Effects of Odd-Even Side Chain Length of Alkyl-Substituted Diphenyl-bithiophenes on First Monolayer Thin Film Packing Structure

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ABSTRACT:

Due to their preferential two-dimensional layer-by-layer growth in thin films, 5,5′bis(4-alkylphenyl)-2,2′-bithiophenes (P2TPs) are model compounds for studying the effects of systematic chemical structure variations on the thin film structure and morphology, which, in turn, impact the charge transport in organic field-effect transistors. For the first time, we observed –by grazing incidence X-ray diffraction (GIXD)– a strong change in molecular tilt angle in a monolayer of P2TP, depending on whether the alkyl chain on the P2TP molecules was of odd or even length. The monolayers were deposited on densely packed ultra-smooth self-assembled alkane silane modified SiO₂ surface. Our work suggests that a subtle change in molecular structure can have a significant impact on the molecular packing structure in thin film, which, in turn, will have a strong impact on charge transport of organic semiconductors. This was verified by quantum-chemical calculations that predict a corresponding odd-even effect in the strength of the intermolecular electronic coupling.
INTRODUCTION

A range of low-cost and large-area applications are emerging from the field of organic electronics, such as solid-state lighting\(^1,2\), solar cells\(^3,4\) and displays\(^5,6\). Different performance parameters such as luminescence efficiency in organic light-emitting diodes (OLEDs), power conversion efficiency in organic photovoltaic cells (OPV) or field-effect mobility in organic field-effect transistors (OFETs) are strongly determined by the morphology of the thin films and by the packing and orientation between individual molecules\(^7,8,9,10,11\). Specifically, molecular packing strongly impacts the electronic coupling between molecules and the resulting charge carrier mobility\(^12,13\). As a result, developing and understanding methods to control molecular packing of organic semiconductors by molecular design as well as processing control have been actively pursued by many research groups\(^14,15,16,17,18,19,20,21,22\). In particular, we have been interested in molecular design rules for controlling thin film packing structures. We have been able to determine the precise molecular packing structures for thin films of pentacene, TIPSE-pentacene, and a series of fluorine-bithiophene oligomers from grazing incidence X-ray diffraction (GIXD) data combined with numerical fitting\(^22,23,24\). This development is important for structure-property relationship studies because it allows us to understand organic semiconductor packing on substrates that are directly used for device fabrication.

One of the common strategies to promote a more favorable two-dimensional growth is to substitute the two ends of a small molecule organic semiconductor with linear alkyl chains. which results in large and more connected grains, Indeed, using this approach, higher thin film transistor mobilities have been reported for sexithiophene (a-6T)\(^25\), flourene-bithiophene-flourene (FTTF)\(^26,27\), acene derivatives\(^28\), [1]benzothieno[3,2-b]benzothiophene (BTBT)\(^29\), naphthalenetetrahydroxylic diimide (NTCDI)\(^30,31\), just to name a few. So far, mostly even-numbered linear alkyl chains have been used. We hypothesized that organic semiconductor molecules with odd- and even-numbered alkyl chain substitutants may result in different molecular packing and tilt angles on a smooth substrate surface – an “odd-even” effect. Indeed, odd-even effects have been observed for self-assembled alkane thiol self-assembled monolayers (SAM) on gold surfaces resulting in different water contact angles\(^32\). These odd and even-numbered SAMs
were also found to give rise to different liquid crystal molecular orientations for films deposited above them\textsuperscript{33}. Therefore, in this work, we investigate the odd-even effects of organic semiconductor side chain length, as well as the dielectric surface modification with alkane silanes, on the molecular packing and the charge transport. Such studies are important since for charge transport, minor changes in molecular packing structures can have a significant impact on charge transport. Therefore, this understanding may provide new insights for molecular design of organic semiconductors.

