

GRAPHICAL ABSTRACT

Nanoscale Functionalization of Surfaces by Graft-Through Sonogashira Polymerization

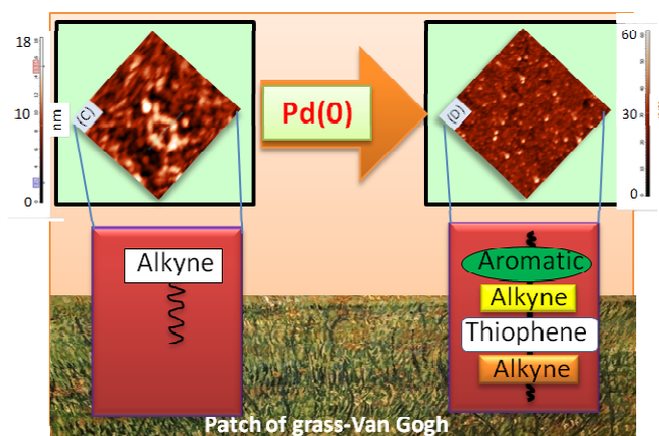
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“Graft-through” Sonogashira polymerization has been performed on functionalized self-assembled monolayer.



Nanoscale Functionalization of Surfaces by Graft-Through Sonogashira Polymerization†

P. Murugan,^a Soundaran Jeevarathinam Ananthakrishnan,^a Narayanasastri Somanathan,^{a,c} Debasis Samanta,^{*a} and Asit Baran Mandal^{*b,c}

Graft through Sonogashira polymerization was used to functionalize various surfaces with conjugate polymers in a dimension of less than 100 nm. Atomic force microscopy measurement revealed a dense surface coverage with several closely packed islands. UV-vis spectroscopy and cyclic voltametry measurements suggested a moderate band gap, which is important for various applications in material science. A device was fabricated using polymer functionalized ITO and deposited aluminium as cathode to determine the current-voltage (I-V) characteristics and charge carrier mobility. Space charge limited current method indicated moderate charge carrier mobility while I-V characteristic data indicated its behaviour as semiconducting material.

Functionalization of surfaces in nanoscale is important for its applications ranging from material science^{1,2} to chemical biology.^{3,4} In this context, utilization of self-assembled monolayer⁵ (SAM) formation technique affords a good control and defined orientation of functional groups—a prerequisite for the improved performance of devices and sensors.⁶ Since a desired, reactive ‘head-group’ may not always be compatible with the functional group necessary for SAM formation on a given surface, post-assembly chemical modification is often unavoidable.^{7,8} The chemical method⁹ to be employed must ensure a quantitative functional group conversion while the reaction condition must be mild enough to retain the structural integrity of the SAM¹⁰ and its bonding to the surface.¹¹ Reactivity issues of different functional groups on self-assembled monolayer and applications have been highlighted in a recent review.¹² Very recently we reported graft-through click polymerization to decorate the surface monolayer with polytriazole.¹³ However, due to the “broken” conjugation because of the presence of triazole¹⁴ moiety between thiophene rings result in higher band gap than the usual requirement for solar cell or molecular electronic device fabrication. Hence we aimed to devise new strategy to immobilize molecules of interest with continuous conjugation using graft-through approach.^{15,16} We were specifically interested in thiophene-based polymers because of its widespread use for various applications.^{17,18}

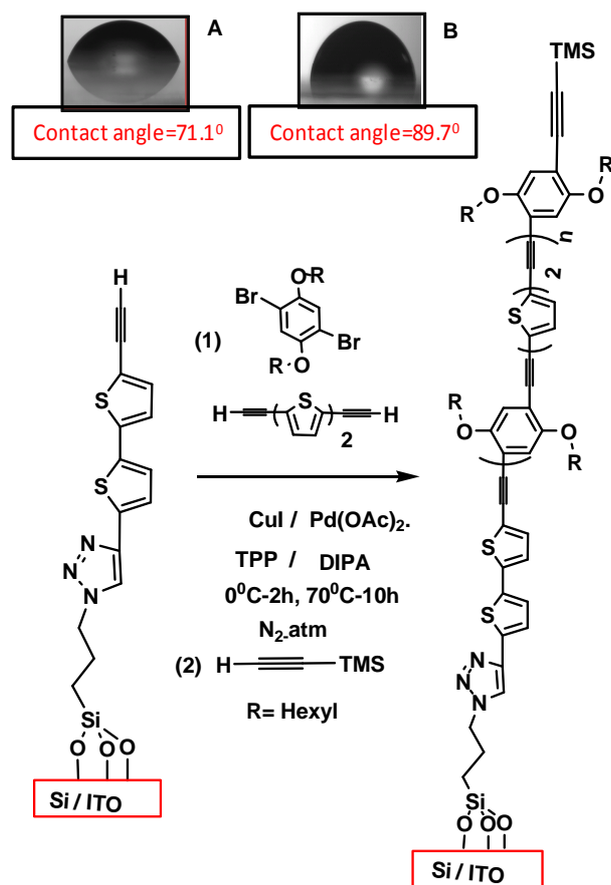
It was recognized that various coupling reactions such as Suzuki coupling, Heck coupling, Sonogashira coupling and Negeshi coupling reactions produce thiophene-based conjugated molecules and polymers.¹⁹ Among them, Sonogashira coupling reactions yield carbon-carbon triple bond between aromatic rings, which provide several unique features.²⁰ For example, the presence of internal triple bonds in the structure results in rigid structure and extended conjugation along with intermolecular cores. Moreover, polyacetylenes have high Q band absorption and higher power conversion efficiency than conjugated polymers with C=C linkage.^{21,22} Recently several researchers described the importance of sonogashira cross coupling reaction for different applications.²³⁻²⁵ Solution processed conjugated sonogashira polymers were found to give wide range of absorption and unique structural features.²⁶

