

US009029499B2

c12) **United States Patent**

Yu et al.

(54) **SEMICONDUCTING POLYMERS**

- (71) Applicant: **University of Chicago,** Chicago, IL (US)
- (72) Inventors: **Luping Yu,** Chicago, IL (US); **Yongye Liang,** Stanford, CA (US)
- (73) Assignee: **University of Chicago,** Chicago, IL (US)
- (*) Notice: Subject to any disclaimer, the term of this patent is extended or adjusted under 35 U.S.C. 154(b) by O days.
- (21) Appl. No.: **14/183,151**
- (22) Filed: **Feb.18,2014**

(65) **Prior Publication Data**

US 2014/0235817 Al Aug. 21, 2014

Related U.S. Application Data

- (63) Continuation of application No. 13/888,738, filed on May 7, 2013, now Pat. No. 8,653,228, which is a continuation of application No. 13/054,719, filed as application No. PCT/US2009/044364 on May 18, 2009, now Pat. No. 8,436,134.
- (60) Provisional application No. 61/082,071, filed on Jul. 18, 2008, provisional application No. 61/082,418, filed on Jul. 21, 2008.
- (51) **Int. Cl.**

(52) **U.S. Cl.** CPC *C08G 75106* (2013.01); *C08G 73118*

(IO) **Patent No.: US 9,029,499 B2**

(45) **Date of Patent: May 12, 2015**

(2013.01); *C08G 73122* (2013.01); *H0JL 5110036* (2013.01); *H0JL 5114253* (2013.01); *Y02E 10/549* (2013.01); *C09B 69/109* (2013.01)

(58) **Field of Classification Search** USPC .. 528/377, 380, 378 See application file for complete search history.

(56) **References Cited**

U.S. PATENT DOCUMENTS

(Continued)

FOREIGN PATENT DOCUMENTS

(Continued)

OTHER PUBLICATIONS

Chen, et al., "Hierarchical Nanomorphologies Promote Exciton Dissociation in Polymer/Fullerene Bulk Heterojunction Solar Cells," *Nano Letters,* vol. 11, pp. 3707-3713 (2011).

Facchetti, "PI-Conjugated Polymers for Organic Electronics and Photovoltaic Cell Application," *Chemistry of Materials,* vol. 23, pp. 733-758 (2011).

(Continued)

Primary Examiner — Shane Fang

(74) *Attorney, Agent, or Firm* - Brinks Gilson & Lione; Ryan L. Marshall

(57) **ABSTRACT**

Novel semiconducting photovoltaic polymers with conjugated units that provide improved solar conversion efficiency that can be used in electro-optical and electric devices. The polymers exhibit increased solar conversion efficiency in solar devices.

19 Claims, 8 Drawing Sheets

(56) **References Cited**

U.S. PATENT DOCUMENTS

FOREIGN PATENT DOCUMENTS

OTHER PUBLICATIONS

International Search Report for PCT/US2009/044364, dated Aug. 27, 2009, 3 pages.

International Search Report for PCT/US2012/033601, dated Sep. 6, 2012, 3 pages.

International Search Report for PCT /US2013/024337, dated Apr. 16, 2013, 3 pages.

Liang, Y., et al., "Control in Energy Levels of Conjugated Polymers for Photovoltaic Application," *JPhys. Chem.* C, vol. 112, pp. 7866- 7871 (2008).

* cited by examiner

Figure 1

Figure 3

Figure 5a

Figure Sb

Figure 6a

Figure 6b

SEMICONDUCTING POLYMERS

REFERENCE TO RELATED APPLICATIONS

This application continuation of U.S. patent application Ser. No. 13/888,738, filed May 7, 2013, now U.S. Pat. No. 8,653,228 issued Feb. 18, 2014, which is a continuation of U.S. patent application Ser. No. 13/054,719, filed Jan. 18, 2011, now U.S. Pat. No. 8,436,134, issued May 7, 2013, which is a 371 national phase of International Application No. 10 PCT/US2009/044364, filed May 18, 2009, which claims the benefit of U.S. Provisional Application Nos. 61/082,071 and 61/082,418, filed Jul. 18, 2008, and Jul. 21, 2008, respectively, both entitled "Semiconducting Polymers," the contents of which are incorporated herein by reference in their entirety.¹⁵

GOVERNMENT INTERESTS

This invention was made with Government support under Grant No. DMR-0703274, awarded by the National Science ²⁰ Foundation. The Government may have certain rights to this invention.

This invention relates to semiconducting polymers based on semiconducting conjugated polymers. This invention also relates to their use in electro-optical and electronic devices.

BACKGROUND

effect, organic solar cells are very promising. Bulk hetero 40 R^2 have the following formula: Solar energy harvest using photovoltaic effect requires active semiconducting materials to convert light into electricity. Currently, solar cells based on silicon are the dominating technology due to their high conversion efficiency. Recently, 35 solar cells based on organic materials showed interesting features, especially on the potential of low cost in materials and processing. Judging from the recent success in organic light emitting diodes based on a reverse effect of photovoltaic junction (BHJ) made from phase separated blends of semiconducting polymers and fullerides is a popular structure that has been adopted for polymer solar cells. Current solar cells exhibit low power conversion efficiency, about 4%-5%, compared to silicon based solar cells, even after sophisticated 45 device optimizations.

bon-nitrogen (C—N) bonds. The conjugated polymers have a $\,$ 50 formula (I): Conjugated polymers have shown some promise in providing a photovoltaic effect. Conjugated polymers are made of alternating single and double carbon-carbon (C-C) or car- δ -bond backbone of intersecting sp² hybrid orbitals. The p_z orbitals on the carbon atoms overlap with neighboring p_z orbitals to provide π -bonds. The electrons that comprise the π -bonds are delocalized over the whole molecule. These polymers exhibit electronic properties similar to those seen in 55 inorganic semiconductors. The semiconducting properties of the photovoltaic polymers are derived from their delocalized π bonds.

There is a need in the art for polymer solar cells that exhibit increased solar conversion efficiency. 60

BRIEF SUMMARY

Described herein are semiconducting photovoltaic polymers that exhibit high solar conversion efficiencies when used in electro-optical and electronic devices. There are also semiconducting polymers that are used as hole transporting mate-

rial with fullerene derivatives as acceptors in electro-optical and electronic devices. The polymers are designed to achieve a low bandgap for broad absorption in the solar spectrum.

The present disclosure describes a semiconducting polymer of the formula (I):

where X^1 and X^2 are independently O, S, N, NH, a crosslinkable group such as an acrylate group, or $CR¹$ or $CR²$, where R^T and $R²$ are independently H, alkyl, alkoxy, aryl, aryloxy, heteroaryl, heteroaryloxy, cross-linkable moiety, TECHNICAL FIELD

oligo (ethylene glycol); Y^1 and Y^2 are independently O, S, Se,
 $\frac{1}{25}$ amino; Z is an ester, amide, cyano, alkyl, or polyfluoroalkyl, amino; Z is an ester, amide, cyano, alkyl, or polyfluoroalkyl, polychloroalkyl, aryl, heteroaryl; W is H, halogen, cyano, dicyanovinyl, or tricyanovinyl; and n is an integer.

> The semiconducting polymer can also be formula (I) where X^1 and X^2 are CH₂, Y^1 and Y^2 are S, Z is COOR", W is H, R¹ 30 and R^2 are C_7H_{15} , and R" has the formula

The semiconducting polymer can also be formula (I) where X^1 and X^2 are O, Y^1 and Y^2S , Z is COOR", W is F or H, R^1 and

Also provided herein is a photovoltaic film comprising a fullerene derivative and a semiconducting polymer of the

where X^1 and X^2 are independently O, S, N, NH, a crosslinkable group such as an acrylate group, or $CR¹$ or $CR²$, where $R¹$ and $R²$ are independently H, alkyl, alkoxy, aryl, aryloxy, heteroaryl, heteroaryloxy, cross-linkable moiety,

(I)

(I)

30

65

oligo (ethylene glycol); Y^1 and Y^2 are independently O, S, Se, amino; Z is an ester, amide, cyano, alkyl, or polyfluoroalkyl, polychloroalkyl, aryl, heteroaryl; W is H, halogen, cyano, dicyanovinyl, or tricyanovinyl; and n is an integer.

