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Unraveling the Main Chain and Side Chain Effects on Thin Film Morphology and Charge Transport in Quinoidal Conjugated Polymers

Xuncheng Liu, Bo He, Andrés Garzón-Ruiz, Amparo Navarro, Teresa L. Chen, Matthew A. Kolaczkowski, Shizhen Feng, Lianjie Zhang, Christopher A. Anderson, Junwu Chen,* and Yi Liu*

Three series of low-bandgap polymers based on a novel guinoidal para-azaquinodimethane (p-AQM) unit are devised and synthesized, enabling an in-depth study of the impact of structural factors such as polymer main chain, branching point of the side chain, and the length of the branch chains on the thin film morphologies and charge transport properties. Morpho-logical studies reveal that the polymers composed of larger repeating units exhibit a stronger tendency to form edge-on lamella. On the other hand, altering the side chain structures of polymers with the same main chain configuration indicates that the branching point position has a more deter-ministic impact than the branch chain length on the interchain interactions and the crystallite orientation. These results demonstrate a compound odd-even effect of the branching point on the chain packing and morphology, which correlates well with the corresponding field effect transistor perfor-mances. The polymer with the branching point at the fourth carbon displays the highest charge carrier mobility over 1.0 cm² V⁻¹ s⁻¹, concurrent with a bimodal texture. This study provides a comprehensive description of the correlations between polymer structures, thin film morphology, and device performances, providing a clear path to desirable bimodal thin film texture for charge transport.

1. Introduction

Conjugated polymers have received 18 great attention for their application in 19 organic field effect transistors (OFETs),^[1] organic photovoltaics (OPVs),^[2] and other 21 area of flexible electronics.^[3] In order to 22 tune the overall optoelectronic proper- 23 ties of conjugated polymers and their 24 device performances, critical materials design considerations have been under-taken to address both intrachain electron delocalization and interchain packing.^[4] Intrachain control is often implemented 29 by the use of properly chosen electronic 30 building blocks to adjust the molecular 31 orbital levels and polymer main chain 32 conformation. On the other hand, inter-chain packing, which governs ordering, crystallinity, and orientation in thin films. is strongly dependent on both the confor-mation of the polymer main chain and the 37 configuration of side chains.^[2d,5a,b-e]

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43 44 45 46 47 48 49 50 51 52 53 54 55 56 57 58 59	Dr. X. Liu, Dr. B. He, T. L. Chen, M. A. Kolaczkowski, C. A. Anderson, Dr. Y. Liu The Molecular Foundry Lawrence Berkeley National Laboratory One Cyclotron Road, Berkeley, CA 94720, USA E-mail: yliu@lbl.gov Dr. X. Liu, S. Feng, Dr. L. Zhang, Prof. J. Chen Institute of Polymer Optoelectronic Materials and Devices State Key Laboratory of Luminescent Materials and Devices South China University of Technology Guangzhou 510640, P. R. China E-mail: psjwchen@scut.edu.cn Dr. B. He, Dr. Y. Liu Materials Sciences Division Lawrence Berkeley National Laboratory One Cyclotron Road, Berkeley, CA 94720, USA	Dr. A. Garzón-Ruiz Department of Physical Chemistry Faculty of Pharmacy Universidad de Castilla-La Mancha Cronista Francisco Ballesteros Gómez, 1, E02071 Albacete, Spain Dr. A. Navarro Department of Physical and Analytical Chemistry Faculty of Experimental Sciences Universidad de Jaén Campus Las Lagunillas E23071, Jaén, Spain M. A. Kolaczkowski, C. A. Anderson Department of Chemistry University of California, Berkeley Berkeley, CA 94720, USA
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Scheme 1. List of p-AQM-containing conjugated polymers grouped into three series



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study of the main chain effect and side chain effect. Herein, 1 2 we have systematically varied the main chain structure, the 3 branching point, and the length of the side chains. Detailed 4 studies of their thin film morphologies using grazing incidence 5 wide-angle X-ray scattering (GIWAXS) reveal strong correlation 6 between the crystallite orientation and both the main chain and 7 the side chain structures. In particular, a compound "odd-even" 8 effect concerning the branching point of the side chains has been 9 observed and understood by computational modeling. Charge 10 transport studies show that the side chain engineering results in enhanced field effect mobilities over $1.0 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$ in thin films 11 featuring bimodal crystallite orientations. Such studies provide 12 unique insight into the relationship between polymer structures, 13 thin film morphologies, and charge transport properties that will 14 help guide the future design of conjugated polymers. 15 16