Herein, we have employed a series of 5,5’bis(4-alkylphenyl)-2,2’-bithiophene (P2TP) molecules, with varying side chain length (3–8 methyl units; see Figure S1 for chemical structures,), as the organic semiconductors. P2TPs exhibit a two-dimensional layer-by-layer growth when thermally evaporated and are known to form thin films with a preferential long axes of the molecule standing upright\textsuperscript{34,35}. Charge carrier mobility greater than 0.1 cm\textsuperscript{2}/Vs has been reported\textsuperscript{34,35,36}. In organic thin film-based field-effect transistors (FETs), the charge transport occurs in the first few nanometers of the semiconductor at the dielectric-organic interface\textsuperscript{34,37}. Therefore, the choice of dielectric material strongly influences the charge transport in an FET, because it determines the morphology of the semiconductor and the possible presence of energetic traps at the interface\textsuperscript{8,38,39}. In particular, the presence of hydroxyl groups on the commonly used Si/SiO\textsubscript{2} substrates strongly influences the mobility of charges in the transistor channel due to charge trapping\textsuperscript{40,41,42}, and limits the operational stability of organic FETs due to a reversible proton migration from the semiconductor to the dielectric\textsuperscript{43,44}. To encapsulate the hydroxyl traps, the Si/SiO\textsubscript{2} substrates are frequently treated with hydrophobic monolayers, such as OTS\textsuperscript{41,45}. Previously, we have shown that a preferable, more two-dimensional, layer growth of thermally evaporated organic semiconductors is achieved when the density of octadecyltrimethoxysilane (OTMS) on the Si/SiO\textsubscript{2} substrate is increased\textsuperscript{46}. Moreover, spin-coated OTMS can form crystalline self-assembled monolayers (SAMs) on amorphous SiO\textsubscript{2}, leading to a highly enhanced two-dimensional growth of the semiconductor\textsuperscript{47}. The crystalline OTMS was shown to result in a much lower surface roughness compared to vapor-deposited OTMS\textsuperscript{47}. The improved organic
semiconductor thin film morphology on crystalline OTMS results in transistors with an increased field-effect mobility by approximately an order of magnitude\textsuperscript{46,47}. Since the dielectric surface is very important for controlling the growth of the organic semiconductor, this study will involve the investigation of varying the organic semiconductor side chain odd-even effects in conjunction with Si/SiO\textsubscript{2} dielectric surfaces treated with crystalline heptadecyltrimethoxysilane (HDTs, C\textsubscript{17}) and octadecyltrimethoxysilane (ODTS, C\textsubscript{18}) SAMs.

**RESULTS AND DISCUSSION**

Fabrication of crystalline SAMs and devices was carried out according to previously published procedures\textsuperscript{46,47}. On highly n-doped silicon wafers with 300 nm thermally grown oxide, a crystalline monolayer of HDTs or ODTS was formed from solution. The crystalline monolayers of HDTs and ODTS exhibit very similar surface properties, with a water contact angle of around 108° and a mean surface roughness (RMS) of ~0.3-0.4 nm (see supplementary information, Figure S2 and S3).

Grazing incidence X-ray diffraction characterizations of the HDTs and ODTS were performed to confirm the crystalline nature of the SAMs. Bragg rods were found at discrete in-plane momentum transfer values \(q_{xy}\) from which the dimensions of the HDTs and ODTS monolayer unit cells were determined. The diffraction images of ODTS and HDTs contain a Bragg rod at \(q_{xy}\) of around 1.50 \(\text{Å}^{-1}\) and 1.51 \(\text{Å}^{-1}\), respectively. This Bragg rod is due to the degenerate \{01\}, \{10\}, and \{-11\} diffraction peaks of the hexagonal lattice formed by the crystalline ODTS and HDTs monolayers (see Figure 2A). The corresponding ODTS and HDTs lattice constants of 4.84 \(\text{Å}\) and 4.81 \(\text{Å}\) are consistent with previous findings\textsuperscript{46}. That the Bragg rod intensity is peaks near \(Q_z=0\) implies that the molecules are standing nearly upright (untilted).

To study the growth of the first layer of P2TP molecules, we evaporated sub-monolayer P2TP films on HDTs and ODTS, respectively. The conditions for vacuum deposition are listed in Table 1. To achieve a highly two-dimensional layered growth, the substrate temperature, \(T_{\text{sub}}\), was chosen to be 60°C for 5,5’bis(4-pentylphenyl)-2,2’-bithiophene (C\textsubscript{5}-P2TP-C\textsubscript{5}). Hereafter, molecules with a side chain
length of $N$ methyl units will be abbreviated as $C_N$-P$\text{TP}$-$C_N$. For the heavier, longer molecules, higher substrate temperatures were chosen (Table 1) to offset the higher molecular weight and thus facilitate thermally activated molecular surface diffusion that is less dependent on the molecular weight.

Figure 1 shows Atomic Force Microscopy (AFM) images of the corresponding films, with images A-F showing the layers evaporated on HDTs, and images G-L evaporated on ODTS, respectively.