In this paper, we report that “graft-through” Sonogashira polymerization can be used to form a conjugated thiophene-based polymers linked by C-C triple bond. Here, we used amenable starting materials like 2-ethynyl-5-(5-ethynylthiophen-2-yl)thiophene²⁷ and 1,4-dibromo-2,5-bis(hexyloxy)benzene monomers.²⁸ This strategy is superior to our previously reported click polymerization, since this Sonogashira polymerization on SAM yielded polymers with continuous conjugation, resulting in reduced band gap.

Scheme 1 depicts the polymerization reaction on an end alkyne functionalized silicon surface using 2-ethynyl-5-(5-ethynylthiophen-2-yl)thiophene and 1,4-dibromo-2,5-bis(hexyloxy)benzene followed by a quenching with ethynyl trimethylsilane.‡ Polymerization reactions were first studied and standardized on Self-assembled monolayer (SAM) on Si surface using FT-IR technique and extended the above approach to SAM on ITO surface.

Azide and alkyne-functionalized self-assembled monolayer on different surfaces were prepared according to literature procedure.²⁹ In brief, the plasma cleaned silicon wafer was treated with 3-azidopropyltrimethoxysilane in anhydrous toluene at 90 °C for 2 hours to form azide functionalized SAM and further click reaction was carried out with 2-ethynyl-5-(5-ethynylthiophen-2-yl)thiophene monomers and CuI/diisopropyl amine/DMF as a catalyst. The untreated and physisorbed monomers and catalysts were removed from the surface by repeated ultrasonication in fresh toluene, CHCl₃, H₂O, acetone and methanol. The contact angle measurement revealed the hydrophobic nature of the end alkyne-functionalized SAM of the surface (contact angle = 71.10°).

The polymerization was performed on the end alkyne functionalized silicon surface using 2-ethynyl-5-(5-ethynylthiophen-2-yl)thiophene and 1,4-dibromo-2,5-bis(hexyloxy)benzene in the presence of Pd(0) catalyst system. We added ethynyltrimethylsilane for quenching the reaction. During the surface polymerization, an appreciable amount of polymers also formed in solution phase. This type of competitive polymerization in solution is common for graft-through approach of polymerization on surfaces.¹⁶ After completion of surface polymerization the unreacted and physisorbed monomers, solution polymers, and catalyst were removed from the surface by repeated ultrasonication in fresh toluene, CHCl₃, H₂O, acetone, and ethanol and dried under N₂ gas. The polymer functionalized surface was characterized by ellipsometry, contact angle, FT-IR, X-ray photoelectron spectroscopy, atomic force microscopy and current-voltage (I-V) characterization. Increase in hydrophobicity after polymerization was observed by contact angle measurement. An appreciable amount of change from 71.1° to 89.7° was clearly observed. This may be attributed to the presence of hydrophobic silicon moiety at the termini of the polymer.³⁰



Scheme 1 ‘‘Sonogashira’’ polymerization reaction on a functionalized silicon surface using a ‘‘graft-through’’ approach. Inset: picture of water droplets on surfaces to measure contact angle: left side: end alkyne functionalized surface, right side: quenched polymer functionalized surface

Fig. 2. (A) shows the IR spectra of the silicon surface after the azide functionalization (magenta curve) and polymerization (red curve), respectively. In IR spectra, complete disappearance of peak at 2100 cm^{-1} was observed indicating the consumption of most of the azide groups.^{31, 32} Appearance of strong peaks at $2850\text{--}3000\text{ cm}^{-1}$ due to the C–H stretching frequency further confirms the successful implementation of ‘‘graft-through’’ Sonogashira polymerization on the surface.

Fig. 2(B) shows the XPS spectra of alkyne functionalized surface (black curve) and polymer functionalized ITO surface (red curve). The peak for S (2p), C (1s), N (1s) and O (1s) was observed at 165 eV, 285 eV, 400 eV and 532.5 eV respectively.³³ Peak for S (2s) appeared as weak signal at 228 eV. However, a clear attenuation of peak for N (1s) and improvement of peak area for S (2p) was observed after polymerization. This may be attributed to the incorporation of more amounts of sulphur and other elements via polymerization. Peak for Si (2s) and (2p) was observed at 152.5 eV and 102 eV respectively. Those peaks may be attributed to both binding silicon on surface before and after reaction as well as silicon from end trimethoxy silane after reaction. Appearance of peaks at 445 eV and 452 eV for In (3d) elements after reaction for the ITO surface confirmed the low value of thickness of the grafted polymer. Attenuation of those peaks after polymerization is indicative of anchoring of several groups after polymerization. Fig. S6, ESI depicts the XPS spectra of alkyne functionalized silicon surface (black curve) and polymer functionalized silicon surface respectively. In those cases peaks were observed at 164 eV, 285 eV, 400 eV and 531 eV respectively for S (2p), C (1s), N (1s) and O (1s).