¹⁸ 2. Results and Discussion

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20 Three groups of polymers are devised to highlight different 21 structural variations. The first series are polymers with the 22 same 2-decyltetradecoxy side chain but different main chain 23 structures featuring the comonomers of p-AQM and thienylvi-24 nylthiophene (2TV), terthiophene (3T), and quarterthiophene 25 (4T) units (Scheme 1a). A nomenclature PAnT(V)-BCn-CxCy 26 is adopted to underline the different composition of the main 27 chain aromatic structure, the position of the branching point, 28 and the number of carbon on each branch. Accordingly, the 29 corresponding polymers are named as PA2TV-BC2-C10C12, 30 PA3T-BC2-C10C12, and PA4T-BC2-C10C12, respectively. A 31 bithiophene (2T) polymer with slightly shorter branched side 32 chains, PA2T-BC2-C8C10, was also included in the series for 33 comparison (boxed structure in Scheme 1a).

34 The second and third series are polymers with the same main 35 chain structure but bearing different side chains, with the former 36 featuring systematically shifted branching points (Scheme 1b) 37 and the latter featuring significantly different branch chain lengths (Scheme 1c). For the second series, we focus on the con-38 39 jugated polymers containing PA3T repeating units in the main chain since it has exhibited higher carrier mobility than other 40 oligothiophene-based polymers.^[9] Alkoxy side chains with sys-41 42 tematically shifted branching point, such as 3-decyltridecoxy, 4-decyltetradecoxy, and 5-decylpentadecoxy, were introduced 1 to give the series PA3T-BC*n*-C10C10, with *n* being 3, 4, and 2 5 to denote the different position of branching point based 3 on number of carbons, respectively. The polymer PA3T-BC2- 4 C10C12, which is listed in Series 1 and has a branching point at 5 C2 position, is included in Series 2 for comparison despite that 6 the two branched alkyl chains are nonequivalent. Series 3 enlists 7 two conjugated PA4T-BC2 polymers bearing side chains with 8 significantly varied lengths, namely, PA4T-BC2-C10C12 (also 9 listed in Series 1) and PA4T-BC2-C16C18, respectively. 10

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2.1. Poly	mer Sy	nthesis
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14 The synthesis of *p*-AOM monomers and all the polymers follows 15 a modular synthetic protocol^[9] and is included in the Supporting 16 Information. Briefly, the *p*-AQM monomers **S2a-e** were furnished 17 by alkylation of the diarylidene piperazinedione S1^[9] with the cor-18 responding alkyl bromides. Pd-catalyzed cross-coupling polym- 19 erization between S2a-d and the corresponding comonomers 20 afforded the desired polymers. PA2T-BC2-C8C10, PA3T-BC2-21 C10C12, and PA4T-BC2-C10C12 in Series 1 were synthesized 22 previously.^[9] The solubility of polymers decreases as the main 23 chain repeating unit becomes larger, as observed for the Series 24 1 polymers, and increases as the side chain becomes longer for 25 the Series 3 polymers. The solubility and molecular weight of the 26 PA3T-BCn polymers in series 2 show a correlation to an odd-even 27 effect of the branching point. The number-averaged molecular 28 weights of PA3T-BC3-C10C10 and PA3T-BC5-C10C10 are around 29 10 000, while these of PA3T-BC2-C10C12 and PA3T-BC4-C10C10 30 are over 26 000. Despite the higher molecular weight of the latter 31 two polymers, they exhibit considerably higher solubility than 32 PA3T-BC3-C10C10 and PA3T-BC5-C10C10. PA3T-BC2-C10C12 33 and PA3T-BC4-C10C10 are readily soluble in solvents such as 34 chloroform and chlorobenzene at room temperature, while the 35 latter two are difficult to dissolve even at elevated temperature. 36

2.2. Optical and Electrochemical Properties

The UV-vis absorption spectra of these polymers are indic- 41 ative of the low-bandgap characteristics (Figure 1 and 42







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40Figure 2. Typical output and transfer characteristics of OFETs based on a,b) PA3T-BC3-C10C10, c,d) PA3T-BC4-C10C10, and e,f) PA3T-BC5-C10C10.4041(a), (c), and (e) Output curves; (b), (d), and (f) transfer curves.41