We found a surprising difference in the initial growth mode between the P$\text{TP}$ layers on HDTs and ODTS. With the exception of $C_8$-P$\text{TP}$-$C_8$, all P$\text{TP}$ molecules on ODTS showed areas of initially flat-lying molecules (indicated by the black regions in Figures 1G-K) in the first layer on ODTS, whereas no indication for flat-lying molecules was found in the P$\text{TP}$ films on HDTs. Since every image of P$\text{TP}$s on ODTS showed clusters of grains comprised solely of standing-up molecules, it is possible that when the molecule-molecule interaction is stronger than the molecule-surface interaction the flat-lying molecules diffuse over the surface and flip upright near the edge of an island of upright oriented molecules. This hypothesis is compatible with the complete absence of any scattering signal in the GIXD images that could be attributed to a second, crystalline phase of lying molecules, though this could also be due to lack of any crystalline ordering in the lying phase. It is also consistent with Molecular Dynamics simulations of similar molecules$^{48}$.

With increasing side chain length—from three methyl units up to eight (indicated by C3-C8)—we observed an increase in apparent grain size. As shown in Figure S4, we observed an increasing grain size of $C_8$-P$\text{TP}$-$C_8$ with substrate temperature from room temperature to 115°C, indicating substrate temperature as a large contributor to the increase in grain size. However, we also observed an increased grain size with molecular chain length for 40 nm films evaporated at 60°C (see Figure S5), clearly indicating an effect of side chain length on grain size. The same trend was previously observed for oligoacenes$^{49,50}$ and oligothiophenes$^{51}$, where higher-aspect molecules tend to form more two-dimensional films with large islands. Larger islands are the result of a lower nucleation density, which, in the case of longer molecules, originates from molecule-molecule interactions that are increased relative to the molecule-substrate interactions. The molecule-substrate interactions are similar for all
P2TP derivatives and dominated by the interactions between the terminal methyl group and the substrate surface.

Grazing incidence X-ray diffraction characterization of the $C_N$-P2TP-$C_N$ films was performed to determine the unit cells and corresponding molecular packing inside these unit cells on HDTS and ODTS substrates. These cell dimensions were obtained from the measured in-plane diffraction peak positions by using a least-square error minimization algorithm. Besides the peaks that were attributed to the $C_N$-P2TP-$C_N$ sub-monolayers, the diffraction images of all investigated films showed an additional broad peak at around 1.50 Å$^{-1}$ that corresponds to a hexagonal lattice of the densely packed HDTS and ODTS films, which confirms that crystalline HDTS and ODTS SAMs were indeed present in all samples and that the crystalline structure of the SAMs were not altered by the growth of the P2TP derivatives. The motif by which the molecules pack inside the $C_N$-P2TP-$C_N$ unit cells was extracted by crystallographic refinement of the diffraction intensity data obtained from these Bragg rods. This was achieved by fitting theoretically calculated diffraction intensities to the measured intensity data, assuming scattering from a smooth thin film is semi-kinematical, see Figure 2B. Several corrections to the theoretical scattering intensity need to be taken into account, as explained in the supplementary information.

Even without the crystallographic refinement, the diffraction data in Fig. 2A show the odd-even effect. For $C_3$-P2TP-$C_3$, the Bragg rod intensity peaks near $Qz=0$ and from this, we can immediately conclude that the molecules are nearly standing upright. For $C_4$-P2TP-$C_4$, the Bragg rod intensity for the (02) peaks near $Qz=0.6$ Å$^{-1}$, which would give a tilt angle of about 22° (atan(0.6/1.5), assuming the molecule tilts along the b-axis, which is justified since the (11) Bragg rod has a maximum intensity at $Qz=0.3$ Å$^{-1}$ ($\frac{1}{2}$ of 0.6 Å$^{-1}$). This approximate calculation ignores complications due to the molecular structure (e.g., it assumes a cylindrical molecule as for many Langmuir monolayer films), but it provides a semi-quantitative picture that supports the more detailed calculations below.