Fig. 2(C) and 2(D) show the atomic force microscope (AFM) pictures of the end alkyne functionalized silicon surface and the silicon surface after polymerization reaction respectively. In this case, the height of the monolayers is more than calculated value. The higher values were observed by other authors also for siloxane terminated monolayer and was attributed to several uncontrolled aggregation of siloxanes in solution.³⁴ After polymerization, a dense pack of islands on surfaces were observed throughout the surfaces indicating a longitudinal grow of polymers typical for ‘‘graft-from’’ or ‘‘graft-through’’ polymerization techniques.³⁵ This differs from a typical ‘‘graft-to’’ methods of immobilization of polymers where polymers collapse to coiled structures.³⁶ Moreover, the resulting surface after polymerization had higher thickness than that of end alkyne monolayer and it showed approximately uniform coverage of an underlying substrate. However, thickness and roughness value increased with increase of polymerization reaction time. For example, as observed in Fig. S4, ESI the AFM image of polymer grafted silicon surface after performing the polymerization reaction for 24h clearly showed a rougher surface with bigger islands of polymers and length scale of heights. In this case, a high value of root mean square roughness of 33.4 nm with an amount of sampling of 25872 was observed. Similar effect of influence of time over the distribution of polymer was observed by others also.²⁹ Fig. S5, ESI showed the distribution profiles of grafted polymer across the polymer functionalized silicon surface when the polymerization reaction was performed for 24h. The

curves further showed the rough surface with uniform nature of surface topography. The large value of surface height indicates a high value of chain length of the polymers. Further, even a more roughness and lack of uniformity was observed when reaction was performed on ITO surface. This may be attributed to the inherent rough nature of pristine ITO surface compared to silicon surface. Similar observation of roughness was observed for “click” polymerization on ITO surface also.¹³ However, in this case, an improvement of surface coverage was clearly observed by performing the polymerization reaction for longer time of 24 h (Fig. S1, B, C).

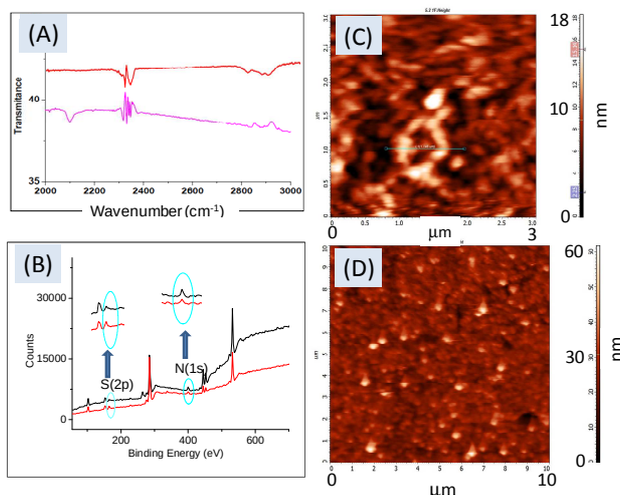


Fig. 2 (A) IR spectra of azide-functionalized (magenta) and polymer functionalized surfaces (red curve) (B) XPS spectra of alkyne-functionalized (black curve) and polymer functionalized (red curve) ITO surfaces. (C) and (D) AFM pictures of alkyne functionalized and polymer grafted (polymerization reaction is 12h) silicon surface respectively

The success of polymerization was also evident from the characterization of the residual polymers in solution by NMR spectroscopy (Fig. 3, Fig. S2, ESI). In this case, the disappearance of end alkyne proton peaks in ¹H NMR (3.41 ppm) and the shift of thiophene protons to 7.70 ppm was observed. Further the peak at 82.6 ppm corresponding to ¹³C alkyne carbon of the reactant disappeared to generate two signals at 87.9 and 85.9 ppm.³⁷ This may be attributed to the unsymmetrical nature of alkynes in the resulting polymer. As a control experiment, polymerization was attempted at cleaned empty surface (not functionalized), in a similar condition which did not yield the polymer functionalized surface as confirmed by ellipsometry, FT-IR, AFM and other studies.

The polymerization method was extended on other surfaces such as ITO which are more useful for device fabrications.³⁸ In a similar fashion, the self-assembled monolayers of azide followed by click reaction and further “graft-through” Sonogashira polymerization was performed using 2-ethynyl-5-(5-ethynylthiophen-2-yl)thiophene and 1,4-dibromo-2,5-bis(hexyloxy)benzene in the presence of Pd(0) catalyst system.

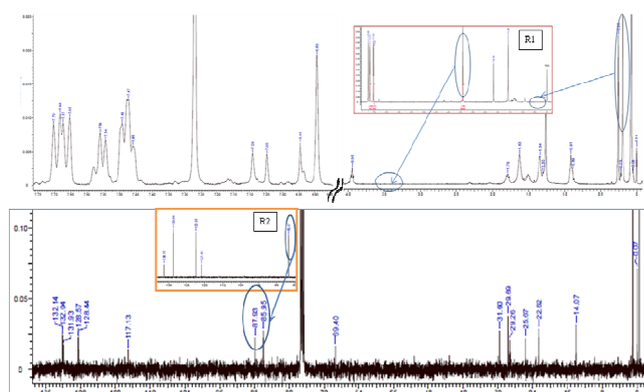


Fig. 3. ¹H (top) and ¹³C (bottom) NMR spectra of polymer. Inset: respective NMR of the reactants.