Table 1). All polymers except PA3T-BC3-C10C10 show a dualband absorption in chlorobenzene solution. The dual-band feature, corresponding to the 0-0 and 0-1 transitions, is indica-tive of preaggregated states in the solutions of these polymers. A comparison of the three Series 1 polymers with the same side chains shows that the bandgap increases as the number of thiophene increases, consistent with previous findings (Figure 1a). The two polymers in Series 3 show nearly iden-tical absorption in both solution and the thin film, suggesting that different branch chain length has little impact on the optical properties (Figure S1, Supporting Information). For the Series 2 polymers, the 0-0 peak of PA3T-BC4-C10C10 is at the longest wavelength. In contrast, PA3T-BC3-C10C10 has no noticeable aggregation peak in its absorption spectrum, and also its absorption maximum is considerably blueshifted even when compared to PA3T-BC2-C10C12. The trend of the solution optical bandgap changes is again in accordance

with an odd-even effect of the branching point. The abnormal behavior of PA3T-BC3-C10C10 may suggest decreased copla-narity and conjugation of the polymer main chain due to the increased steric hindrance imposed by the specific branching pattern, which is discussed in further details later. A new shoulder peak arises in the thin film absorption spectrum of PA3T-BC3-C10C10, suggesting reinforced coplanarity and stronger chain aggregation in the solid state. The dual band absorption feature for all other polymers is preserved in the corresponding thin film absorption spectra, together with slight redshift of the 0-0 transition peaks, also consistent with enhanced coplanarity and stacking in the solid state. Overall, the optical bandgaps of these polymers are strongly dependent on the main chain structure but much less so on the side chains, as can be seen from the apparent shift of 0-0 transition peaks from 1.53 to 1.36 eV in the spectra of Series 1 polymers, while these for the Series 2 polymers

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Table 1.	Summar	v of the c	optical and	d electrochemical	properties	and fr	ontier orbit	al energies
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Polymer		UV–vis						
	Solution ^{a)}			Film				
	λ_{\max} [nm]	$\lambda_{ m onset}$ [nm]	E _g ^{opt} [eV]	λ _{max} [nm]	$\lambda_{ m onset}$ [nm]	E_{g}^{opt} [eV]	HOMO ^{b)} [eV]	LUMO ^{c)} [eV]
PA2TV-BC2-C10C12	824	896	1.38	834	910	1.36	-5.0	-3.6
PA3T-BC2-C10C12	742	810	1.53	758	832	1.49	-5.0	-3.5
PA4T-BC2-C10C12	731	805	1.54	731	810	1.53	-5.1	-3.5
PA3T-BC3-C10C10	655	802	1.55	745	836	1.48	-4.9	-3.4
PA3T-BC4-C10C10	752	832	1.49	766	858	1.45	-4.9	-3.4
PA3T-BC5-C10C10	748	809	1.53	763	841	1.47	-4.9	-3.4
PA4T-BC2-C16C18	726	805	1.54	726	820	1.51	-5.1	-3.5

15 ^{a)}Measured in chlorobenzene; ^{b)}Calculated using $E_{HOMO} = -(4.8 - E_{ox})$ eV. The E_{ox} was measured by cyclic voltammetry using conventional three-electrode setup at a scan 16 rate of 100 mV s⁻¹; c)Calculated using the corresponding E_{HOMO} and E_g^{opt} . 17

19 are within 1.45 and 1.49 eV based on the absorption edges of 20 the respective thin films. Approximate frontier orbital energy 21 levels^[11] of the polymers were estimated from cyclic voltammetry (Figure S2, Supporting Information) and optical gaps 22 23 and summarized in Table 1. Not unexpectedly, the highest 24 occupied molecular orbital (HOMO) and lowest unoccupied 25 molecular orbital (LUMO) energy levels for the Series 2 poly-26 mers vary only slightly in the range between -5.0 and -4.9 eV 27 and between -3.5 and -3.4 eV, respectively.

2.3. Thin Film FET Measurements

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Charge transport properties of all the polymers were evaluated 33 in thin film field effect transistors, using the bottom gate/top 34 contact OFET configuration with octadecyltrichlorosilane (OTS)-35 modified SiO₂ as the dielectric layer. Gold (Au) was used as the 36 source/drain electrodes, and scratched n-doped Si was used as 37 the gate electrode. The devices were fabricated using as-cast 38 thin films or those thermally annealed at 200 or 225 °C. All the 39 devices displayed distinct p-type transporting behavior. PA2TV-BC2-C10C12 had a hole mobility (μ) of 3.9 × 10⁻³ cm² V⁻¹ s⁻¹, 40 which was lower than that of the other polymers in Series 1 that 41 42 were reported previously (Table S1, Supporting Information).^[9] The two polymers in Series 3 displayed very comparable mobil-43 ities around $8.0 \times 10^{-2} \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$, indicating that the branch 44 45 chain length variation has little impact on the charge transport 46 properties (Table S1, Supporting Information).