BRIEF DESCRIPTION OF THE DRAWINGS

FIG. 1 is a graph of the current-voltage characteristics of a device made with a semiconducting photovoltaic polymer prepared as described herein. 10

FIG. 2 is a bar graph of the HOMO and LUMO energy levels of certain polymers and fullerene described herein.

FIG. **3** is a line graph of the $J^{0.5}$ vs V plots for the polymers described herein with solid lines being fits of the data points and the thickness of the films indicated in the plots.

FIGS. *4a* through *4i* are TEM images of polymer/fullerene blend films and polymer PC_{61} BM blend films prepared from mixed solvents.

FIGS. **Sa** and **Sb** are CAFM images of polymer/fullerene blend films: LY-14 and LY-18, with the white regions in 20 CAFM (low current) corresponding to the fullerene-rich areas, consistent with hole current images.

FIG. *6a* is a graph of the current-voltage characteristics of polymer/fullerene solar cells prepared from mixed solvents. FIG. 6b is a graph of the external quantum efficiency of 25 LY-16/PC $_{61}$ BM device prepared from mixed solvents.

DETAILED DESCRIPTION OF THE DRAWINGS AND THE PREFERRED EMBODIMENTS

Unless defined otherwise, all technical and scientific terms used herein have the same meaning as commonly understood to one of ordinary skill in the art to which this invention belongs. When describing the compounds, compositions, methods and processes of this invention, the following terms 35 have the following meanings, unless otherwise indicated.

"Alkyl" by itself or as part of another substituent refers to a hydrocarbon group which may be linear, cyclic, or branched or a combination thereof having the number of carbon atoms designated (i.e., C_{1-8} means one to eight carbon atoms). 40 Examples of alkyl groups include methyl, ethyl, n propyl, isopropyl, n butyl, t butyl, isobutyl, sec butyl, cyclohexyl, cyclopentyl, (cyclohexyl)methyl, cyclopropylmethyl, bicyclo [2 .2 .1]heptane, bicyclo[2.2.2]octane, etc. Alkyl groups can be substituted or unsubstituted, unless otherwise indi- 45 cated. Examples of substituted alkyl include haloalkyl, polyhaloalkly, such as polyfluoroalkyl and polychloroalkyl, aminoalkyl, and the like.

group include methoxy, ethoxy, n-propoxy etc.

"Ary!" refers to a polyunsaturated, aromatic hydrocarbon group having a single ring (monocyclic) or multiple rings (bicyclic), which can be fused together or linked covalently. Ary! groups with 6-10 carbon atoms are preferred, where this number of carbon atoms can be designated by C_{6-10} , for 55 example. Examples of aryl groups include phenyl and naphthalene-1-yl, naphthalene-2-yl, biphenyl and the like. Ary! groups can be substituted or unsubstituted, unless otherwise indicated. "Aryloxy" refers to ---O-aryl; and "heteroaryloxy" refers to the group ---0-heteroaryl. 60

The term "amino" refers to -NRR' where Rand R' are independently selected from hydrogen, alkyl, aryl, aralkyl and alicyclic, all except H are optionally substituted; and R and R' can form a cyclic ring system.

"Cyano" refers to $-CN$.

"Ester" refers to $R' C(O)$)— where R' is a hydrogen atom, an alkyl group, an aryl group or an arylheterocyclic ring, as

defined herein. "Arylheterocyclic ring" refers to a bi- or tricyclic ring comprised of an aryl ring, as defined herein, appended via two adjacent carbon atoms of the aryl ring to a heterocyclic ring, as defined herein. Exemplary arylheterocy-5 clic rings include dihydroindole, 1,2,3,4-tetra-hydroquinoline, and the like.

"Halo" or "halogen," by itself or as part of a substituent refers to a chlorine, bromine, iodine, or fluorine atom.

"Haloalkyl," as a substituted alkyl group, refers to a monohaloalkyl or polyhaloalkyl group, most typically substituted with from 1-3 halogen atoms. Examples include 1-chloroethyl, 3-bromopropyl, trifluoromethyl and the like.

The term "heteroaryl," as used herein, refers to a mono-, bi-, or tri-cyclic aromatic radical or ring having from five to ten ring atoms of which at least one ring atom is selected from S, 0, and N; zero, one or two ring atoms are additional heteroatoms independently selected from S, 0, and N; and the remaining ring atoms are carbon, wherein any N or S contained within the ring may be optionally oxidized. Heteroaryl includes, but is not limited to, pyridinyl, pyrazinyl, pyrimidinyl, pyrrolyl, pyrazolyl, imidazolyl, thiazolyl, oxazolyl, isooxazolyl, thiadiazolyl, oxadiazolyl, thiophenyl, furanyl, quinolinyl, isoquinolinyl, benzimidazolyl, benzooxazolyl, quinoxalinyl, and the like. The heteroaromatic ring may be bonded to the chemical structure through a carbon or heteroatom.

"Heteroatom" is meant to include oxygen (0), nitrogen (N), sulfur (S) and silicon (Si).

The photovoltaic property of the polymers described herein was studied in polymer solar cells. Simple solar cells were used with semiconducting polymers of the formulas described herein. The power conversion efficiency (PCE) was greater than 3% in some embodiments. In embodiments that may be preferred, the semiconducting polymers yielded a PCE greater than 5%. The PCE reached 5.6% to 6.1% in some preferred embodiments.

The high conversion efficiency exhibit by the polymers described herein may be the result of an effective light harvest. The cells using the polymers absorb light in almost the entire visible spectrum. The polymer morphology also favors charge separation and charge transport. This can lead to high fill factors.

The polymers described herein are based on the concept that a thienothiophene moiety can support a quinoidal structure and lead to a narrow polymer bandgap that may have an effect on harvesting solar energy. Because the thienothiophene moiety is electron-rich, an electron-with- "Alkoxy" refers to $-$ O-alkyl. Examples of an alkoxy drawing ester group can be introduced to stabilize the result-
oup include methoxy, ethoxy, n-propoxy etc.

The semiconducting polymers described herein may be represented by formula (I):

where X_1 and X_2 are independently O, S, N, NH, a crosslinkable group, or $CR¹$ or $CR²$, where $R¹$ and $R²$ are indepen-

(I)

dently H, alkyl, alkoxy, aryl, aryloxy, heteroaryl, heteroaryloxy, a cross-linkable moiety, or oligo (ethylene glycol); $Y¹$ and Y^2 are independently O, S, Se, amino; Z is an ester, amide, cyano, alkyl, or polyfluoroalkyl, polychloroalkyl, aryl, heteroaryl; and W is H, halogen, cyano, dicyanovinyl, or tricyanovinyl. "n" is an integer or can be an integer from 1 to 200. X^1 and X^2 may be the same or different. Y¹ and Y² may be the same or different. $R¹$ and $R²$ maybe the same or different.

Another embodiment may also comprise semiconducting polymers having the formula (I) where W represents F or H. X^1 and X^2 may independently represent CH₂ or O and Y¹ and $Y²$ may independently represent S in some embodiments that may be preferred.

In some embodiments, Z may represent an ester or polyfluoroalkyl. Preferably, Z may represent COOR" where R" is alkyl, alkoxy, aryl, aryloxy, heteroaryl, heteroaryloxy. More preferably, R" may represent C_mH_{2m+1} where m is an integer from 1 to 30. R" may be branched or, preferably, have the following formulae (II) and (III):

Z may also represent C_nF_{2n+1} where n is an integer from 1 to 12 in some embodiments or C_nH_{2n+1} where n is an integer from 1 to 12 in other embodiments. There are also embodi- 35 ments where R^1 , R^2 , and R'' are different.

In embodiments that may be preferred, the semiconducting polymers have the formula (I) where X^1 and X^2 can independently represent O, Y^1 and Y^2 can independently represents S, Z is COOR", W represents H , R^1 and R^2 independently rep- 40 resent $C_{12}H_{25}$, and R" is the formula (III):

The semiconducting polymers may preferably have the formula (I) where X^1 and X^2 independently represent CH_2 , Y^1 and Y^2 independently represent S, Z represents COOR", W represents H, R^1 and R^2 independently represent $C_{12}H_{25}$, and R" represents C_8H^{17} . The semiconducting polymers may have the formula (I) where X^1 and X^2 independently represent $_{55}$ $CH₂$, $Y¹$ and $Y²$ independently represent S, Z represents COOR", W represents H, $R¹$ and $R²$ independently represent C_7H_{15} , and R" has the formula (III):

6

and Y^2 independently represent S, Z represents COOR", W represents F. R", $R¹$ and $R²$ can independently represent the same formula (III):

The semiconducting polymers may have the formula (I) where X^1 and X^2 independently represent O, Y^1 and Y^2 independently represent S, Z represents COOR", W represents F or H, R ["], R ¹ and R ² can independently represent the formula (III):

and R" is C_8H_{17} . In some preferred embodiments, X^1 and X^2 ²⁰ independently represent \tilde{O} , Y^1 and Y^2 independently represent S, Z represents C_8F_{17} , W represents H, R^1 and R^2 can independently represent $C_{12}H_{25}$, and R" represents C_8H_{17} .