Fig. 4A shows the UV-vis spectra of the ITO surface after end alkyne functionalization (black curve) and polymerization (red curve), respectively. Absorption maxima (λ_{\max}) appeared at 414 nm for the polymer functionalized surfaces. This value is typical for a conjugated aromatic system. The band gap was 1.78 eV calculated from the Tauc plot from absorption data (Fig S7 ESI[†]). This low band gap should be important for the designing of solar cell or molecular electronics materials. Similar UV-vis spectra were shown in reflection mode for polymer functionalized ITO surface (Fig. 4(B)).

The devices were fabricated by using above mentioned polymer functionalized Indium Tin oxide (ITO) substrate (thickness 110 nm, 10 Ω /square) followed by the deposition of aluminum as cathode at 10⁻⁵ torr (Fig. 4 (C), inset) for charge carrier mobility study

and I-V characteristics. The Current-Voltage (I-V) characteristics were measured using Keithley 2400 source meter respectively. Current-Voltage (I-V) characteristic curve (Fig. 4(C)) showed that the polymer had characteristics of a semiconducting polymer which is a clear indication of the surface functionalization by “graft-through” approach. Similar I-V characteristic was observed when the ITO surface was spin coated with the similar polymer obtained in solution (Fig. S8, ESI). However, when control experiment was performed on ITO surface (polymerization in presence of pristine ITO without grafted alkyne) similar sandwich structure did not produce characteristic I-V curve for semi conductive material (Fig. S9, ESI). Further the surface functionalized ITO substrates were studied for charge carrier mobility by Space Charge Limited Current (SCLC) method.^{39, 40} The charge carrier mobilities of quenched polymer functionalized self assembled mono layer on ITO was found to be $1.23 \times 10^{-6} \text{ cm}^2 \text{ V}^{-1} \text{ S}^{-1}$. The band structure of the device (Fig S3, ESI) indicated that the hole and electron injection barriers are 0.52 and 0.56 eV respectively. Hence, the mobility value corresponds to ambipolar mobility.

The surface bound conjugated polymers on ITO surface were also characterized via cyclic voltammetry (CV) (Figure 4(D)) using Ag/AgCl as reference electrode to find the band gap, HOMO-LUMO energy levels. The equations⁴¹: $\text{HOMO} = - (j_{\text{ox}} + 4.71)$ and $\text{LUMO} = - (j_{\text{red}} + 4.71)$ was used to calculate the band gap. The oxidation process was observed at 1.07 V and onset of the reduction process was observed at -0.67 eV. Hence the corresponding HOMO and LUMO level was determined as -5.78 eV and -4.04 eV respectively. Thus the band gap (HOMO-LUMO gap) was calculated to be 1.74 eV, much lower than our previously reported polytriazole system.¹³

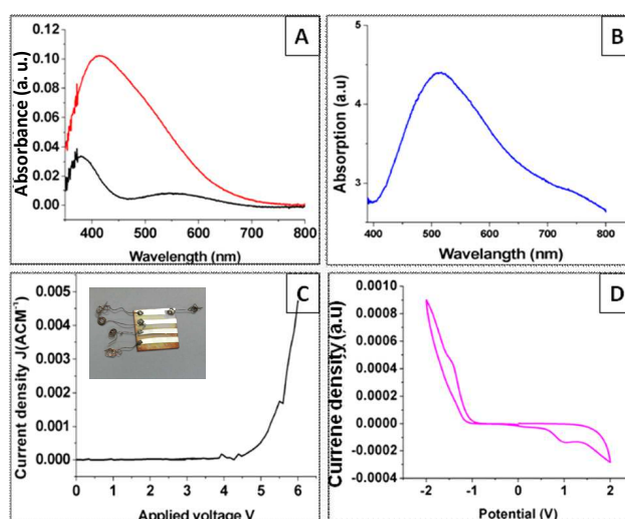


Fig. 4. (A) UV-vis absorbance spectra of ITO surfaces: alkyne functionalized (black) and polymer functionalized (red curve). (B) UV-vis spectra (reflection mode) on polymer functionalized ITO surfaces. (C) Current Voltage characteristics of the polymer grafted ITO surface in the fabricated device using aluminium as cathode in a sandwich structure. Inset: typical fabricated device using aluminium as cathode for this study (D) Cyclic voltammogram of polymer functionalized ITO surface

In conclusion, we demonstrated that “graft-through” Sonogashira polymerization can be used to functionalize various surfaces with conjugated thiophene-based polymers. The process is mild, requires only three-steps for getting an appreciable thickness of the polymer anchored on the surface. It is also superior to our previously reported click polymerization method in terms of continuous conjugation and lower band gap, determined from cyclic voltammeter studies and UV-Vis spectroscopic studies. The fabricated device with functionalized ITO as anode and deposited aluminium as cathode showed an I-V characteristic typical of a semiconducting material with moderate charge carrier mobility. The material further indicated that the methodology should be useful for the fabrication of conjugated polythiophene-based devices. We plan to pursue some of our research efforts in that direction. We thank Dr. A. Dhathathreyan of CLRI for assisting with AFM and contact angle data acquisition; Dr. Chakravarty of IACS, Kolkata for assisting with XPS. Financial support from CSIR network project NWP54 is gratefully acknowledged. CSIR-CLRI communication no. 1101.