For the Series 2 polymers, PA3T-BC3-C10C10 displayed the 47 48 lowest hole mobility among all four polymers, with a maximum 49 and average mobility of 7.0 \times 10⁻³ and $5.0 \times 10^{-3} \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$, respectively (Table 2). 50 51 This value is nearly two orders lower than that 52 of the previously reported PA3T-BC2-C10C12 53 (0.54 cm² V⁻¹ s⁻¹). PA3T-BC5-C10C10 showed 54 a comparable mobility to PA3T-BC2-C10C12 55 of 0.32 cm² V⁻¹ s⁻¹. PA3T-BC4-C10C10 displayed the highest transporting property 56 57 among the series, with the mobility up to 1.05 cm² V⁻¹ s⁻¹ and an average mobility 58 of 0.92 cm² V⁻¹ s⁻¹. The overall trend of 59

mobilities follows an odd-even relationship with respect to the 19 branching positions. Devices based on PA3T-BC4-C10C10 and 20 PA3T-BC5-C10C10 also feature low threshold voltages, which 21 suggest low charge trapping sites in the active layer. 22

2.4. Morphological Studies

GIWAXS measurements were conducted to probe molecular 27 packing within the spun-cast films. Thin films were either as 28 cast or annealed at 200 °C for 30 min unless stated otherwise 29 before being subjected to synchrotron X-ray studies. Greater 30 details regarding the molecular packing and orientation are 31 revealed from the comparative studies of the different series, as 32 itemized below 33

2.4.1. The Main Chain Effect

37 Studies of the Series 1 polymers revealed that for the polymers 38 bearing the same side chain, as the size of the repeating unit 39 becomes larger, the crystallites tend to adopt an edge-on orienta-40 tion over the face-on one. This can be seen from the textures of 41 both the room temperature and the annealed thin films of the 42 polymers. As shown in Figure 3, the as-cast thin films of PA2TV-43 BC2-C10C12 and PA3T-BC2-C10C12 have a bimodal texture, 44 showing the presence of both face-on and edge-on lamellas, while 45 PA4T-BC2-C10C12 has a pure edge-on orientation. The face-on 46 component decreases upon thermal annealing, and completely 47 disappears at 200 °C for PA2TV-BC2-C10C12 and at 250 °C 48

Table 2. Summary of OFET characteristics of Series 2 polymers.

Polymers	$T_{\text{annealing}} [^{\circ} C]$	$\mu_{\rm h}{}^{\rm avg,a)} \; [\rm cm^2 \; V^{-1} \; s^{-1}]$	$\mu_{\rm h}^{\rm max,b)} [{\rm cm}^2 {\rm V}^{-1} {\rm s}^{-1}]$	$V_{\rm th} [V]$	$I_{\rm On}/I_{\rm Off}$
PA3T-BC2-C10C12 ^{c)}	200	0.47	0.54	-14	$5 imes10^5$
PA3T-BC3-C10C10	200	0.005	0.007	-10	$3 imes10^3$
PA3T-BC4-C10C10	225	0.92	1.05	-3	$2 imes 10^5$
PA3T-BC5-C10C10	225	0.26	0.32	-4	$8 imes 10^5$

^{a)}Average mobility; ^{b)}Maximum mobility; ^{c)}Reported previously.

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Figure 3. GIWAXS patterns of thin films of a,b) PA2TV-BC2-C10C12, c,d) PA3T-BC2-C10C12, and e,f) PA4T-BC2-C10C12. (a), (c), and (e): as-cast; 29 (b) and (f): 200 °C; (d) 250 °C. 30

for PA3T-BC2-C10C12, respectively. This is consistent with the 32 literature report that polymers with lower side chain density 33 34 have a stronger tendency of forming edge-on oriented crystal-35 lites due to tighter packing between lamellas.^[12] Tighter packing 36 is clearly observed in this series, as the interlamella distance 37 (d_{100}) for the thermally annealed polymers decreases from 2.63 nm for PA2TV-BC2-C10C12, to 2.23 nm for PA3T-BC2-C10C12 38 39 and to 2.05 nm for PA4T-BC2-C10C12. It is worth noting that

the polymer PA2T-BC2-C8C10 with an even smaller main chain 32 repeating unit (A2T) than that of PA2TV-BC2-C10C12,^[9] despite 33 slightly shorter side chains, adopts a pure face-on orientation at 34 both room temperature and after thermal annealing (Figure S3, 35 Supporting Information). The solid-state packing behavior com-36 plements the range of orientation preferences that changes pro-37 gressively from face-on to bimodal to pure edge-on as the side 38 chain density decreases, as illustrated in Figure 4. 39

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2.4.2. Branching Side Chain Effect

Several notable trends could be summarized from the GIWAXS studies of the Series 2 polymers.