 (II) The semiconducting polymers of the present invention also can have the formula (I) where X^1 and X^2 independently 25 represent O, Y^1 and Y^2 independently represent S, Z represents COOR", W represents H , $R¹$ and $R²$ can independently represent C_8H_{17} , and R" represents $C_{12}H_{25}$. Also, there are embodiments where X^1 and X^2 independently represent O, Y^1 and Y^2 independently represent S, Z represents COOR", W ³⁰ represents H, R¹ and R² can independently represent C₈H₁₇, and R" is the formula (II):

$$
\sum_{\mathbf{C}_2\mathbf{H}_S} C_4\mathbf{H}_9.
$$

In some embodiments that may be preferred, X^1 and X^2 independently represent O, Y^1 and Y^2 independently represent S, Z represents COOR", W represents H, $R¹$ and $R²$ can independently represent C_8H_{17} , and R" has the following formula

$$
\left\langle \vphantom{\sum_{\sum_{i=1}^{k}}^N}\right\rangle_{C_{12}H_{25}}
$$

45

60

65

The semiconducting polymers of the present invention can also be of the formula (IV):

where R represents polyfluoralkyl, polychloroalkyl, or an ester; R' represents alkyl, alkoxy, aryl, aryloxy, heteroaryl, or heteroaryloxy; X represents an integer from 1 to 12; and m is an integer from 1 to 200.

In some embodiments that may be preferred R represents C_nF_{2n+1} where n is an integer from 1 to 12 or -COOC_nH_{2n+1} where n is an integer from 1 to 30. R can represent C_6F_{13} , R^1 can represent C_nH_{2n+1} where n is an integer from 1 to 12. In a preferred embodiment, R^1 represents C_6H_{13} , R represent C_6F_{13} , and X is 8.

Gel permeation chromatography studies showed a number averaged molecular weight of 18.3 k and weight averaged 5 molecular weight of 22.9 k with a polydispersity of 1.25.

The present invention also comprises the conjugates of the semiconducting polymers (I) and (IV) described herein. The conjugates can be joined with electron accepting materials including, but not limited to, Buckminsterfullerene ("fullerene") derivatives. Any electron accepting fullerene can be used with the semiconducting polymers described herein. Such fullerenes can be from C_{60} to C_{90} . In preferred embodiments the fullerene can be C_{61} , C_{71} , C_{81} , or C_{91} . The $_{15}$ fullerene can be [6,6]-phenyl-C61-butyric acid methyl ester (PC61BM), having the following formula:

The polymer and fullerene can blend as a mixture. In another aspect of the invention, the conjugates can be joined with the fullerenes as shown below.

where

 R' =-(CH2)m-, m=2-16

 $X=O, S, NH, -ONH-$

 $X¹$ and $X²$ can be independently O, S, N, NH, a crosslinkable group, or $CR¹$ or $CR²$, and $R¹$ and $R²$ can be independently H, alkyl, alkoxy, aryl, aryloxy, heteroaryl, heteroaryloxy, cross-linkable moiety, or oligo (ethylene glycol).

One aspect of the present invention includes the use of the semiconducting polymers and the conjugates described herein in devices such as a solar cell, an optical device, an electroluminescent device, a photovoltaic cell, semiconducting cell, or photodiode. Another aspect of the present invention includes the use of the conjugates of the semiconducting polymers described herein in a solar cell, an optical device, an electroluminescent device, a photovoltaic cell, semiconducting cell, photodiode or polymeric field effect transistors.

A method of incorporating the semiconducting polymers described herein onto a substrate comprises dissolving one or more of the polymers described herein with a fullerene derivative in a solvent and applying the resulting composite onto a substrate.

The one or more of the polymers can be co-dissolved with a fullerene derivative in 1,2-dichlorobenzene, chlorobenzene, 20 chloroform, or toluene, for example, in a 1:1 weight ratio, respectively. 1,8-diiodooctance can also be used in combination with above-mentioned solvents. Polymers involving LY-12, LY-14, or LY-18 (described below) can be in concentrations in a range of about 8 to about 12 mg/ml or any 25 combination or subcombination therein. They can also be about 10 mg/ml. Polymers involving LY-15, LY-16, or LY-17 (described below) can be in concentrations in a range of about 10 to about 15 mg/ml or any combination or subcombination therein. They can also be about 13 mg/ml. LY-15, LY-16, and 30 LY-17 can be dissolved in a combination of3% by volume of 1,8-diiodooctance and 1,2-dichlorobenzene. From about 2% to about 4% by volume of 1,8-diiodooctance can be mixed with 1,2-dichlorobenzene.

The polymer/fullerene composite can then be applied onto 35 a substrate. Suitable substrates such as Indium Tin Oxide (ITO)-coated glass that are known in the art can be used. The composite can be applied onto the substrate using any variety of processes suitable for layer application or casting. Spin casting can be used to apply a thin layer or film of the com-40 posite blend onto a substrate. The layer of composite blend can be from about 80 nm to about 150 nm thick. Layer thickness will vary depending on the application of the composite blend and the substrate. Therefore, provided herein is a photovoltaic film comprising a semiconducting polymer as 45 described herein with a fullerene derivative.

Semiconducting polymers described herein can have alternating thieno[3,4-b]thiophene and benzodithiophene units. The physical properties of these polymers can be finely tuned for photovoltaic application. The HOMO energy levels of the 50 polymer can be lowered by substituting alkoxy side chains to the less electron donating alkyl chains in LY-15 or introducing electron withdrawing fluorine into the polymer backbone in LY-16, leading to significant increase in Voe (28%) for polymer solar cells. The side chains and substitute groups 55 also affect polymer's absorption and hole mobility, as well as the miscibility with fulleride, all influencing polymer solar cell performances. Films prepared from mixed solvent exhibit finely distributed polymer/fulleride interpenetrating network and significantly enhanced solar cell conversion effi-60 ciency. A power conversion efficiency over 6% can be achieved in solar cells based on fluorinated LY-16/PC61BM composite films prepared from mixed solvents.

Throughout this specification, various indications have been given as to preferred and alternative embodiments of the 65 invention. However, it should be understood that the invention is not limited to any one of these. It is therefore intended that the foregoing detailed description be regarded as illustrative rather than limiting, and that it be understood that it is the appended claims, including all equivalents, that are intended to define the spirit and scope of this invention.

EXAMPLES

Monomer Synthesis

4,6-Dihydro-thieno[3,4-b]thiophene-2-carboxylic acid methyl ester (3): A solution of 4.8 g (20 mmol) (2) in 240 mL methanol was heated up and kept mild boiling. Sodium sulfide solution $(1.7 g, 22 mmol)$ in 160 mL methanol was added 5 dropwise in 2 h. After cooling of the reaction mixture, the precipitate was filtered off. The solvent was removed by evaporation and the residue was washed by water. Colunm chromatography on silica gel using dichloromethane yielded the pure title compound as a white solid 2.0 g (50%). ¹HNMR ¹⁰ (CDCl₃): δ 3.87 (3H, s), 4.05-4.06 (2H, t, J=3 Hz), 4.19-4.20 (2H, t, J=3 Hz), 7.48 (1H, s). ¹³CNMR (CDCl₃): 633.12, 33.47, 52.19, 127.90, 137.30, 143.90, 146.64, 162.51.

15 tion of2.0 g (10 mmol) (3) in 30 mL ethyl acetate was stirred ²⁰ the residue contained a crude product of (4) and 3-chloroben-Methyl thieno[3,4-b]thiophene-2-carboxylate (5): A soluand cooled down in dry ice bath. MCPBA (1.75 g, 10 mmol) in 30 mL ethyl acetate was added dropwise to the reaction solution. After the addition, the mixture was kept stirring for overnight. Then the solvent was removed by evaporation and zoic acid.