Notes and references

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† Electronic Supplementary Information (ESI) available: [experimental procedure and instrument details and various characterization data]. See DOI: 10.1039/b000000

‡ Typical procedure for Sonogashira polymerization: We took hot and cleaned 10 ml 2 neck round bottom flask, degassed with N₂ gas and added alkyne-functionalized ITO surface, freshly distilled 10 ml diisopropylamine. After that, we added 2-ethynyl-5-(5-ethynylthiophen-2-yl)thiophene [25mg, 0.116 m.mol], 1,4-dibromo-2,5-bis(hexyloxy)benzene [43mg, 0.10 m.mol], CuI [5 mg, 0.026 m.mol], palladium(II) acetate [5mg, 0.022 mmol] and triphenylphosphine [15mg, 0.057 m.mol]. After that the reaction mixture was heated at 75 °C for 12 or 24h. Then, we added 0.1 ml of ethynyltrimethylsilane, and refluxed for another 12 h. The ITO plate was taken out and washed with CHCl₃, toluene, H₂O, acetone, ethanol [each one 15 minutes] in a sonicator, dried over by N₂ atm.

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Supplementary Information for:
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1. Materials

3-chloropropyltrimethoxysilane, sodium azide, CuI, diisopropylamine, N,N-dimethylformamide, 2-bromo-5-(5-bromothiophen-2-yl)thiophene, hydroquinone, bromohexane, bromine were purchased from Sigma-Aldrich. 3-Azidopropyltrimethoxysilane was prepared from 3-(chloropropyl)trimethoxysilane and sodium azide using literature procedure.¹ 2-ethynyl-5-(5-ethynylthiophen-2-yl)thiophene (1) was freshly prepared from 2-bromo-5-(5-bromothiophen-2-yl)thiophene using the method described by Wong and others. 1,4-dibromo-2,5-bis(hexyloxy)benzene was prepared from hydroquinone using the literature procedure and used immediately.

2. Imaging and spectroscopy

UV-vis spectra were recorded on Cary 50 Bio spectrometer. Cyclic voltammetry was performed on CHI600D electrochemical workstations with ITO substrate as working electrode, Ag/AgCl as reference electrode and support electrolyte used was tetrabutylammonium hexafluorophosphate. Atomic force micrographs were obtained from Nova 1.026 RCI atomic force microscope with NT-MDT solver software analysis. Contact angle measurement was done using Holmarc HO-IAD-CAM-01 instrument. Thickness measurement was performed using Filmetric F20-UV thin film analyzer. XPS measurements were done in Omicron Nanotechnology, GmbH XPS (ESCA).

3. Formation of azide functionalized ITO, Si surface

The ITO plate was sequentially washed with toluene, water, acetone, ethanol (each one 10 minutes), followed by plasma cleaning for 10 minutes. The ITO, Si plate was placed in (H₂O:H₂O₂:NH₃) in the ratio of (5:1:1) at 70 °C for 3h. After 3h, the ITO, Si plate was washed with distilled water for 3 times.

The ITO or Si plate was placed into 10 ml hot and cleaned round bottom flask and added 0.1 ml of 3-azido propyl (trimethoxysilane) and 5 ml of dry toluene in N₂ atm at room temperature. It was then refluxed at 100 °C for 2.5 h. After the completion of reaction, the ITO or Si plates were washed and used further.

4. Formation of triazole and end alkyne functionalized ITO, Si surface.

We took 10 ml 2 neck round bottom flask degassed with N₂ gas, and we added end azide functionalized ITO or silicon plate and 5 ml of dry toluene under N₂ atm at 0 °C. Then, we added N₂ purged solution of [DMF 4ml+CuI-16mg + freshly distilled diisopropylamine -16 microliter] and 2-ethynyl-5-(5-ethynylthiophen-2-yl)thiophene [35mg, 0.163 m.mol] at 0 °C. After that the reaction mixture was heated at 60 °C for 24h. After the completion of reaction the ITO, Si plate was washed with toluene, CHCl₃, water, acetone and ethanol by ultrasonication method and kept under N₂ atm.

5. Sonogashira polymerization on ITO surface

We took hot and cleaned 10 ml 2 neck round bottom flask, degassed with N₂ gas and added alkyne-functionalized ITO surface, freshly distilled 10 ml diisopropylamine. After that, we added 2-ethynyl-5-(5-ethynylthiophen-2-yl)thiophene [25mg, 0.116 m.mol], 1,4-dibromo-2,5-bis(hexyloxy)benzene [43mg, 0.10 m.mol], CuI [5 mg, 0.026 m.mol], palladium(II) acetate [5mg, 0.022 mmol] and triphenylphosphine [15mg, 0.057 m.mol]. After that the reaction mixture was heated at 75 °C for 12h. After the completion of 12h, we added 0.1 ml of ethynyltrimethylsilane, and refluxed for another 12 h. The ITO plate was taken out and washed with CHCl₃, toluene, H₂O, acetone, ethanol [each one 15 minutes] in a sonicator, dried over by N₂ atm. Fig. S1 represents the AFM pictures of ITO surface before polymerization (A) and after polymerization (B and C).