(1) Face-on orientation is observed in the as-cast films of all the PA3T polymers, and the interlamellar spacing in the face-on crystallites increases monotonically as the side chain length becomes larger. In the case of PA3T-BC2-C10C12, it coexists with a significant fraction of edge-on orientation (Figure 5). The interlamellar distances of the face-on and edge-on crys-1 tallites are 2.53 and 2.23 nm, respectively. In contrast, for 2 the polymers with branching point more than two carbons 3 away from the main chain, they all adopt a pure face-on ori- 4 entation, with PA3T-BC3-C10C10 being the least crystalline. 5 The interlamellar *d*-spacings of the face-on crystallites are 6 2.27, 2.60, and 2.71 nm for PA3T-BC3-C10C10, PA3T-BC4-7 C10C10, and PA3T-BC5-C10C10, respectively. It should be 8 noted that the side chains in PA3T-BC2-C10C12 are com- 9 posed of two alkyl branches with two carbon differences, 10



(d), (g), and (j): as-cast; (b), (e), (h), and (k): 100 °C; (c), (f), (i), and (l): 200 °C.





and the overall length of the side chains should be close to
that of P3AT-BC4-C10C10. Indeed, the corresponding faceon interlamellar spacing of PA3T-BC2-C10C12 (2.53 nm) is
larger than that of PA3T-BC3-C10C10 (2.27 nm) and close to
that of PA3T-BC4-C10C10 (2.60 nm).

(2) Edge-on is the thermodynamically more preferred orienta-31 32 tion for all the PA3T polymers except PA3T-BC3-C10C10. 33 As can be seen from Figure 5, PA3T-BC3-C10C10 thin film 34 retains its face-on orientation with low crystallinity after 35 thermal annealing, with no changes of the interchain d-spac-36 ing. This is in contrast to all other polymer films that show 37 clear transition from face-on to edge-on upon annealing to 100 and 200 °C. A mixed texture is observed in the annealed 38 39 thin films of PA3T-BC2-C10C12 and PA3T-BC4-C10C10. The edge-on oriented crystallites are evidenced by the strong out-40 of-plane (n00) peaks (n up to 4) while the out-of-plane (001) 41 peak indicates the coexisted face-on crystallites. In the case 42 of PA3T-BC5-C10C10, the strong out-of-plane (n00) peaks 43 (n up to 4) together with the absence of out-of-plane (001) 44 45 peak clearly indicate a pure edge-on phase and the absence of face-on component. For the 200 °C annealed samples, the in-46 terlamellar *d*-spacings for edge-on crystallites are 2.32, 2.34, 47 48 and 2.49 nm for PA3T-BC2-C10C12, PA3T-BC4-C10C10, 49 and PA3T-BC5-C10C10, respectively (no edge-on for PA3T-50 BC3). On the other hand, the interlamellar d-spacings for 51 face-on crystallites, which remain nearly constant upon an-52 nealing, are 2.53, 2.24, and 2.58 nm for PA3T-BC2-C10C12, 53 PA3T-BC3-C10C10, and PA3T-BC4-C10C10 (no face-on for 54 annealed PA3T-BC5). In both orientations, the interlamellar 55 *d*-spacings increase as the length of the side chains becomes larger. In all cases when both face-on and edge-on oriented 56 57 crystallites are present, the interlamellar *d*-spacing of the 58 face-on is always larger than that of the edge-on crystallites, 59 which is indicative of tighter chain packing in the latter.