 $_{25}$ evaporation. The residue was purified by flash chromatogra-The residue was refluxed in acetic anhydride for 2.5 h. The mixture was cooled down and the solvent was removed by phy on silica gel to give crude product of (5) 1.62 g (82%) . ¹HNMR (CDCl₃): δ 3.90 (3H, s), 7.29-7.30 (1H, d, J=3 Hz), 7.60-7.61 (lH, d, J=3 Hz), 7.71 (lH, s).

4,6-dibromothieno[3,4-b]thiophene-2-carboxylic acid (7): NBS ³⁰ To a solution of 1.20 g (6.0 mmol) (5) in 15 mL DMF was
added dropwise a solution of NBS (2.70 g, 15.0 mmol) in 15 added dropwise a solution of NBS (2.70 g, 15.0 mmol) in 15 mL DMF under nitrogen protection at dark. The reaction mixture was heated up to 40° C. and kept stirring for 24 h. Then it was cooled down, poured to saturated sodium sulfite 35 solution at ice-water bath, and extracted with dichloromethane. The organic phase was collected and dried by sodium sulfate. Removal of the solvent, and colunm purifi-

> The free carboxylic acid (7) was obtained upon hydrolyzing the ester (6) by refluxing overnight with lithium hydroxide in THF/H₂O solution, followed by acidification in 1 N HCl solution. ¹HNMR ((CD₃)₂SO): δ 7.73 (1H, s), 13.46 (lH, s). MS (EI): Calcd, 339.8. found (M+l)+, 340.7.

> > B-1

4,5-Bis-chloromethyl-thiophene-2-carboxylic acid methyl ester (2): To a mixture of 4.0 g (28 mmol) of (1) and 11.0 g (136 mmol) of chloromethyl methyl ether was added dropwise 8.4 g (42 mmol) of $TiCl₄$ in an ice-water bath, and the mixture was stirred for 5 h at RT after the addition. 50 mL 60 $CH₂Cl₂$ was added, and the reaction mixture was poured to ice-water. The organic phase was washed with brine, dried and evaporated in vacuo. Recrystallization of the residue from hexane yield the title product as colorless needles 5 .5 g (82%). ¹HNMR (CDCl₃): δ 3.89 (3H, s), 4.60 (2H, s), 4.79 ϵ 5 $(2H, s)$, 7.72 (1H, s). ¹³CNMR $(CDC1₃)$: δ 37.07, 37.29, 52.44, 132.85, 134.83, 137.04, 161.85.

B-2

Dodecyl 4,6-dibromothieno[3,4-b]thiophene-2-carboxylate (8-1): 0.68 g (2.0 mmol) of 6, 0.50 g (2.4 mmol) DCC, 84 mg (0.69 mmol) DMAP were added to a 25 mL round bottom flask with 10 mL CH_2Cl_2 . 1.86 g (10.0 mmol) of ROH (dode- 15 can-1-ol) was added to the flask and then kept stirring for 20 h under N_2 protection. The reaction mixture was poured to 60 mL water and extracted with $CH₂Cl₂$. The organic phase was dried by sodium sulfate and the solvent was removed. Column chromatography on silica gel using hexane/CH₂Cl₂=4/1 $_{20}$ yielded the pure title compound as light orange solid 0.75 g (74%). ¹HNMR (CDCl₃): δ 0.86-0.89 (3H, t, J=7 Hz), 1.20-1.45 (18H, m), 1.74-1.77 (2H, m), 4.30-4.33 (2H, t, J=7 Hz), 7.54 (lH, s).

2'-Methyltetradecyl-2'4,6-dibromothieno[3,4-b] thiophene-2-carboxylate (8-2): 0.68 g (2.0 mmol) of 8, 0.50 g (2.4 mmol) DCC, 84 mg (0.69 mmol) DMAP were added to 25 a 10 mL round bottom flask with 5 mL CH₂Cl₂. 2.28 g (10.0) mmol) of 2-methyltetradecan-2-ol was added to the flask and then kept stirring for 20 h under N_2 protection. The reaction $_{30}$ mixture was poured to 30 mL water and extracted with $CH₂Cl₂$. The organic phase was dried by sodium sulfate and the solvent was removed. Column chromatography on silica gel using hexane/ $CH_2Cl_2=4/1$ yielded the pure title compound as light orange solid 0.63 g (72%). δ 0.86-0.89 (3H, t, 35 J=7 Hz), 1.24-1.40 (12H, m), 1.56 (6H, s), 1.83-1.87 (2H, m), 7.44 (lH, s).

2'-Ethylhexyl 4,6-dibromothieno [3,4-b]thiophene-2-carboxy late (8-3): 0.68 g (2.0 mmol) of 8, 0.50 g (2.4 mmol) DCC, 84 mg (0.69 mmol) DMAP were added to a 10 mL 40 round bottom flask with 5 mL CH₂Cl₂. 1.3 g (10.0 mmol) of 2-methyltetradecan-2-ol was added to the flask and then kept stirring for 20 h under N_2 protection. The reaction mixture was poured to 30 mL water and extracted with CH_2Cl_2 . The organic phase was dried by sodium sulfate and the solvent 45 was removed. Colunm chromatography on silica gel using hexane/ $CH_2Cl_2=4/1$ yielded the pure title compound as light orange solid 0.63 g (72%). δ 0.90-0.96 (6H, m, J), 1.32-1.46 (SH, m), 1.66-1.75 (lH, m), 4.21-4.27 (2H, m), 7.53 (lH, s).

Ml

M2

Synthesis of other electronegative thieno [3,4-b] thiophene monomers. To further lower the HOMO energy level in the polymers, we propose the synthesis of several monomers that are more electronegative based on ester substituted thieno [3,4-b]thiophene. M1 is replacing the carboxylic ester with a cyanide group, cyanide is more electronegative than an ester. M2 is to change the thiophene ring to a thiazole ring, a thaizole is more electronegative because of the nitrogen. The last two (M3, M4) are introducing a more electron withdrawing functional group in the proton position. On group would be fluorine and the other would be cyanide. Synthesis of these monomers is outlined below.

45 4,8-Dioctoxybenzo[l,2-b:4,5-b']dithiophene (9): 1.0 g (4.5 mmol) of benzo $[1,2$ -b:4,5-b']dithiophene-4,8-dione was mixed with 0.65 g (10 mmol) of zinc dust in a flask. Then, 4 ml of ethanol and 15 ml of 20% NaOH was added and the mixture was refluxed for 1 h. 4.3 mL octyl p-toluenesulfonate 50 was added in portions with stirring until the color changed to red. The resulting precipitate was filtered, the filtrate was diluted with 100 mL water and extracted with 100 mL chloroform. The organic extraction was dried with anhydrous sodium sulfate and evaporated in vacuo. Column chromatog-55 raphy on silica gel using dichloromethane and hexanes mixed solvents yielded the pure title compound as a white solid $(1.24 \text{ g}, 62\%)$. ¹HNMR (CDCl₃): $\delta 0.88$ -0.91 (6H, t, J=7 Hz), 1.31-1.38 (16H, m), 1.54-1.58 (4H, m), 1.84-1.89 (4H, m), 4.26-4.29 (4H, J=7 Hz), 7.36-7.37 (2H, d, J=6 Hz), 7.47-7.48 60 (2H, d, J=6 Hz).

(10): 0.62 g (1.4 mmol) of (9) was dissolved in 20 mL anhydrous THF and cooled in acetone/dry ice bath under nitrogen protection. 1.4 mL (3.5 mmol) butyllithium solution (2.5M in hexanes) was added dropwise with stirring, after the 65 addition the mixture was kept in dry ice bath for 30 mins and RT for 30 mins. The mixture was cooled into the dry ice bath and 4.2 mL (4.2 mmol) trimethyltin chloride solution (IM in hexanes) was added and kept stirring at RT for overnight. The mixture was quenched with 50 mL water and extracted with hexanes. The organic extraction was dried with anhydrous sodium sulfate and evaporated in vacuo. Recrystallization of the residue from isopropanol yields the title product as col- ⁵ orless needles $0.70 \text{ g} (65\%)$. ¹HNMR (CDCl₃): $\delta 0.44$ (18H, s), 0.88-0.91 (6H, t, J=7 Hz), 1.31-1.38 (16H, m), 1.54-1.58 (4H, m), 1.86-1.90 (4H, m), 4.28-4.31 (4H, J=7 Hz), 7.51 (2H, s).