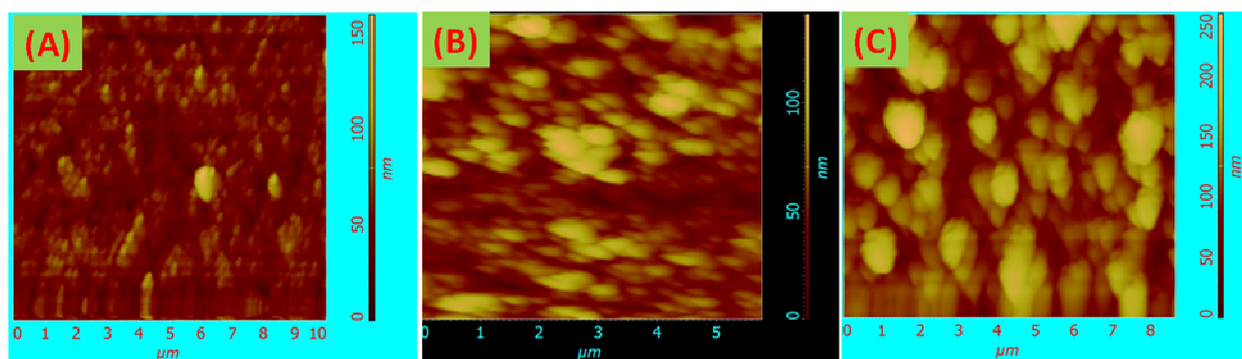


Fig. S1 AFM images of (A) alkyne-functionalized ITO plate, (B) ITO plate after the polymerization reaction for 12 h (C) ITO plate after the reaction for 24h.

6. Study of polymerization by NMR spectroscopy in residual solution

After the “graft through” polymerization on surface, the residual solution was studied by ^1H and ^{13}C NMR (Fig. S2). In ^1H NMR, the peak at 3.4 ppm (Inset, R1) disappeared while new peak at 0.26 ppm appeared. In ^{13}C NMR, peak characteristic at ~ 88 ppm (Inset, R2) was changed.

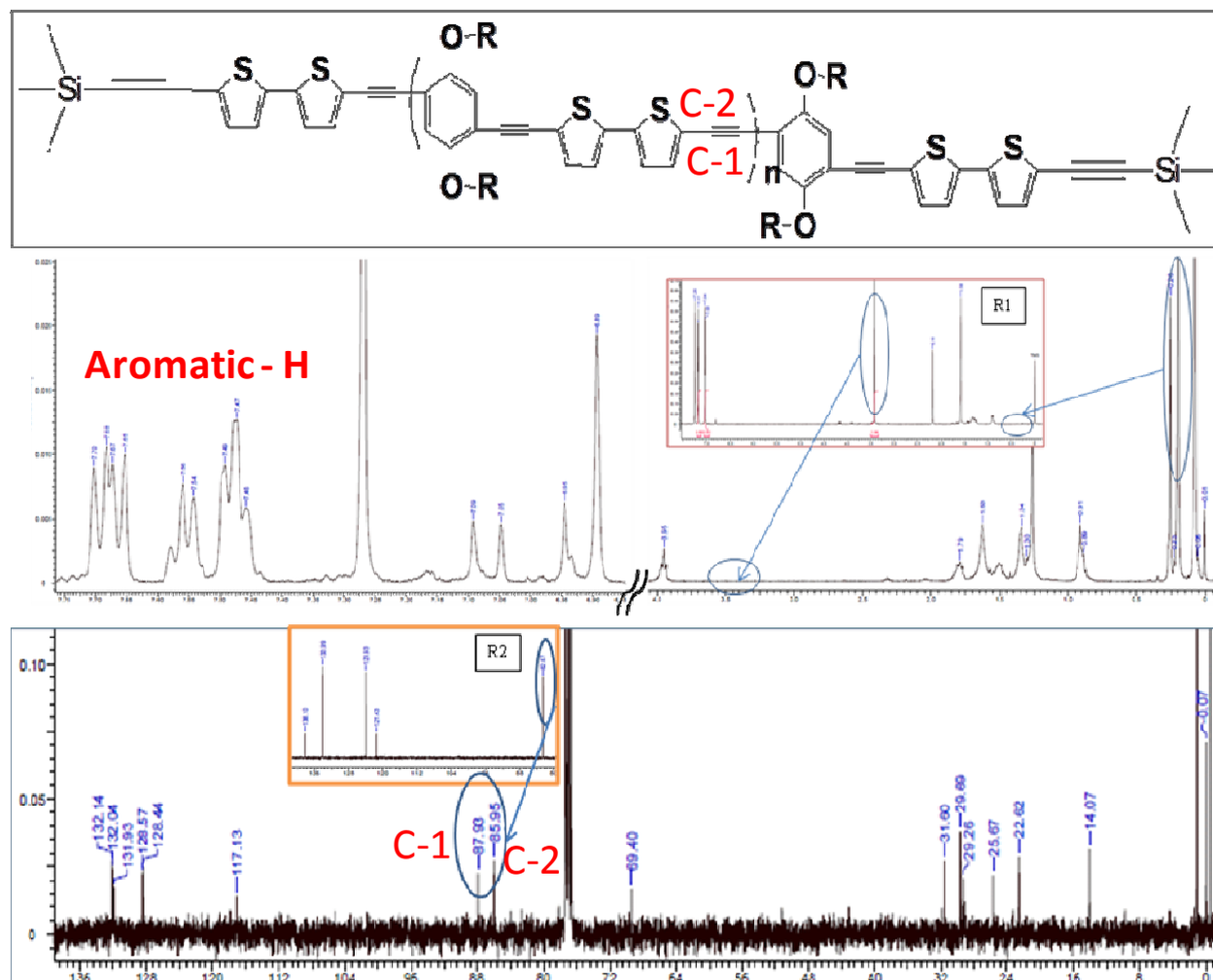


Fig. S2. ^1H (top) and ^{13}C NMR (bottom) of the Sonogashira polymer from the solution after “graft through” polymerization. Inset (R1) and (R2): NMR of the starting material (2-ethynyl-5-(5-ethynylthiophen-2-yl)thiophene).