Minimization of the crystallographic residual was performed by in-house software POWERGRID which implements the Monte Carlo Simulated Annealing method. For this refinement, integrated
intensities $I(qz)$ were measured along the Bragg rods, and the molecular alignment varied until the best agreement between measured and calculated intensities is achieved. Only the clearly measurable rods were included in the fit (typically {11}, {02}, {12}, but in some cases only {11}, and {02}. In addition, owing to the significance of the absence of the {01} and {10} for the symmetry in the molecular packing motif, the {01} and {10} rods were included in the calculation and set to zero. For the computation to be feasible, the $C_N$-P2TP-$C_N$ molecules were treated as rigid objects. This assumption is justified only in case of the shorter alkyl chain derivatives since alkyl chains beyond a certain length (we estimate $> 4$ methyl groups) possess inherent flexibility that is not properly captured by the rigid molecule approximation. Specifically, while good structural refinements could be obtained for $C_N$-P2TP-$C_N$ with $N = 3, 4, 5$, the fits became increasingly worse for the longer derivatives ($N = 6, 7, 8$). However, for two reasons, the crystallographic refinement can still be performed, even for the longer chain derivatives: i) a significant portion of the peak intensity is determined by the heavy sulfur atoms on the essentially rigid P2TP backbone (geometry-stabilized by the $\pi$-$\pi^*$ interactions between the aromatic carbon atoms) that scatter far stronger than the carbon atoms, and ii) the alkyl chains, especially the longer ones, do not necessarily contribute significantly to the peak intensity (due to disorder) but rather create a diffuse scattering background. Therefore, we assumed the intensity along the Bragg rods to be mostly determined by the alignment of the P2TP core alone. The quality of the fit between theoretical and experimental intensity profiles when the fitting was performed for the P2TP core only (see Figure 2B) supports this conclusion. The molecular geometry of the P2TP core was calculated using the Parametric Model number 3 (PM3) semi-empirical method. Besides an overall scaling factor, 10 degrees of freedom were considered in the optimization: 3 Euler angles for each P2TP molecule, 3 coordinates describing the position of the second molecule relative to the first one, and the molecular mean square displacement $\langle U^2 \rangle$. Thus, no artificial constraint was placed on the molecular positions in the unit cell.

The best-fit theoretical $I_{hk}(qz)$ functions and the corresponding experimental $I_{hk}(qz)$ values are displayed in Figure 2B; the corresponding arrangement of the two $C_N$-P2TP-$C_N$ molecules after 2.5
million Monte Carlo steps is shown in Figure 2C. The optimization was repeated numerous times, leading to identical best-fit configurations in all runs. For all chain lengths where the Bragg profiles were refined (N=3-8), the unit cell contains two P2TP molecules that are aligned in the well-known herringbone pattern (see center of Figure 3C) whose two glide-symmetry planes explain the absences of diffraction intensity in the {01} and {10} rods. This arrangement is common in most linear aromatic molecules, for example, pentacene\textsuperscript{23}. Note the weak intensity of the {20} Bragg rod in the GIXD images in Figure 2. This results from the rather uniform electron density along the a-direction as can be seen in the top-down view in Figure 2.

A clear odd-even effect is obtained for both the long unit cell axis b, and the tilt angle of the P2TP core with respect to the substrate surface normal (see Table 2). For the three odd-length derivatives, in which N = 3, 5 or 7, the b axis is shorter, at around 7.8 Å, and the tilt angle is small, i.e., the P2TP core is nearly vertical relative to the substrate. In contrast, the b axis in the case of the even length molecules, in which N = 4, 6, or 8 is substantially longer (8.2-8.4 Å), and the molecules are significantly more tilted, with tilt angles around 20°, as shown for C\textsubscript{4}-P2TP-C\textsubscript{4} in the right part of Figure 3C. The concurrent increase in b axis length and tilt angles for the even-chain length derivatives is consistent with the finding that the tilt of the P2TP cores is almost exclusively in the direction of the b axis. Because of the tilt, the projection of the molecules in the substrate surface plane is larger and, consequently, the unit cell is expanded in the tilt direction, i.e., the b axis. The GIXD intensity profiles, and consequently the P2TP molecular packing structure of C\textsubscript{N}-P2TP-C\textsubscript{N} sub-monolayers on HDTS, are nearly identical to those on ODTS, suggesting identical growth of C\textsubscript{N}-P2TP-C\textsubscript{N} on the two substrates. This was verified in fits for C\textsubscript{3}-P2TP-C\textsubscript{3} and C\textsubscript{4}-P2TP-C\textsubscript{4} on HDTS that yielded tilt angles and packing motifs that were practically identical to those on ODTS.

To provide insights into the mechanism leading to the odd-even effect, we performed simulations of the system using Molecular Dynamics (MD) computations. Experimental length scales are inaccessible using MD simulations; a tractable-sized system in the simulation is composed of a herringbone lattice consisting of 60-80 P2TP molecules. Periodic boundaries were employed in the lateral x- and y-
directions, which simulate infinite 2D lattice structures efficiently and remove edge effects. In this system, alkylated P2TP molecules with an optimal lattice configuration (described in the supplementary information) were placed on top of a thermalized ODTS surface at a distance between the long-axis of the P2TP and the ODTS surface of approximately 3Å, corresponding to the minimum of the van der Waals interactions. At the start of the simulation, all the atoms were fixed, corresponding to a temperature of 0 K, before the ensemble was thermalized at 300 K for about 100 ps, which is enough for the SAM surface to equilibrate. The density of the simulated ODTS monolayer was chosen to be 5 molecules/nm² (the same as the experimental density), and resulted in a tilt angle of 6 ± 2° for the ODTS molecules. This value matches closely the experimentally observed tilt angle. Based on the nearly identical GIXD diffraction signals from both ODTS and HTDS films, the same molecular density was also chosen for HDTDS. The deposition of a P2TP molecule on a SAM can, in principle, involve different possible outcomes, such as the insertion of the molecule in the SAM, adsorption on the surface, or scattering of the molecule. However, due to the high density of the ODTS monolayer, insertion is unlikely. Scattering does not affect growth habit, and hence only the surface adsorption effect is important here. Additional details of the simulations are provided in the supplementary information.