The transition from face-on in PA3T-BC3-C10C10 to 26 bimodal in PA3T-BC4-C10C10, and to edge-on in PA3T-BC5-27 C10C10 in thermally treated thin films indicates a preference 28 for edge-on conformation as the branching point moves further 29 away from the polymer main chain. A comparison of the thin 30 films of PA3T-BC*n*-C10C10 (n = 3-5) annealed at 100 °C shows 31 that at this intermediate temperature, both PA3T-BC3-C10C10 32 and P3AT-BC4-C10C10 remain face-on, while PA3T-BC5-33 C10C10 clearly has developed strong edge-on texture (Figure 5). 34 Counting PA3T-BC2-C10H12 in the series, the overall trend 35 observes that, with the exception of PA3T-BC3, the farther away 36 the branching point is from the polymer main chain, the more 37 edge-on it becomes upon thermal annealing, as illustrated in 38 Figure 6. 39

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Tapping-mode atomic force microscopic (AFM) studies of 40 the thermally annealed thin films of the Series 2 polymers 41 revealed additional information about the surface morphology 42 and roughness (Figure 7). PA3T-BC3-C10C10 has a very rough 43 surface with a root-mean-square (RMS) roughness of 19.4 nm. 44 This large roughness is in accordance with its low solubility 45 and inferior mobility. On the other hand, PA3T-BC4-C10C10 46 thin film is very smooth with a roughness of only 1.05 nm and 47 a fibril network structure. In the case of PA3T-BC5-C10C10, it 48 has good film forming ability despite its low solubility, giving 49 rise to a similar fibril network structure with slight larger 50 roughness of 1.53 nm. These features are comparable to 51 PA3T-BC2-C10C12, which also exhibits a fibril network with a 52 roughness of 1.63 nm. 53

2.4.3. Side Chain Length Effect

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As shown in Figure S4 (Supporting Information), both polymers in the Series 3 behave very similarly and adopt an edge-on 59



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Figure 7. AFM images of the thin films of a) PA3T-BC2-C10C12, b) PA3T-BC3-C10C10, c) PA3T-BC4-C10C10, and d) PA3T-BC5-C10C10 after annealing 12 at 200 °C (3.5 μm × 3.5 μm). 13

15 orientation throughout the thermal treatment. The interlamella distance remains constant during the thermal annealing while 16 17 small changes of the diffraction pattern were observed, which 18 indicates more ordered side chain organization upon equi-19 librium. More notably, the interlamella distance varies only 20 slightly from 2.05 nm for PA4T-BC2-C10C12 to 2.08 nm for 21 PA4T-BC2-C16C18. Compared to the significant disparity of the 22 branch chain lengths that differ by six carbon atoms, this vari-23 ation is very insignificant, which suggests less steric crowded-24 ness between lamellas as a result of more void space created 25 by the larger π -surface of PA4T. The very similar interlamella 26 packing between the two polymers is also consistent with their 27 comparable device behavior. 28

30 2.5. Correlations between Mobility and Morphology 31

32 The study of the Series 1 polymer indicates that the thermally annealed PA3T polymer has the highest mobility, the thin film 33 34 of which displays a bimodal texture. On the other hand, com-35 parison of the Series 3 polymer suggests that side chain length 36 has minimal effect on the film morphology and related charge 37 transport. The behavior of Series 2 polymers further indicates that the side chain modulated fluctuation of mobilities of PA3T 38 39 polymers correlates well with the observed morphological trend, 40 which bodes well for the importance of bimodal orientation for facile charge transport. The lowest mobility PA3T-BC3-C10C10 41 42 maintains a face-on orientation throughout the annealing process. All three other PA3T polymers prefer a face-on orientation 43 in the as-cast thin films but display different phase purity upon 44 45 thermal annealing. PA3T-BC2-C10C12 remains a mixed face-on 46 and edge-on texture throughout the annealing (up to 200 °C). PA3T-BC4-C10C10 transitions from a face-on orientation to 47 48 a mixed bimodal texture upon annealing to give the highest 49 mobility. Moving the branching point further away from the 50 polymer main chain led to pure edge-on lamellar stacking after 51 annealing, concomitant with a lower charge transport mobility 52 for PA3T-BC5-C10C10. Such correlations provide consistent experimental evidences that highlight the benefit of developing 53 54 bimodal orientations in the thin films, which facilitate the formation of an effective 3D conduction network for more effi-55 cient charge transport.^[13] 56

57 When taking molecular weight into consideration, we note 58 that there lacks a direct correlation between molecular weight 59 and charge carrier mobility. Polymer PA3T-BC4-C10C10 has a significantly lower M_n than PA3T-BC2-C10C12, yet the mobility 15 of the former is much higher. On the other hand, polymers 16 PA3T-BC3-C10C10 and PA3T-BC5-C10C10 have comparable 17 $M_{\rm n}$ yet the former has a much lower mobility. These results 18 suggest that the charge transport mobility of such series of 19 polymers is less contingent on molecular weight but more so 20 on aggregation order and morphology, which is consistent with 21 previous findings.^[14] 22 23