7. Device studies

The device was fabricated as a sandwiched structure to measure I-V characteristics with eight sets of experiments. Measurements were carried out in ambience with the thickness of 100 to 150 nm and the area of 0.6 cm^2 ($0.3\text{ cm} \times 2.0\text{ cm}$). The charge carrier mobility of polymer functionalized layer was determined under space charge limited current (SCLC) condition.

It is clear from the band structure of the device (Fig S3) that the hole and electron injection barriers are 0.52 and 0.56 eV respectively. Hence, the mobility value corresponds to ambipolar mobility.

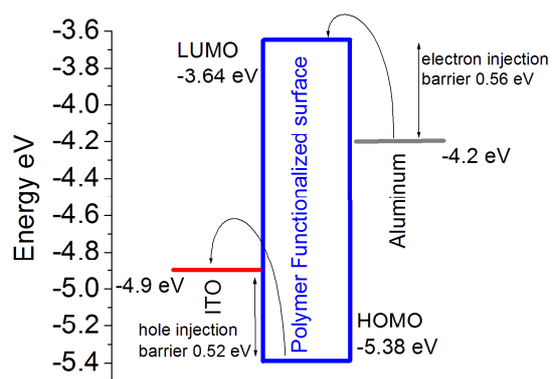


Fig. S3. Energy level diagram of polymer functionalized layer in the device configuration of ITO/grafted polymer/Al

8. AFM study of the polymer functionalized silicon surface

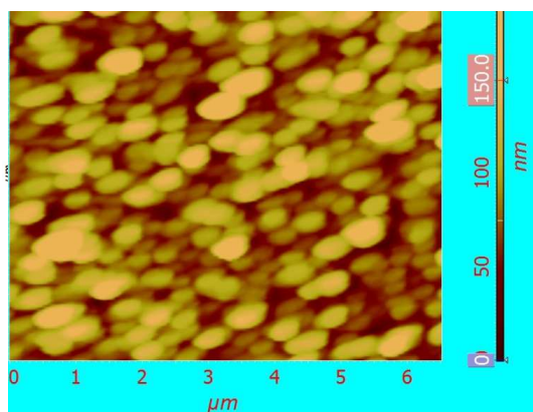
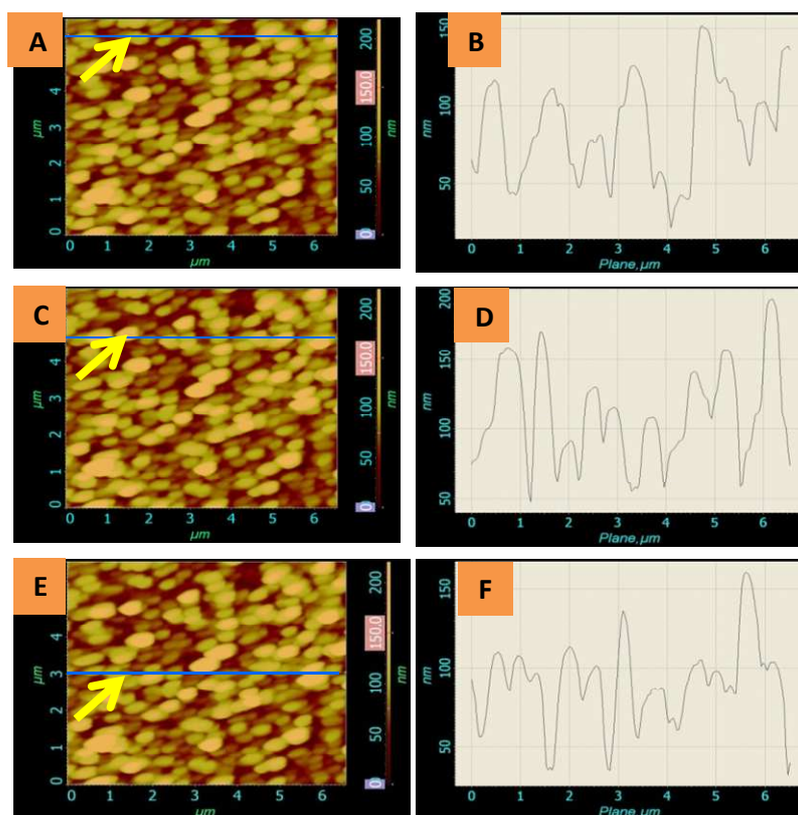


Fig. S4. AFM image of polymer functionalized silicon surface after performing the polymerization reaction for 24h

9. Study of distribution profile of surface by AFM



9.

Fig. S5. Distribution profiles of grafted polymer across the polymer functionalized silicon surface studied by AFM using NOVA software when the polymerization reaction was performed for 24h

10. XPS studies on silicon surface

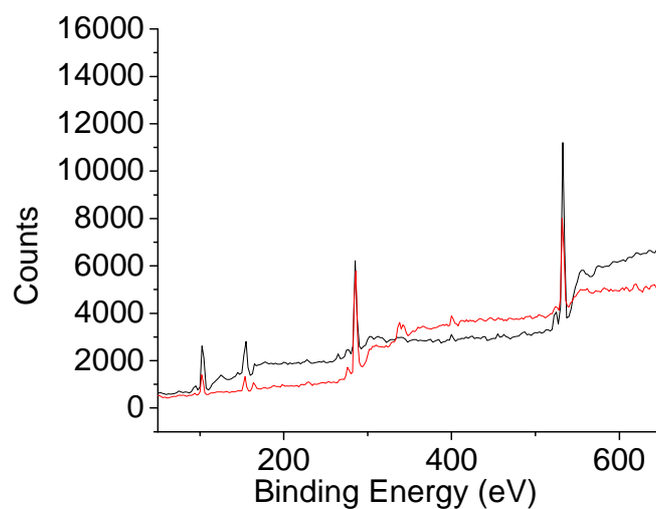


Fig. S6. XPS spectra of alkyne-functionalized (black curve) and polymer grafted (red curve) silicon surfaces (polymerization time 24 h).