Table 3 shows the lattice parameters and tilt angles obtained from simulations. The simulations show a clear difference in tilt angle between odd- and even- side chain lengths, that closely mirrors experimental findings by GIXD. Predictions of a and b determined by simulation may not reproduce each individual experimental value, but the mean values of a and b are well within experimental error. More importantly, a distinct trend for b to oscillate between smaller and larger values for odd and even N values is seen in both experiments and simulation. This odd-even oscillation occurs because there is a significant energetic advantage for each set of C₉-P2TP-C₉ molecules to orient themselves to a characteristic tilt angle. For example, for N = 4, i.e., C₄-P2TP-C₄, the lowest energy occurs when the molecule is tilted by 19° (a 2 kcal/mol advantage relative to a tilt-free alignment, and a difference in the central energy maximum of 11 kcal/molecule). Furthermore, the position of lowest energy for both odd-
and even-numbered side chain molecules lies in-between the terminal methyl groups of ODTS (not on top of ODTS). However, the even-numbered side chain molecules tilt away from the substrate surface normal in order to maximize the interactions of the last methyl group of the ODTS with the terminal methyl group of the alkyl side chain of the P2TP molecule (shown in Figure 3) with the consequent change in the P2TP-P2TP interactions.

A question that remains is whether this odd-even effect is due to the C₅₋P2TP-C₅ - SAM interactions alone, or due to preferences for distinctly different crystal structures (say, herringbone and honeycomb) between molecules with odd- and even-numbered side chains. To answer this question, 2D crystals of the C₅₋P2TP-C₅ molecules without SAMs present were simulated. The simulation protocol described in the SI included a section in which changes to the lattice parameters adopted by the P2TP molecules were permitted. Despite the additional freedom of the P2TP molecules to reorient if energetically advantageous, the crystals continued to exhibit herringbone packing for both odd- and even-length chains. Table 4 shows the predicted lattice parameters and tilt angles for these 2D crystals with no interactions with the SAM. While the lattice parameters and tilt angles in the absence of the SAM are significantly different from those when the substrate is present, the odd-even effect on the tilt angles persists even without the presence of the SAM. Figure 4 compares the tilt angles of the two systems mentioned here. Overall, the selection of a tilt angle preferred by each P2TP molecule is a cumulative effect of the interactions of the P2TP molecules among themselves, accentuated by interactions between the P2TP crystal and the SAM, and the topology of the surface (if a substrate is present). Thus, the odd-even effect is observed whether the SAM is present or not (see SI for more details). However, the odd-even effect is enhanced in the presence of an underlying SAM substrate.

The significant odd-even effect that we observed in the tilt angle of alkylated P2TP molecules on both HDTS and ODTS monolayers can potentially result in a similar odd-even effect in the strength of the electronic coupling between molecules along the plane of the substrate. This is the plane in which charge transport takes place in organic field-effect transistors (OFETs). To study the effect on the charge transport in OFETs, 40 nm thick films of C₅₋P2TP-C₅ were evaporated, after which the
transistors were completed by evaporating 40 nm thick Au source-drain electrodes through a shadow mask. The AFM images showed two-dimensional grains with monolayer terraces (see Figure S5). The OFETs were measured in an N₂ glove box to exclude the effect of air on the devices. The electrical measurements and mobility calculations are provided in the supporting information. Figure 5 shows the saturated field-effect mobility obtained from the transfer curves, where the source-drain current ($I_{sd}$) is plotted versus gate voltage ($V_G$). No clear dependence of the mobility was observed with increasing chain length and no differences in charge transport between OFETS using HDTs and ODTS treated substrates were observed, i.e. no odd-even effect was observed within the error of measurement.