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2.6. Modeling of the Branching Chain Effect

In the Series 2 polymers, PA3T-BC3-C10C10 appears as an out-27 liner that behaves differently from the rest, alluding to subtle 28 steric factors that affect the chain packing behavior. More 29 detailed DFT modeling of that series of polymer chains was 30 thus conducted to provide more insight into the branching 31 chain effect (see the Supporting Information for more detailed 32 information on the previous conformational study of short oli-33 gomers as well as setting up of the starting structures for the 34 modeling). **Figure 8** shows the most stable structures computed 35 for the titled polymer chains for which two different starting 36 conformations, generated by slight rotation of the prebranching 37 point side chains, were employed as calculation starting point, 38 that is, intercrossed (IC) and separated (SC) conformations (see 39 the Supporting Information for more details). 40

In polymer PA3T-BC2, the short prebranching point side 41 chains avoid significant steric hindrances between postbranching 42 point chains. Hence, both the starting points SC and IC led to 43 similar molecular conformations with close energies (see dihe-44 dral angles defined in Figure 9 and relative energies shown in 45 Table 3). The polymer main chain is not fully planar and the 46 calculations yielded a value of 20° for the dihedral angle $|\tau_1|$. Dihe-47 dral angles, $|\tau_2|$ and $|\tau_3|$, between subsequent thiophene rings 48 amount to 166°-177°. In general, similar conformations were 49 observed for the rest of polymer chains studied here. Notably, the 50 postbranching point chains of PA3T-BC2 are oriented out of 51 the plane of the polymer main chain (Figure 8). In contrast, the 52 postbranching point chains are oriented toward the cofacial area 53 of the polymer main chain in both conformers of PA3T-BC3 54 (Figure 8). The change of orientation in the postbranching point 55 chains is reflected in the values of the dihedral angles $|\tau_4|$ which 56 are within 119°-130° for PA3T-BC2 and 17°-51° for PA3T-BC3. 57 The orientation of the postbranching point chains in PA3T-BC3 58 can hinder the π -stacking interactions between neighbor polymer 59





Figure 8. Infinite polymer chains optimized for PA3T-BC2 (conformer
 IC), PA3T-BC3 (conformer IC), PA3T-BC4 (conformer IC), and PA3T-BC5
 (conformer SC) at the B3LYP/3-21G* level of theory. The repeat unit is
 highlighted on the polymer chain. The translation vector of the periodic calculation is shown in red.

39 chains, reducing the charge-hopping-based carrier transport 40 mobility. In the bulk polymer, postbranching point chains gen-41 erally can fold to accommodate the 3D structure. In the case of 42 PA3T-BC3, however, the branching point (the zone with larger 43 density of atoms of the side chain) is near the polymer main 44 45 chain and significant steric hindrance can always be expected. The branch chains in PA3T-BC4 behave similarly as these in 46 PA3T-BC2, with the postbranching point chains orienting out of 47 48 the plane of the polymer main chains as indicated by the large 49 dihedral angle ($|\tau_4| = 109^\circ - 141^\circ$), from which reduced steric inter-50 actions between neighbor polymer chains are expected (Figure 8). 51 Structural conformations calculated for PA3T-BC4 and PA3T-BC3 52 and their implication on the interchain interactions are consistent 53 with the highest and lowest charge mobilities observed for these 54 compounds, respectively. In the case of PA3T-BC5, the post-55 branching point chains are again oriented toward the polymer 56 main chain, as in the case of PA3T-BC3, however the branching 57 point is further away from the polymer chain than for the PA3T-58 BC3. As shown in Figure 8, the side chains of PA3T-BC5 could 59 induce certain hindrance in between polymer main chains,



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Figure 9. Repeating unit of a polymer chain showing the dihedral angles selected to analyze the molecular structure of the studied compounds. The values calculated for these angles are collected in Table 3.

however this steric effect is alleviated by the distant position of 26 the branching point from the polymer main chain that allows the 27 folding of the side chains. The polymer main chain of PA3T-BC5 28 29 is also the most planar of the series on the basis of $|\tau_{1-3}|$ values since the large separation of the branching point reduces the 30 steric effect on the polymer chain. The computation results out-31 line a compound odd-even effect: at odd or even branching point, 32 the postbranching point chains are oriented toward or away from 33 the polymer chain. In the former cases, the steric hindrance is 34 disruptive to interchain packing. However, as the branching point 35 is further moved away, the steric effect is alleviated and becomes 36 less disruptive. PA3T-BC3 thus experiences the most hindrance 37 among all and is the least well-packed in the thin film. 38