11. TAUC plot

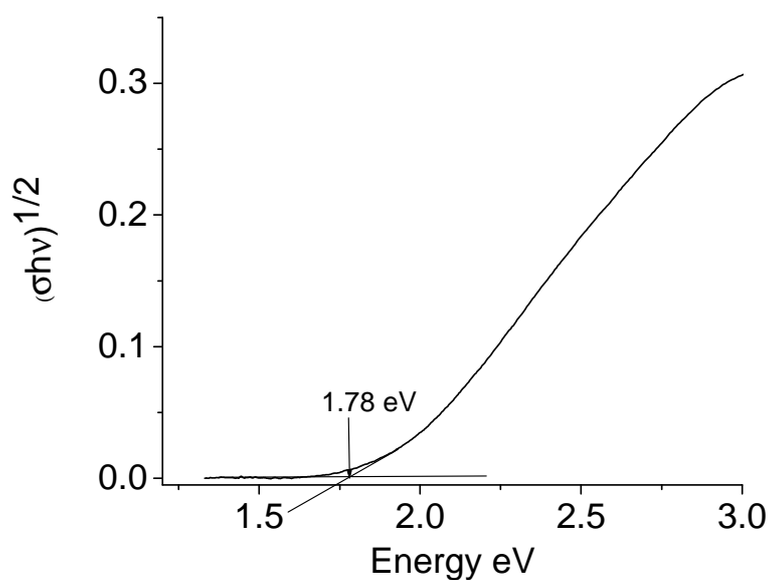


Fig. S7. TAUC plot for the calculation of band gap for polymer functionalized surface

12. Control experiments for I-V characterization on ITO plate

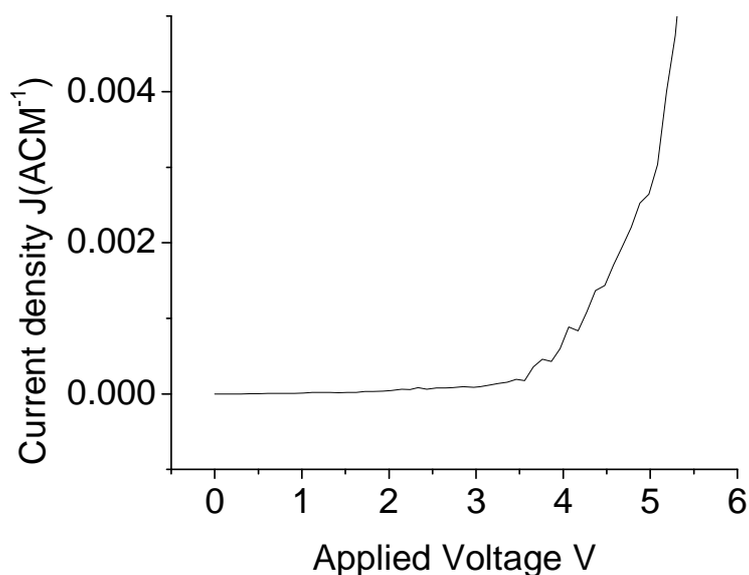


Fig. S8. Current Voltage characteristics of the ITO surface spin coated with polymer in the fabricated device using aluminium as cathode in a sandwich structure.

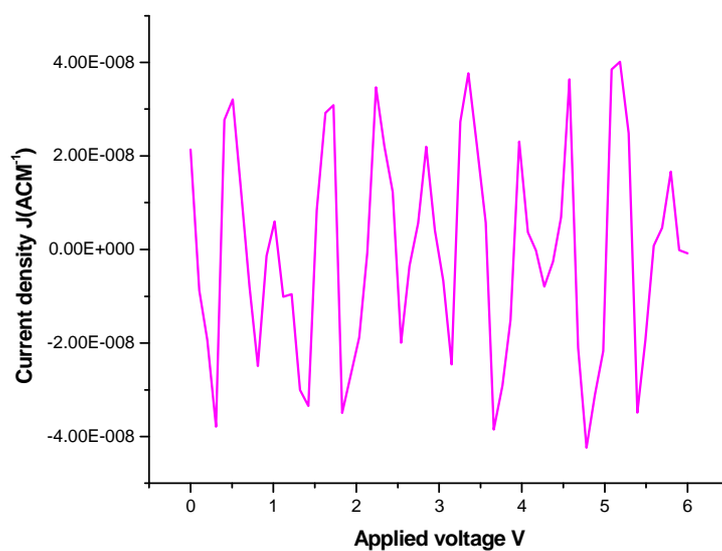


Fig. S9. Control experiment: Current-Voltage characteristics of the ITO surface (without polymer) in the fabricated device using aluminium as cathode in a sandwich structure.

1 P. Paoprasert, J. W. Spalenka, D. L. Peterson, R. E. Ruther, R. J. Hamers, P. G. Evans, P. Gopalan, *J. Mater. Chem.*, 2010, **20**, 2651.