The charges in the transistor channel are confined to the first few nanometers and 2 to 3 layers of organic semiconductor dominate the charge transport. While the available X-ray data do not permit a structural refinement of a multilayer film comprised of very few layers, the similarity of the X-ray data between the sub-monolayer films and 2-3 layer thick films strongly suggests that the odd-even effect persists for at least the first 2-3 layers. However, factors such as the density and type of grain boundaries are known to play a significant role in determining the overall device performance. Therefore, while an odd-even effect could still exist in the intrinsic charge carrier mobility, i.e., in individual first monolayer grains of the different C₅₇-P2TP-C₅₇ films, this effect is not necessarily reflected in the aggregate device performance. Nonetheless, we attempted to fabricate monolayer dimensional devices which would allow us to measure the transport solely in the first layer of the organic semiconductor. However, due to a strong dependence of the mobility on the second layer coverage, the results showed a large variation and were inconclusive.

In order to estimate the impact of the odd-even effect on the intrinsic electronic coupling between the molecules in the C₅₇-P2TP-C₅₇ layer, DFT calculations were performed for the packing obtained from the GIXD refinement. The transfer integrals for the HOMO orbitals were calculated for the neighboring dimers at the B3LYP/6-31G(d,p) level of theory as implemented in Q-Chem 3.2 quantum chemistry package. The solubilizing alkyl chains are known to have no significant effect on the transfer integrals, and were therefore not included. The transfer integral values were computed for all
four inequivalent next-nearest neighbor transitions (dimer pairs) in the unit cell, but only the three transitions indicated in Figure 2C (P, T1, T2) yielded non-zero transfer integral values. The results for these transitions are summarized in Table 5. An odd-even effect occurs in all three dimer transitions, most pronounced in the diagonal transfer elements T1 and T2. Overall, these values suggest that there should indeed be an odd-even effect in the intrinsic hole conductivity (HOMO level). This effect was not observed experimentally due to the challenges discussed above.

**CONCLUSIONS**

The layered growth of C$_N$-P2TP-C$_N$, in which $N = 3$ to 8, was studied on crystalline monolayers of HDTs and ODTS. For the first monolayer of C$_N$-P2TP-C$_N$, an odd-even effect in tilt angle with increasing side chain length was observed by GIXD, with no observable differences between either HDTs- or ODTS-treated surfaces. The odd-even effect was theoretically verified by MD computations, where we observed that the tilting of the molecules oscillated between odd- and even- length alkyl chains even in the absence of a substrate. This can only mean that the phenomenon occurs as a result of the composite inter- and intra-molecular interactions among molecules. The effect is increased by the presence of the SAM whose interactions of the P2TP molecules alter the lattice parameters and the magnitude of the tilting. We observed no tendency for molecules in this series to alter their crystal structure from the initial herringbone crystal structure; hence this is not the origin of the odd-even effect. In the initial layer growth of C$_N$-P2TP-C$_N$ on HDTs an additional phase of lying molecules was observed in AFM measurements, but not during the GIXD investigations, suggesting that it might not be stable and could over time be converted to the respective standing (tilted or untilted) phases that is found on both HDTs and ODTS. Despite the pronounced odd-even effect in the film structure, no clear odd-even effect or dependence of side chain length on mobility was observed in FET measurements, which is likely due to the contribution of several layers to the charge transport and other limiting factors for charge transport, such as grain boundaries. In conclusion, we showed that small differences in dielectric surface or molecular structure can lead to significant effects during the growth of the first
layers of organic compounds by thermal evaporation. The origins of these differences were further understood by MD simulations. Additional studies need to cover a different range of molecules with similar systematic variations to broaden the understanding between molecular structure and macroscale thin film morphology.

METHODS

Highly n-doped Si wafers with 300nm thermally grown oxide were cleaned in a piranha solution (70:30, H$_2$SO$_4$:H$_2$O$_2$- caution highly reactive with organic compounds) and UV-O$_3$ treated for 5 minutes prior to spin coating. A solution of 3 mM hepta- or octa-decyltrimethoxysilane in trichloroethylene was deposited on the Si/SiO$_2$ wafers and allowed to partially self-assemble for 10 s, after which the substrate was spun at 3000 rpm. The substrates were subsequently placed in a closed container together with a small vial with hydrochloric acid for a period of >5 hours to promote the bonding to SiO$_2$ by the hydrolysis of the anchoring group. Afterwards, the samples were rinsed with toluene and water. The thin films of alkyl substituted P2TPs were thermally evaporated in a vacuum chamber (< 5×10$^{-6}$ Torr) at elevated substrate temperatures at a rate of 0.3-0.5 Å/s. For the field-effect transistors Au source-drain electrodes were thermally evaporated through a shadow mask at room temperature. Electrical characterization was done using a Keithley 4200 Semiconductor Analyzer.

