an increase of the PCE up to 5.80% \((V_{OC} = 0.82\) V, \(J_{SC} = 10.80\) mA cm\(^{-2}\), \(FF = 0.65\)).

The impact of the N atom and its position on \(184\) analogues was deeply investigated in a further work of Bazan (Scheme \(93\)) and compared to the non-nitrogen analogue \(187.\)\(^{127}\) The position of the N-atom affects the self-assembling properties in the BHJ film and might be correlated to the dipole-dipole induced-orientation in the solid-state. Chromophores \(183-186\) exhibit similar optical behaviors in solution and in the solid-state with a pronounced vibronic splitting compared to \(187.\) Optical band gap are determined to be 1.50eV for \(185\) and \(186,\) 1.52eV for \(184\) and 1.58eV for \(187.\) The small molecules BHJ solar cells were fabricated with the structure of ITO/MoO\(_3\)/BM/Al and tested under the same conditions and same donor: acceptor weight ratios. The devices were evaluated without any treatment giving PCEs of 4.52% for \(185 (V_{OC} = 0.80\) V, \(J_{SC} = 12.50\) mA cm\(^{-2}\), \(FF = 0.45),\) 2.47% for \(186 (V_{OC} = 0.75\) V, \(J_{SC} = 8.90\) mA cm\(^{-2}\), \(FF = 0.37),\) 1.78% for \(184 (V_{OC} = 0.73\) V, \(J_{SC} = 7.40\) mA cm\(^{-2}\), \(FF = 0.33)\) and 0.19% for \(187 (V_{OC} = 0.83\) V, \(J_{SC} = 0.90\) mA cm\(^{-2}\), \(FF = 0.26).\) Using 0.25% v/v DIO as solvent additive an increase of the photovoltaics efficiencies were obtained in most cases leading to the following PCEs 6.70% for \(185 (V_{OC} = 0.78\) V, \(J_{SC} = 14.40\) mA cm\(^{-2}\), \(FF = 0.59),\) 2.47% for \(186 (V_{OC} = 0.73\) V, \(J_{SC} = 12.70\) mA cm\(^{-2}\), \(FF = 0.60),\) 3.61% for \(184 (V_{OC} = 0.72\) V, \(J_{SC} = 9.80\) mA cm\(^{-2}\), \(FF = 0.45)\) and 0.18% for \(187 (V_{OC} = 0.78\) V, \(J_{SC} = 0.90\) mA cm\(^{-2}\), \(FF = 0.25).\) These results imply that the design of new organic materials for SMOSCs needs to consider not only the electronic structure and the nature of the side-chains but also to take into account other interactions that can enhance/direct the self-assembly in the film.

Further improvement has been achieved by the introduction of a fluorine atom in place of the N-atom (Scheme \(94.\) This structural modification renders the chromophore less sensitive to acidic medium and may increase its stability. Photovoltaic devices were fabricated using the general structure of ITO/PEDOT: PSS/\(188\):PC\(_3\)BM/Ca/Al and evaluated.\(^{128}\)

Scheme \(94\)

As expected compounds \(189\) and \(190,\) that are oligomeric analogues of \(185,\) exhibit a red-shift absorption spectra in solution and in the solid-state. From the absorption onsets, the optical band-gaps have been determined to be 1.44eV and 1.41eV respectively which agrees with the HOMO (-5.17eV, -5.04eV respectively) and LUMO energy levels determined by cyclic

Scheme \(95\)

As expected compounds \(189\) and \(190,\) that are oligomeric analogues of \(185,\) exhibit a red-shift absorption spectra in solution and in the solid-state. From the absorption onsets, the optical band-gaps have been determined to be 1.44eV and 1.41eV respectively which agrees with the HOMO (-5.17eV, -5.04eV respectively) and LUMO energy levels determined by cyclic
voltammetry. In addition, the presence of a vibronic fine structure is associated to specific intermolecular interactions resulting from π–π overlaps in the thin film. Moreover, a thermal annealing (100°C) has shown to be benefit for improving the molecular organization as more intense π–π interactions were observed after a thermal treatment, which is more pronounced for 189. These features are important to ensure efficient charge transport within the film. Hence, relatively high hole-mobilities were measured from the films \( \mu_h = 1.00 \times 10^{-2} \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1} \) (189) and \( \mu_h = 6.00 \times 10^{-3} \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1} \) (190) even after thermal treatment beyond 200°C moderate hole-mobilities are maintained \( \mu_h = 3.00 \times 10^{-3} \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1} \) at 230°C (189) and \( \mu_h = 2.60 \times 10^{-3} \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1} \) at 250°C (190).