12

14

45

 C_6F_{13}

 Rr

of2.3 g (5.0 mmol) (12) in 40 mL ethyl acetate was cooled in a dry ice bath, then 1.1 g (5.0 mmol) 3-Chloroperbenzoic acid in 20 mL ethyl acetate was added dropwise. After the addition, the mixture was kept stirring in dry ice bath for 2 h, then RT 6 h. Removal of solvent under vacuum produced a white solid. The obtained solid was refluxed in 20 mL acetic anhydride for 2 h. Then the mixture was cooled down and solvent was removed under vacuum. Column chromatography on ¹⁰ silica gel using hexanes as fluent yielded the title compound as colorless liquid 1.8 g (80%). ¹HNMR (CDCl₃): δ 7.33-7.36 (2H, m), 7.57-7.59 (lH, d J=2 Hz).

4,6-Dibromo 2-perflurohexyl-thieno[3,4-b]thiophene (14): To a solution of 2.75 g (6.0 mmol) (5) in 15 mL DMF was added dropwise a solution of NBS (2.70 g, 15.0 mmol) in 15 mL DMF under nitrogen protection at dark. The reaction mixture was heated up to 40° C. and kept stirring for 24 h. Then it was cooled down, poured to saturated sodium sulfite solution at ice-water bath, and extracted with dichlo-²⁰ romethane. The organic phase was collected and dried by sodium sulfate. Removal of the solvent, and colunm purification on silica gel using hexane as fluent yielded the target product (14) 3.32 g (90%). ¹HNMR (CDCl₃): δ 7.20 (1H, s) MS (EI): Calcd, 616.1. found $(M+1)^{+}$, 616.7.

2,3-Bis-chloromethyl-5-perflurohexyl-thiophene (11): To a mixture of 4.0 g (10 mmol) of 2-perflurohexyl thiophene and4.0 g (50mmol) of chloromethyl methyl ether was added dropwise 4.6 g (18 mmol) of $SnCl₄$ in an ice-water bath, and the mixture was stirred for 5 hat RT after the addition. 50 mL $CH₂Cl₂$ was added, and the reaction mixture was poured to ice-water. The organic phase was washed with brine, dried and evaporated in vacuo. Colunm chromatography on silica gel using hexanes yielded the titled compound as light yellow oil 4.4 g (88%). ¹HNMR (CDCl₃): δ 4.61 (2H, s), 4.80 (2H, s), 7.38 (lH, s). 50

2-Perflurohexyl-4,6-dihydro-thieno[3,4-b]thiophene (12): A solution of 5.0 g (10 mmol) (11) in 100 mL methanol was heated up and kept mild boiling. Sodium sulfide solution (2. 7 g, 11 mmol) in 80 mL methanol was added dropwise in 1 h. After cooling of the reaction mixture, the precipitate was filtered off. The solvent was removed by evaporation and the residue was washed by water. Colunm chromatography on silica gel using hexanes/dichloromethane (3/1) yielded the 1 HNMR (CDCl₃): δ 4.07-4.09 (2H, t, J=3 Hz), 4.21-4.23 (2H, m), 7.13 (lH, s). pure title compound as light yellow liquid 2.5 g (53%). 65

17

Oligomer (18): 331 mg (0.50 mmol) of monomer 14 was weighted into a 25 mL round-bottom flask. 2.13 g (1.20 mmol) Tributylstannyl octithiophene and 25 mg of Pd(PPh₃)₄ were added. The flask was subjected to 3 successive cycles of evacuation followed by refilling with argon. Then, 4 mL anhydrous DMF and 16 mL anhydrous toluene were added via syringe. The reaction was carried out at 120° C. for 24 h. The raw product (16) was collected by precipitating in methanol and the precipitate was dissolved in 20 mL THF and cooled in dry ice bath. 1.8 mL of IM TBAF solution was added dropwise and kept stirring in dry ice bath for 1 h then 25 RT for overnight. The raw product (17) was collected by precipitating in hexanes. The product was further purified by colunm chromatography on silica gel using hexanes/methylene chloride as fluent yielded the title compound 1.09 g (70%). ¹HNMR (CDCl₃): δ 0.86-0.91 (48H, m), 1.26-1.44 (96H, m), 1.63-1.71 (32H, m), 2.55-2.65 (4H, t, J=8 Hz), 2.73-2.82 (28H, m), 6.90 (lH, s), 6.96-7.00 (llH, m), 7.02- 7.03 (2H, d, J=4 Hz), 7.07 (lH, s), 7.11 (lH, s), 7.61 (lH, s). 30

To a solution of 0.93 g (0.30 mmol) oligomer (17) in 30 mL $_{35}$ THF was added dropwise a solution of NBS (0.133 g, 0.75 mmol) in 5 mL THF under nitrogen protection at dark. The reaction mixture was kept stirring for 24 h. The raw product (18) was collected by precipitating in hexanes. The product was further purified by column chromatography on silica gel $_{40}$ using hexanes/methylenechloride as fluent yielded the title compound 0.83 g (85%). ¹HNMR (CDCl₃): δ 0.90-0.91 (48H, m), 1.25-1.54 (96H, m), 1.63-1.72 (32H, m), 2.55-2.59 (4H, t, J=8 Hz), 2.73-2.80 (28H, m), 6.69 (2H, s), 6.82-7.11 (12H, m), 7.61 (lH, s). 45

18

-continued

2'-Methyltetradecyl-2'4,6-dibromothieno[3,4-b] thiophene-2-carboxylate (276 mg, 0.50 mmol) was weighted into a 25 mL round-bottom flask. 331 mg (0.50 mmol) 2,5- 10 Bis(tributylstannyl)thiophene and 25 mg of $Pd(PPh₃)₄$ were added. The flask was subjected to 3 successive cycles of evacuation followed by refilling with argon. Then, 2 mL anhydrous DMF and 8 mL anhydrous toluene were added via syringe. The polymerization was carried out at 120° C. for 24 h. The raw product was collected by precipitating in methanol and the precipitate was dissolved in chloroform, filtered with Celite to remove the metal catalyst. The final polymers were obtained by precipitating in hexanes, and drying in vacuum for 12 h, got solid 142 mg (60%). ¹HNMR (CDCI₃): δ 0.80-
20 \pm 0.80-2011 \pm 0.80-2011 \pm 0.80-2012 $\frac{1}{2}$ 1.00 (3H, br), 1.00-1.80 (26H, br, m) 1.82-2.10 (6H, br), 7.28-7.40 (2H, br), 7.58-7.70 (1H, br). GPC: Mw $(98\times10^{3}$ g/mol), PDI (1.76).

 $C_{12}H_{25}OOC$

LY-12: Monomer(8-2) (255 mg, 0.50 mmol) was weighted into a 25 mL round-bottom flask. Bis(trimethylstannyl)benzodithiophene (386 mg, 0.50 mmol) and $Pd(PPh₃)₄$ (25 mg) were added. The flask was subjected to 3 successive cycles of 65 vacuum followed by refilling with argon. Then, anhydrous DMF (2 mL) and anhydrous toluene (8 mL) were added via a syringe. The polymerization was carried out at 120° C. for 12 h under nitrogen protection. The raw product was collected by precipitating in methanol. The precipitate was dissolved in chloroform, filtered with Celite to remove the metal catalyst. The final polymers were obtained by precipitating in hexanes, and drying in vacuum for 12 h, get solid 338 mg (85%). 5 ¹HNMR (CDCl₂CDCl₂): δ 0.70-2.40 (53H, br), 3.82-4.83 (6H, br), 6.70-7.90 (3H, br). GPC: Mw (23×10³ g/mol), PDI (1.29).

LY-14: Polymer was synthesized by similar method as 25 LY-12. Monomer(8-3) (227 mg, 0.50 mmol) was weighted into a 25 mL round-bottom flask. Bis(trimethylstannyl)benzodithiophene (386 mg, 0.50 mmol) and $Pd(PPh₃)₄$ (25 mg) were added. The flask was subjected to 3 successive cycles of vacuum followed by refilling with argon. Then, anhydrous 30 polymerization with ditin monomers through a palladium DMF (2 mL) and anhydrous toluene (8 mL) were added via a synnge. The polymerization was carried out at 120° C. for 12 h under nitrogen protection. The raw product was collected by precipitating in methanol. The precipitate was dissolved in chloroform, filtered with Celite to remove the metal catalyst. 35 The final polymers were obtained by precipitating in hexanes, and drying in vacuum for 12 h, get solid 296 mg (80%). ¹HNMR (CDCl₂CDCl₂): δ 0.70-2.42 (45H, br), 3.90-4.80 (6H, br), 6.70-8.00 (3H, br).GPC: Mw (27×10³ g/mol), PDI $(1.67).$ 40

A device was made with the polymer LY-14 and an electron acceptor in a 1:1 mass ratio, 14 mg/mL polymer solution in chlorobenzene. The current density versus voltage is shown in FIG. 1. J_{sc} =17.3 mA/cm², V_{oc}=0.60 V, FF=0.53, $PCE=5.5\%$. 45

20

Synthesis of other electronegative thieno [3,4-b] thiophene polymers. The dibrominated monomers will then undergo catalyzed Stille polycondensation reaction.