3. Conclusion

We have carried out a comprehensive study on the structure- 43 property relationships of several low-bandgap polymers based on 44

Table 3. Values calculated at the B3LYP/3-21G* level of theory for the
dihedral angles (in degrees) shown in Figure 9. These angles were
computed for infinite polymer chains with C10 side chains beyond
the branching point. E_R (in kcal mol⁻¹) is the relative energy between
the conformer IC (intercrossed chains) and the conformer SC (separated
chains) for each polymer and calculation method ($E_R = 0$ corresponds to
the most energetically stable conformer).45
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Polymer	Conformer IC					Conformer SC				
	E _R	$ \tau_1 $	$ \tau_2 $	$ \tau_3 $	$ \tau_4 $	E _R	$ \tau_1 $	$ \tau_2 $	<i>T</i> ₃	$ \tau_4 $
BC2	0.00	20	166	177	119–130	0.21	20	167	176	125–127
BC3	0.00	4	178	175	17–45	1.94	22	167	174	30–51
BC4	0.00	20	171	171	121–132	2.65	17	172	172	109–141
BC5	-	-	_	-	-	-	0.9	172	168	29–35

Adv. Funct. Mater. 2018, 1801874



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Conflict of Interest

The authors declare no conflict of interest.

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3 devising several series of polymers that allow for independent 4 investigation of the main chain effect and two side chain effects 5 involving both the branching point and the chain length. In the 6 PAnT-BC2 series, the polymers bearing the same 2-decyltet-7 radecoxy side chains but with increasing size of the repeating 8 unit in the polymer main chain have a stronger tendency to 9 form edge-on oriented lamellas in the as-cast thin films. Upon 10 thermal annealing, the face-on fraction is significantly reduced to 11 give pure edge-on lamellas. For the PA3T-BCn (n = 3-5) series of polymers that consist of the same conjugated backbone but with 12 systematically varied branched alkoxy side chains, the branching 13 point of the side chains has a dramatic impact on the interchain 14 interactions as well as the orientation of the crystallites on the 15 substrate. A face-on orientation is observed in the as-cast thin 16 17 films of all four PA3T polymers. Thermal annealing induces ori-18 entation transition from face-on to edge-on lamellas at different 19 degrees, varying significantly across the differently branched 20 polymers and is clearly related to the position of the branching 21 point. When the branching point is moved away from the polymer main chain, the polymers are more inclined to transform 22 23 from face-on to edge-on. Once beyond this point, the polymer chains have a stronger tendency to form edge-on lamellas with 24 25 no face-on component. These results demonstrate a compound 26 odd-even effect of the branching point, which amounts to a 27 subtle steric effect as illustrated using detailed computational 28 modeling. We have further shown that the length of the branch 29 chains has little effect on the lamellar packing and the charge transport properties. Among these polymers, PA3T-BC4-C10C10, 30 31 the one with branching point at the fourth carbon away from the main chain has the highest charge carrier mobility up to 32 33 1.05 cm² V⁻¹ s⁻¹ in bottom gate, top-contact OFET devices.

a novel *p*-AQM quinoidal unit, with further insight provided by

detailed theoretical modeling. This study was implemented by

34 The high carrier mobility highlights the great potential of 35 p-AQM-based quinoidal conjugated polymers as novel high-perfor-36 mance organic semiconductors. On the other hand, the revealed 37 correlation between the polymer structures (main chain structure, 38 the side chain branching point, and the length of the branched 39 chains), the thin film morphology, and device performances will 40 serve as design principles to guide the development of high-per-41 forming organic semiconductors for anisotropic charge transport. The bimodal orientation in the highest performing polymer also 42 43 highlights the importance to strike a balance between the two orientations:^[15] while the edge-on oriented crystallites appear to be 44 45 more favorable for in-plane charge transport, in semicrystalline 46 polymer thin films there are abundant lattice defects and disor-47 dered aggregates that throttle the charge carrier movement. A 48 mixed orientation is needed in order to form a 3D conduction net-49 work for optimal charge transport. While this study identifies dis-50 cernible and consistent trends for predictive design and control of 51 morphologies in conjugated polymer films, further studies should 52 be directed toward revealing details of the interconnects between 53 domains with different π -stacking orientations.^[5b]

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56 57 Supporting Information

58 Supporting Information is available from the Wiley Online Library or 59 from the author.

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