Owing to these considerations, BHJ solar cells have been prepared with the standard structure ITO/MoOx/PCBM:PC70BM/Al and tested. An optimal weight ratio of 60:40 for 189:PCBM blend followed by a thermal treatment at 100°C (10 min.) afford a PCE of 5.80% (\( V_{oc}= 0.71 \text{V}, J_{sc}= 13.60 \text{mA} \cdot \text{cm}^{-2}, \) FF= 0.60) while a PCE of 6.50% (\( V_{oc}= 0.66 \text{V}, J_{sc}= 15.20 \text{mA} \cdot \text{cm}^{-2}, \) FF= 0.65) is reached for 190:PCBM blend at ratio 50:50 without any post-deposition treatment. Indeed, it is worth noticing that for 190:PCBM blends thermal annealing does not affect the photovoltaic performances, which are also quite insensitive to the blend ratio composition. The general trends observed for 189 and 190 will be useful to design other materials with improved performances that can withstand major environmental changes.

**Conclusions**

The last decade has witnessed a rapid progress of organic photovoltaics boosted by the design and synthesis of novel π-conjugated small donor molecules, the control and optimization of both device processing and fabrication. Although some important progress has been reached, current challenges remain to further improve efficiency, durability and cost-effectiveness in order to compete with silicon-based solar cells. Within this article we aimed at suggesting some guiding principles for designing new efficient and emerging alternatives to π-conjugated polymers that are solution-processable, facile to synthesize, simple to purify and monodisperse for a batch-to-batch reproducibility. To date, efficiencies up to 7% have been reported and important understandings were found. The main approaches consist in either a modification of the photoactive layer itself or addition of extra transport/blocking layers to facilitate the contacts. As the injection and the photovoltage critically depend on the interface, molecular modifications or physical treatments have been proposed to appropriately modulate the injection barriers either at the electron or hole collecting electrodes (for instance, LiF or PEDOT: PSS respectively). As seen, optimized light harvesting material is also required to match with the solar spectrum and to enhance the photoconversion efficiency. Substantial increases of the solar photon harvesting have led to the design and synthesis of suitable organic low band-gaps materials with appropriate side-groups. Another way is to replace the PCBM by PCBM or by changing the nature of the acceptor with the aim to increase the absorption range. In addition, novel solutions can be applied to achieve the same effect, for instance the combination of small molecules with complementary absorption spectrum might be tested in the future. More importantly, the nanoscale morphology of the blend dramatically influences the photovoltaic properties.

Hence, in order to increase the efficiency it is crucial to improve the miscibility of the mixed materials leading to finer phase segregation. This can be achieved by adding plasticizer additive, choosing the best solvents, controlling the evaporation time, their wettability and surface interactions, applying thermal annealing and varying the D-A composition. It is of crucial interest to develop also new methodologies and concepts to control the nanomorphology of the bulk heterojunction based on our supramolecular chemistry knowledge, crystallisation processes and solid-sate controlled orientation.

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**Notes and references**

Thiophene-based push-pull chromophores for small molecule organic solar cells (SMOSCs).
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Scheme 1
Scheme 8

Scheme 9

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Scheme 12

Scheme 13
Scheme 17

Scheme 18
Scheme 19

Scheme 20
Scheme 23

Scheme 24
Scheme 31

Scheme 32
Scheme 48

Scheme 49

$102a$: n = 1, R = H
$102b$: n = 2, R = C$_6$H$_{13}$
$102c$: n = 3, R = C$_6$H$_{13}$

Scheme 50

$103a$: n = 1
$103b$: n = 2
$103c$: n = 3

Scheme 51

Scheme 52
Scheme 53

Scheme 54

Scheme 55
Scheme 63

Scheme 64
Scheme 65

132a: R₁ = n-dodecyl, R₂ = n-hexyl
132b: R₁ = n-dodecyl, R₂ = H
132c: R₁ = 2-ethylhexyl, R₂ = H

133a: n = 0
133b: n = 1
133c: n = 2

Scheme 66
Scheme 78

Scheme 79

Scheme 80
Scheme 84

Scheme 85

Scheme 86
Scheme 87

Scheme 88

Scheme 89