Polymer was synthesized by a similar procedure described 60 above. Oligomer 18 (327 mg, 0.10 mmol) was weighted into a 25 mL round-bottom flask. 66 mg (0.10 mmol) 2,5-Bis (tributylstannyl)thiophene and 5 mg of $Pd(PPh₃)₄$ were added. The flask was subjected to 3 successive cycles of evacuation followed by refilling with argon. Then, 2 mL 65 anhydrous DMF and 8 mL anhydrous toluene were added via syringe. The polymerization was carried out at 120° C. for 24 h. The raw product was collected by precipitating in methanol and the precipitate was dissolved in chloroform, filtered with Celite to remove the metal catalyst. The final polymers were obtained by precipitating in hexanes, and drying in vacuum for 12 h, got solid 288 mg (90%). ¹HNMR (CDCl₃): δ 0.80-0.91 (48H, m), 1.20-1.56 (96H, m), 1.60-1.82 (32H, m), 2.68-2.85 (32H, m), 6.69 (2H, s), 6.90-7.13 (18H, m), 7.60 (1H, s). GPC: Mw (19×10^3 g/mol), PDI (1.37). Polymer Synthesis

LY-12: $X = H$, $R_1 = n$ -dodecyl, $R_2 = n$ -octyloxy LY-14: $X = H$, $R_1 = 2$ -ethylhexyl, $R_2 = n$ -octyloxy LY-15: $X = H$, $R_1 = 2$ -ethylhexyl, $R_2 = n$ -octyl LY-16: $X = F$, $R_1 = n$ -octyl, $R_2 = 2$ -ethylhexyloxy LY-17: $X = H$, $R_1 = n\text{-octyl}$, $R_2 = 2\text{-ethylhexyloxy}$ LY-18: $X = H$, $R_1 = 2$ -butyloctyl, $R_2 = n$ -octyloxy

 PC_{61} BM

LY-16: Octyl-6-dibromo-3-fluorothieno[3,4-b]thiophene-2-carboxylate (6) (236 mg, 0.50 mmol) was weighted into a $_{60}$ 25 mL round-bottom flask. 2,6-Bistrimethyltin-4,8-Di(2 ethyl)hexyloxybenzo[l,2-b:4,5-b']dithiophene (9) (386 mg, 0.50 mmol) and $Pd(PPh_3)_4$ (25 mg) were added. The flask was subjected to 3 successive cycles of vacuum followed by refilling with argon. Then, anhydrous DMF (2 mL) and anhydrous 65 toluene (8 mL) were added via a syringe. The polymerization was carried out at 120° C. for 12 h under nitrogen protection.

The raw product was precipitated into methanol and collected by filtration. The precipitate was dissolved in chloroform, filtered with Celite to remove the metal catalyst. The final polymers were obtained by precipitating inhexanes, and drying in vacuum for 12 h, yielding LY-16 (309 mg, 82%). ¹HNMR (CDC1₂CDC1₂): δ 0.80-2.40 (45H, br), 3.90-4.70 $(6H, br), 7.00-7.90 (2H, br), GPC: Mw (19.3×10³ g/mol), PDI$ (1.32).

LY-14, LY-15, LY-17 and LY-18 are synthesized according to the same procedure as LY-16 with respective monomers. The ¹ HNMR and GPC data of the polymers are listed below:

LY-14: ¹HNMR (CDC1₂CDC1₂): δ 0.70-2.42 (45H, br), 3.90-4.80 (6H, br), 6.70-8.00 (3H, br). GPC: Mw (23.2xl0³ 15 g/mol), PDI (1.38).

LY-15: ¹HNMR (CDC1₂CDC1₂): δ 0.70-2.35 (45H, br), 2.90-3.40 (4H, br), 4.20-4.70 (2H, br), 6.70-8.20 (3H, br). GPC: Mw (23.7x103 g/mol), PDI (1.49).

20 LY-17: ¹ HNMR (CDC $\{zCDC\}$ z): δ 0.90-2.40 (45H, br), 3.90-4.70 (6H, br), 7.00-7.60 (2H, br), 7.60-8.10 (lH, br). GPC: Mw $(22.7 \times 10^3 \text{ g/mol})$, PDI (1.41).

LY-18: ¹HNMR (CDC1₂CDC1₂): δ 0.70-2.42 (53H, br), 3.90-4.80 (6H, br), 6.70-8.00 (3H, br). GPC: Mw (25.0×10³ 25 g/mol , PDI (1.50).

The polymerization was carried out via the Stille polycondensation reaction. The corresponding monomers were synthesized according to the following routes.

6

5

 $EH = 2$ -ethylhexyl

To shorten the dodecyl ester chain in LY-12, a n-octyl side 15 chain substituted polymer was synthesized. Soluble LY-14 was synthesized with shortened and branched side chains. For comparison, a more bulky branched side chain, 2-butyloctyl was used in LY-18. The branched side chain can also be $_{20}$ grafted to the benzodithiophene, which lead to LY-17 with two 2-ethylhexyloxy side chain attached to the benzodithiophene ring. The alkoxy groups grafted on benzodithiophene ring are strong electron-donating groups that can raise the HOMO energy level of the polymer. This will 25 lead to the reduction in Voe, detrimental to the performance of SnMe_3 polymer solar cells. In order to further adjust the polymer's electronic properties, LY-15 with less electron-donating alkyl chains in benzodithiophene was synthesized.

To further lower the HOMO level, a second electron with-30 drawing group can be introduced to the 3 position of the thieno[3,4-b]thiophene ring. Fluorine is a good candidate to functionalize the 3 position because the fluorine has a high electronegativity. The size of the fluorine atom is small, which will introduce only small steric hindrance for the configura-35 tion and packing of the polymer. The fluorinated thieno[3,4 b]thiophene was synthesized via a modified route previously reported for ester substituted thieno[3,4-b]thiophene (above scheme). The fluorine was introduced to the fused ring unit from 4,6-dihydrothieno[3,4-b]thiophene-2-carboxylic acid 40 after deprotonation by using BuLi and reacting with PhS02NF. The fluorinated acid was first converted to ester and then dibromosubstituted thieno[3,4-b]thiophene. Initially, we attempted to introduce fluorine atom to LY-12 $(R_1 = n$ -dodecyl, $R_2 = n$ -octyloxy). The obtained polymer 45 exhibited poor solubility and only dissolves in dichlorobenzene over 100° C., which makes it difficult to prepare uniform films. To increase the solubility, benzodithiophene substituted with branch side chains was used and the fluorinated polymer, LY-16, was obtained.

50 The structures of polymers were characterized with ¹ HNMR spectroscopy, all consistent with the proposed ones. Gel permeation chromatography (GPC) studies showed that these polymers have similar weight-averaged molecular weights between 19.3-25.0 kg/mol with a relatively narrow 55 polydispersity index (PDI) between 1.25 and 1.50. The results indicate that the changes in monomer structures did not lead to significant changes in polymerization reaction. These polymers have good solubility in chlorinated solvents, such as chloroform and chlorobenzene. Thermogravimetric 60 analyses (TGA) indicate that the polymers are stable up to about 200° C.

Characterization
¹ HNMR spectra were recorded at 400 or 500 MHz on Bruker DRX-400 or DRX-500 spectrometers, respectively. 65 Molecular weights and distributions of polymers were determined by using gel permeation chromatography (GPC) with a Waters Associates liquid chromatograph equipped with a 5

Waters 510 HPLC pump, a Waters 410 differential refractometer, and a Waters 486 tunable absorbance detector. THF was used as the eluent and polystyrene as the standard. The optical absorption spectra were taken by a Hewlett-Packard 8453 UV-Vis spectrometer.