Grazing incidence X-ray diffraction of the C$_N$-P2TP- C$_N$ films was performed at the Stanford Synchrotron Radiation Lightsource (SSRL) on beam line 11-3. The diffracted intensity was recorded on a 2-D image plate (MAR-345) with a pixel size of 150 μm (2300 × 2300 pixels). The samples were ~10 mm long in the direction of the beam path, and the detector was located at a distance of 400 mm from the sample center (distance calibrated using a Lanthanumhexaboride standard). The incidence angle was chosen in the range of 0.10–0.12° to optimize the signal-to-background ratio. The beam size was 50 μm × 150 μm, which resulted in a beam exposure on the sample 150 μm wide over the entire length of the 10 mm long sample. The data were distortion-corrected (θ-dependent image distortion
introduced by planar detector surface) before performing quantitative analysis on the images using the software WxDiff\textsuperscript{22}.

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Supporting Information Available:

1. Chemical Structures.

2. Characterization HDTs and ODTs self-assembled monolayers

3. Influence of substrate temperature on apparent grain size of P2TPs

4. Grazing incidence X-ray diffraction

5. Computer simulations

6. Transfer characteristics FETs

This material is available free of charge via the Internet at http://pubs.acs.org.
Table 1. Thermal evaporation conditions for half a monolayer coverage of C₃-P2TP-C₃.

<table>
<thead>
<tr>
<th>Molecule</th>
<th>Substrate temperature (°C)</th>
<th>Evaporated thickness (nm)</th>
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<tr>
<td>C3-P2TP-C3</td>
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<tr>
<td>C4-P2TP-C4</td>
<td>53</td>
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<tr>
<td>C7-P2TP-C7</td>
<td>76</td>
<td>1.55</td>
</tr>
<tr>
<td>C8-P2TP-C8</td>
<td>84</td>
<td>1.7</td>
</tr>
</tbody>
</table>

Table 2. Unit cell geometries and P2TP core off-normal tilt angle on ODTs by GIXD (experimental uncertainty < 0.01 Å, for both a and b).

<table>
<thead>
<tr>
<th>Side chain length N</th>
<th>a / Å</th>
<th>b / Å</th>
<th>gamma / °</th>
<th>tilt angle / °</th>
</tr>
</thead>
<tbody>
<tr>
<td>3</td>
<td>5.66</td>
<td>7.85</td>
<td>89.9</td>
<td>1.01</td>
</tr>
<tr>
<td>4</td>
<td>5.70</td>
<td>8.46</td>
<td>90.0</td>
<td>22.79</td>
</tr>
<tr>
<td>5</td>
<td>5.70</td>
<td>7.86</td>
<td>90.0</td>
<td>5.95</td>
</tr>
<tr>
<td>6</td>
<td>5.71</td>
<td>8.42</td>
<td>90.0</td>
<td>20.41</td>
</tr>
<tr>
<td>7</td>
<td>5.74</td>
<td>7.81</td>
<td>90.2</td>
<td>1.20</td>
</tr>
<tr>
<td>8</td>
<td>5.72</td>
<td>8.21</td>
<td>90.6</td>
<td>18.8</td>
</tr>
</tbody>
</table>
Table 3. Unit cell geometries and P2TP off-normal tilt angle by simulations on SAMs.

<table>
<thead>
<tr>
<th>Side chain length N</th>
<th>a / Å</th>
<th>b/ Å</th>
<th>tilt angle / °</th>
</tr>
</thead>
<tbody>
<tr>
<td>3</td>
<td>5.77 ± 0.3</td>
<td>7.90 ± 0.3</td>
<td>3.5 ± 1.5</td>
</tr>
<tr>
<td>4</td>
<td>6.12 ± 0.3</td>
<td>8.07 ± 0.3</td>
<td>19 ± 1.5</td>
</tr>
<tr>
<td>5</td>
<td>5.85 ± 0.3</td>
<td>7.60 ± 0.3</td>
<td>4.2 ± 1.5</td>
</tr>
<tr>
<td>6</td>
<td>5.61 ± 0.3</td>
<td>9.37 ± 0.3</td>
<td>19 ± 1.5</td>
</tr>
<tr>
<td>7</td>
<td>5.85 ± 0.3</td>
<td>7.65 ± 0.3</td>
<td>4.5 ± 2</td>
</tr>
<tr>
<td>8</td>
<td>5.63 ± 0.3</td>
<td>9.64 ± 0.3</td>
<td>22 ± 2</td>
</tr>
</tbody>
</table>

Table 4. Unit cell geometries and P2TP off-normal tilt angle by simulations without SAMS.