Cyclic voltammetry (CV) was used to study the electrochemical properties of the polymers. For calibration, the redox potential of ferrocene/ferrocenium (Fc/Fc^+) was measured in the same condition, and it's located at 0.09 V to the Ag/Ag^+ electrode. It is assumed that redox potential of Fc/Fc^+ 10 has an absolute energy level of -4.80 eV to vacuum. The energy levels ofHOMO and the lowest unoccupied molecular orbital (LUMO) were then calculated according to the following equations:

$$
E_{HOMO} = -(\phi_{ox} + 4.71)
$$
 (eV)

$$
E_{LUMO} = -(\phi_{red} + 4.71)(\text{eV})
$$

where ϕ_{ox} is the onset oxidation potential vs Ag/Ag⁺ and ϕ_{red} is the onset reduction potential vs $A\mathcal{Q}/A\mathcal{Q}^+$.

Hole mobility was measured according to a similar method described in literature, using a diode configuration of ITO/ poly(ethylenedioxythiophene) doped with poly(styrenesulfonate) (PEDOT:PSS)/Polymer/Al by taking current-voltage current in the range of 0-6 V and fitting the results to a 25 space charge limited form, the SCLC is described by:

 $J=9\in_{o}\in_{n} u V^{2}/8L^{3}$

where \in_{\circ} is the permittivity of free space, \in_{\circ} is the dielectric constant of the polymer, μ is the hole mobility, V is the voltage 30 drop across the device, and L is the polymer thickness, $V=V_{app1}-V_r-V_{bi}, V_{app1}$ is the applied voltage to the device, V_r is the voltage drop due to contact resistance and series resistance across the electrodes, V *bi* is the built-in voltage due to the difference in work function of the two electrodes. The resistance of the device was measured using a blank configuration ITO/PEDOT:PSS/Al and was found to be about 10-20 Ω . The V_{bi} was deduced from the best fit of the $J^{0.5}$ versus V_{appl} plot at voltages above 2.5 V and is found to be about 1.5 \overline{V} . The dielectric constant, \in , is assumed to be 3 in 40. our analysis, which is a typical value for conjugated polymers. The thickness of the polymer films is measured by usingAFM.

Device Fabrication

The polymers LY-12, LY-14, LY-15, LY-16, LY-17, and 45 LY-18 were co-dissolved with $PC_{61}BM$ in 1,2-dichlorobenzene (DCB) in the weight ratio of 1:1, respectively. LY-12, LY-14, and LY-15 concentrations were 10 mg/ml while LY-15, LY-16, and LY-17 concentrations were 13 mg/ml. For the last three polymer solutions, mixed solvent effect was 50 studied with about 3% (volume) 1,8-diiodooctance to further improve the final device performance.

Indium Tin Oxide (ITO)-coated glass substrates $(15! 1/sq)$ were cleaned stepwise in detergent, water, acetone and isopropyl alcohol under ultrasonication for 15 minutes each and 55 subsequently dried in an oven for 5 hours. A thin layer $(\sim 30$ nm) of PEDOT:PSS (Baytron PVP Al 4083) was spin-coated onto ITO surface which was pre-treated by ultraviolet ozone for 15 min. Low conductivity PEDOT:PSS was chosen to minimize measurement error from device area due to lateral conductivity ofPEDOT:PSS. After being baked at 12° C. for \sim 20 min, the substrates were transferred into a nitrogen filled glove box(<0.1 ppm O2 & H₂0). A polymer/PCBM composites layer (ca.100 nm thick) was then spin-cast from the blend solutions at 1000 rpm on the ITO/PEDOT:PSS substrate 65 Hole Mobility without further special treatments. Then the film was transferred into thermal evaporator which is located in the same

glove box. A Ca layer (25 nm) and an Al layer (80 nm) were deposited in sequence under the vacuum of 2×10^{-6} Tor. The effective area of film was measured to be 0.095 cm². Current-Voltage Measurement

The fabricated device was encapsulated in nitrogen filled glove box by UV epoxy (bought from Epoxy Technology) and cover glass. The current density-voltage (J-V) curves were measured using Keithley 2400 source-measure unit. The photocurrent was measured under AM 1.50 illumination at 100 mW/cm² under Newport Thermal Oriel 91192 1000W solar simulator $(4"$: $.4"$ beam size). The light intensity was determined by a mono-silicon detector (with KG-5 visible color filter) calibrated by National Renewable Energy Laboratory (NREL) to minimize spectral mismatch.

¹⁵ External Quantum Efficiencies (EQEs) were measured in UCLA by using lock-in amplifier (SR830, Stanford Research Systems) with current preamplifier (SR570, Stanford Research Systems) under short circuit condition. The devices were illuminated by monochromatic light from a xenon lamp passing through a monochromator (SpectraPro-2150i, Acton Research Corporation) with a typical intensity of $10 \mu W$. The monochromic incident beam is chopped with a mechanical chopper prior to reaching the device connected to lock-in amplifier and then focused on the testing pixel of the device. The photocurrent signal is then amplified by SR570 and detected with SR830. A calibrated mono silicon diode with known spectral response is used as a reference. Conductive atomic force microscopy measurement

All CAFM measurements were done under ambient conditions using a commercial scanning probe microscope (Asylum Research, MFP-3D). Platinum-coated, contact-mode AFM cantilevers with spring constant of 0.2 N/m and tip radius of ca. 25 nm (Budget Sensors) were used to map out the hole-current of films in the dark using contact mode. The deflection set point are 0.3 V and bias voltage are -2 V for all the sample measurements, and the conditions used to prepare the films are same to make the solar cell device. Electrochemical and Optical Properties

The HOMO and LUMO energy level of the polymers were determined by cyclic voltammetry (CV) and the results were summarized in FIG. 2. The HOMO energy levels of the polymers are very close except for LY-15 and LY-16. From the comparison ofLY-14 and LY-15, it was noticed that the substitution of octyloxy side chain to octyl side chain lowered the HOMO energy level of the polymer from -4.94 eV to -5.04 eV. Comparing LY-16 and LY-17, polymers with same side chain patterns, it is clear that the introduction of the electron withdrawing fluorine in polymer backbone significantly lowered the HOMO level. Characteristics of the polymer absorption are summarized in the table below. All these polymers show very similar absorption spectra, the changes of the absorption peak and onset point among the polymers are within 25 nm.

The hole mobility of the polymers is measured according to method based on space charge limited current (SCLC) model, and the results are plotted in FIG. **3.** The hole mobility of 4.7×10^{-4} cm₂/V·s, 4.0×10^{-4} cm₂/V·s, 7.1×10^{-4} cm₂/V·s, 7.7×10^{-4} cm₂/V·, 4.0×10^{-4} cm₂/V·s, 2.6×10^{-4} cm₂/V·s are found for LY-12, LY-14, LY-15, LY-16, LY-17 and LY-18 respectively. A small decrease of the polymer hole mobility is 5 observed after the introduction of bulky branched side chains to the polymer backbones. It is expected that the bulky side chains may increase the steric hindrance for intermolecular packing, so the hole mobility decreases. This explains that the largest decrease of the hole mobility happens to LY-18, which 10 has the most bulky 2-butyloctyl side chain on the ester group. It is interesting to note that the alkyl grafted LY-15 has higher mobility than the alkoxy grafted LY-14, though they both have similar side chain patterns. LY-16 has the largest hole mobility of $7.7 \times 10-4$ cm2/V·s among these polymers. It has 15 been reported that there is a strong n-stacking among interaction between the electron-deficient fluorinated aromatic rings and the electron-rich non-fluorinated ones in the fluorine-substituted aromatic moieties. The increase of mobility in fluorinated LY-16 is probably due to the increase in inter- 20 molecular packing between fluorinated backbone. Detailed studies by using grazing angle x-ray diffraction are in progress to elucidate polymer structures and will be presented Photovoltaic Properties

Photovoltaic properties of the polymers were investigated in solar cell structures of ITO/PEDOT:PSS/polymer: $PC_{61}BM(1:1, %$ wt ratio)/Ca/Al. The polymer active layers with nanoscale domain structure observed from TEM in solar were spin-coated from a dichlorobenzene solution. Representative characteristics of the solar cells are summarized in the 30 table below. Generally, the bulky side chain grafted polymers show larger V_{oc} than LY-12, as they have lower HOMO energy levels. The alkyl substituted LY-15 has an enhanced V_{oc} compared to LY-14, which is expected from the HOMO energy level difference. The fluorinated polymer LY-16 35 devices showed a larger Voe than LY-17. However, except for LY-14 and LY-15, the other polymer solar cells suffer obvious decrease in short circuit current (Js) and fill factor (FF) compared to the LY-12 solar cell. Further studies by using transmission electron microscopy (TEM) indicated that the poor 40 solar cell performances in LY-15 to LY-18 are related to the non-optimized morphology, which has a large effect on the BHJ polymer solar cell performance. The TEM images of polymer/PC₆₁BM blend films are shown in FIGS. 4a-4i: LY-12 (a); LY-14 (b); LY-15 (c); LY-16 (d); LY-17 (e); LY-18 45 (f). Polymer/PC $_{61}$ BM blend films prepared from mixed solvents dichlorobenzene/diiodooctance (97/3, v/v): LY-15 (g); LY-16 (h); LY-17 (i).

The TEM images of LY-14/PC $_{61}$ BM blend film shows finer features comparable to LY-12 one, which may be due to the 50 increase in the miscibility of the polymer with PC_{61} BM after shortening the dodecyl side chain into 2-ethylhexyl side chain. As a result, Jsc and FF in LY-14 solar cell are slightly larger than LY-12 one. However, large domains are observed in their PC_{61} BM blend films of LY-17 or LY-18. The bulky 55 side chains reduce the miscibility of polymer with PC_{61} BM, leading to better phase separation between polymer chains and PC_{61} BM molecules. As a result, the interfacial areas of charge separation in LY-17 or LY-18 are reduced and the polymer solar cell performances diminished. It is not coinci- ⁶⁰ dent that the LY-18 has the largest feature sizes (150-200 nm) in TEM image and its solar cell performance is the worst. With the same side chain patterns as LY-17, the fluorinated LY-16 also suffer the non-optimized morphology, as shown the large features (over 100 nm) in the TEM image of LY-16/ PC_{61} BM blend film. Although LY-16 shows the lowest HOMO energy level and the largest hole mobility, its photo-

voltaic performance in simple polymer/ PC_{61} BM solar cells is modest (3.10%). Comparing LY-14 and LY-15 with similar side chain patterns, LY-15 has a larger Jsc than LY-14, which is due to the increase of hole mobility in LY-15. However, the better packing ability in LY-15 may reduce its miscibility with PC_{61} BM, and there are a few large features (about 50 nm) in the TEM images of LY-15/PC $_{61}$ BM blend film. Due to the non-optimized morphology, LY-15 solar cell suffers a slight decrease on FF compared to LY-14. But with the increase in Jsc and Voe, LY-15 devices still show 5.53% PCE.

Insightful information on the effect of morphology on in the future publication. $_{25}$ charge transport behaviors of composite films was obtained from studies using conductive atomic force microscopy (CAFM). The two-dimensional current maps correlate well cell films. We select LY-14 and LY-18 systems as example to illustrate this point. FIGS. **Sa** and **Sb** show the CAFM current images at fixed bias voltage of $-2V$ for films LY-14/PC₆₁BM and LY-18/PC $_{61}$ BM. Consistent with TEM study, we observe donor/acceptor interfaces throughout the films with interpenetrating networks of donor/acceptor material in the LY-14 film. The high hole current features (dark areas) are contributions from polymer rich domains and are distributed uniformly over whole LY-14/PC $_{61}$ BM film surface, which makes the solar cells high current density. In contrast, there are low current areas in the images and largely phase separated features throughout the LY-18/PC $_{61}$ BM films. The much lower current (see the difference in scale bar) of LY-18 reflects the lower hole mobility of LY-18 than LY-14.

> This morphological problem can be remedied by using mixed solvents in preparing polymer/fullerides spin-coating solution. Primary study showed that the LY-15/PC $_{61}$ BM blend film exhibited improved morphology by using dichlorobenzene/1,8-diiooctane (97/3, v/v) as solvent: there are no large features and it shows similar morphology as LY-12 or LY-14 blend film in TEM image. (FIG. 4g) The PCE of such polymer solar cell reaches to 5.85%. Similar morphology change for LY-16/PC $_{61}$ BM and LY-17/PC $_{61}$ BM blend films was observed (FIGS. *4h* and *4i).* Dramatic enhancement in solar cell performance can be observed both in LY-16 and LY-17 solar cells. (FIG. *6a)* Besides the increase of the Voe, LY-16 solar cell shows larger Jsc than LY-17 one, possibly due to the higher hole mobility in fluorinated LY-16. It can also explain the slight increase of the FF in the LY-16 solar cell. Therefore, we have successfully improved the Voe from $0.58V$ to $0.74V$ (\sim 28%) over LY-12 polymer system without scarification in photocurrent. The PCE from the LY-16/ PC₆₁ BM solar cell reached to 5.9%. FIG. 6b shows the EQE spectrum of LY-16/PC $_{61}$ BM solar cell prepared from mixed solvent. It can very efficiently harvest the light in the maximum photon flux region (680 nm), showing over 50% from 550 nm to 750 nm. A spectral mismatch factor (M) of 0.965 can be calculated by inserting AM1.5G standard spectrum, Oriel solar simulator (with 1.5G filter) spectrum and EQE

10

5

15

30

45

50

data. The efficiency of LY-16/PC $_{61}$ BM solar cell thus reaches to 6.1%, which is the highest value so far for single layer polymer solar cells.

The invention claimed is:

1. A semiconducting polymer of the formula (I)

- where X^1 and X^2 are independently O, S, N, NH, CR¹, CR², 20 CHR¹, or CHR², where R¹ and R² are independently H, alkyl, alkoxy, aryl, aryloxy, heteroaryl, heteroaryloxy, oligo(ethylene glycol);
- $Y¹$ and $Y²$ are independently O, Se, or amino;
- **Z** is an ester, amide, cyano, alkyl, or polyfluoroalkyl, polychloroalkyl, aryl, or heteroaryl;
- W is H, halogen, cyano, dicyanovinyl, or tricyanovinyl; and
- n is an integer greater than 0.

2. The semiconducting polymer of claim **1**, where $X¹$ and X^2 are independently O, S, NH, CHR¹, or CHR².

3. The semiconducting polymer of claim **2,** where Wis For H.

4. The semiconducting polymer of claim 2 , where $X¹$ and X^2 are independently CH₂ or O. 35

5. The semiconducting polymer of claim **2,** where Z is O_nF_{2n+1} or C_nH_{2n+1} , and n is an integer from 1 to 12.

6. The semiconducting polymer of claim **2,** where Z is ⁴⁰ -COOR", where R" is alkyl, alkoxy, aryl, aryloxy, heteroaryl, or heteroaryloxy.

7. The semiconducting polymer of claim **6,** where R" is

8. The semiconducting polymer of claim **2** where X^1 and X^2 are CH₂, Z is COOR", WisH, R^1 and R^2 are C_7H_{15} , and

R" has the formula

9. The semiconducting polymer of claim **2** where X^1 and X^2 are O, Z is COOR", W is F or H, $R¹$ and $R²$ have the following formula:

30

and R" is C_8H_{17} .

10. A conjugate of a semiconducting polymer according to claim **1.**

- **11.** A conjugate of a semiconducting polymer according to claim 2 and an acceptor material.
- **12.** The conjugate of semiconducting polymer of claim **1** where the acceptor material is a fullerene derivative.
- **13.** The conjugate of a semiconducting polymer of claim **2** where the fullerene derivative is C_{61} or C_{71} .
- **14.** A conjugate of the semiconducting polymer according to claim **1,** wherein Z is of the formula:

and R' is $-(CH_2)_m$ and m is an integer between 2 and 16; X is selected from O, S, NH, and $-\text{ONH}$.

15. The conjugate of claim **14**, where X^1 and X^2 are independently O, \tilde{S} , $\tilde{N}H$, CHR¹, or CHR².

16. The semiconducting polymer of claim **2** in a solar cell, an optical device, an electroluminescent device, a photovol-

taic cell, semiconducting cell, or photodiode. 55 **17.** The conjugate of claim **1** in a solar cell, an optical device, an electroluminescent device, a photovoltaic cell, semiconducting cell, photodiode or polymeric field effect transistors.

18. A photovoltaic layer, comprising a semi-conducting 60 polymer according to claim **2.**

19. The semiconducting polymer of claim **6,** where Z is C_mH_{2m+1} and m is an integer from 1 to 30.

* * * * *

(III)