<table>
<thead>
<tr>
<th>Side chain length N</th>
<th>a / Å</th>
<th>b/ Å</th>
<th>tilt angle / °</th>
</tr>
</thead>
<tbody>
<tr>
<td>3</td>
<td>5.68 ± 0.3</td>
<td>8.67 ± 0.4</td>
<td>16 ± 2.9</td>
</tr>
<tr>
<td>4</td>
<td>5.63 ± 0.3</td>
<td>10.19 ± 0.7</td>
<td>26 ± 4.3</td>
</tr>
<tr>
<td>5</td>
<td>5.66 ± 0.3</td>
<td>8.59 ± 0.4</td>
<td>8 ± 2.3</td>
</tr>
<tr>
<td>6</td>
<td>5.15 ± 0.3</td>
<td>6.55 ± 0.5</td>
<td>10 ± 2.7</td>
</tr>
<tr>
<td>7</td>
<td>5.63 ± 0.3</td>
<td>8.54 ± 0.5</td>
<td>3 ± 2.1</td>
</tr>
</tbody>
</table>
Table 5. Transfer integral values for chain lengths 3-7, calculated at the B3LYP/6-31G(d,p) level. The dimer types (P, T1, T2) are indicated in the top-view unit cell in Figure 2C.

<table>
<thead>
<tr>
<th>Dimer Type</th>
<th>P / meV</th>
<th>T1 / meV</th>
<th>T2 / meV</th>
</tr>
</thead>
<tbody>
<tr>
<td>3</td>
<td>1.64</td>
<td>4.23</td>
<td>4.20</td>
</tr>
<tr>
<td>4</td>
<td>2.32</td>
<td>7.25</td>
<td>7.25</td>
</tr>
<tr>
<td>5</td>
<td>1.21</td>
<td>0.62</td>
<td>1.02</td>
</tr>
<tr>
<td>6</td>
<td>2.76</td>
<td>4.23</td>
<td>4.34</td>
</tr>
<tr>
<td>7</td>
<td>1.23</td>
<td>2.90</td>
<td>2.76</td>
</tr>
</tbody>
</table>
Figure 1. Images by atomic force microscopy (AFM) on a half monolayer coverage of $C_N$-P2TP-$C_N$, with $N$ ranging from 3 to 8 methyl units in length. A-F depict the layers evaporated on HDTs treated substrates and G-H the layers of ODTs. The non-covered substrate is shown as blue. An increase in grain size (red) with increasing side chain length is observed. A second flat lying phase (black) on ODTs substrates is present but does not form on HDTs treated substrates.
**Figure 2** A. GIXD spectra of half a monolayer of C\textsubscript{3}-P2TP-C\textsubscript{3} and C\textsubscript{4}-P2TP-C\textsubscript{4} on ODTS. Significant differences in Q\textsubscript{z} are visible in the position of the \{11\}, \{02\}, and \{12\} peaks, indicating a structural odd-even effect. B. Best fit of the theoretically calculated diffraction intensities to the experimentally obtained data of half a monolayer of C\textsubscript{3}-P2TP-C\textsubscript{3} and C\textsubscript{4}-P2TP-C\textsubscript{4} on ODTS, respectively. C. Best-fit packing from the calculated GIXD spectra of C\textsubscript{3}-P2TP-C\textsubscript{3} and C\textsubscript{4}-P2TP-C\textsubscript{4} on ODTS. In all cases, the P2TP cores adopt a herringbone packing in the unit cell (see Top-down view of C\textsubscript{3}-P2TP-C\textsubscript{3}). However, while in C\textsubscript{3}-P2TP-C\textsubscript{3} films the P2TP core is perpendicular to the substrate, the P2TP core in C\textsubscript{4}-P2TP-C\textsubscript{4} films is tilted about 23°, mainly in the direction of the b unit cell axis.
Figure 3. Lowest energy position of P2TP on top of ODTS. A $C_3$-P2TP-$C_3$ molecules with a tilt angle of $5^\circ$. B $C_4$-P2TP-$C_4$ under a tilt angle of $19^\circ$.

Figure 4. A comparison of P2TP off-normal tilt angles by simulation, between the presence of an ODTS SAM (blue) and that with the absence of a SAM (red).
Figure 5. The field-effect mobility obtained on transistors with 40 nm thick films of C₆-P2TP-C₆. No clear trend with side chain length is observed within the error of measurement, which is likely due to other aspects that are the main limiting mechanism for charge transport: e.g., grain boundaries and/or the packing of the second and third molecular layer.
REFERENCES

(18) MacGillivray, L. R. CrystEngComm 2004, 6, 77.
(21) Yang, J.; Yan, D. Chemical Society Reviews 2009, 38, 2634.

0.5 monolayer

Molecular tilt angle (deg) vs. alkyl side chain length (N):

- Minimum tilt angle at N = 3 and N = 7
- Maximum tilt angle at N = 4 and N = 6