



Cite this: *J. Mater. Chem. C*, 2025, 13, 16590

Manipulation of intramolecular hydrogen bonds in conjugated pseudoladder polymer for semiconductivity and solution-processability†

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The conformational coplanarity and local rigidity of π -conjugated backbones are critical for the semiconducting performance of organic electronic materials. While fusing the aromatic system into a ladder-type structure effectively enhances these properties, it also often results in poor solution processability and hence limits their transition to device application. To address this challenge, an intramolecularly hydrogen-bonded pseudoladder polymer (HPLP) system based on alternating hydrogen bond donating benzobisimidazole (BBI) and hydrogen bond accepting benzodifuran (BDF) units, is designed and synthesized. A Boc-protected precursor of HPLP allows for feasible solution processing of the polymer into thin films. Subsequently, *in situ* thermal Boc-deprotection generates the HPLP polymer, in which intramolecular hydrogen bonds form between each pair of neighboring repeating units, inducing coplanarity and rigidity throughout the entire backbone. This process is accompanied by a significant red-shift of the absorption spectrum, reduced bandgap, and enhanced rigidity, as confirmed by NMR, UV-Vis, and density functional theory analyses. HPLP films exhibit a three-order-of-magnitude enhancement in charge carrier mobility compared to the Boc-protected precursor and demonstrate excellent solvent resistance in organic thin-film transistors.

Received 31st March 2025,
Accepted 4th July 2025

DOI: 10.1039/d5tc01365h

rsc.li/materials-c

Introduction

The conformational coplanarity and local rigidity of π -conjugated backbones have been identified as two critical factors for the semiconducting performances of organic electronic materials.^{1–3} A coplanar and rigid backbone often facilitates charge transport through enhanced intramolecular π -conjugation, stronger intermolecular electronic coupling, and lower reorganization energy during charge transfer. A common strategy to induce rigid coplanar conformation is to employ a fused-ring, ladder-type backbone constitution, which restricts torsional rotation between polymer subunits. This

strategy is utilized, for example, by installing a second strand of covalent bonds along the backbone of conjugated polymers,^{4–8} affording conjugated ladder polymers as in the case of **BBL**⁹ and **BTI**¹⁰ (Fig. 1(a)). However, the application of these rigid, coplanar ladder polymers often presents challenges, primarily due to their poor solution processability, which hinders the fabrication of high-quality thin films for device applications. This represents a major dilemma in developing and optimizing semiconducting polymers that possess both excellent electronic performance and processability.^{11–13}

A widely used approach to address the aforementioned challenge involves the incorporation of weak intramolecular interactions in conjugated polymers to produce high performance semiconductors,¹⁴ such as **DPPT-TVTOET**, **PTTz**, and **F-P3HT** (Fig. 1(c)).¹⁵ These weak interactions include non-classical hydrogen bonds (CH–N, CH–O, CH–F) that are in the range of 1–2 kcal mol^{–1} and heteroatom interactions (S–S, S–O, S–N, S–F) falling below 1 kcal mol^{–1}.^{16,17} This strategy induces a coplanar conformation without forming an overly rigid, poorly soluble ladder-type backbone, and has enabled the development of high-performance, solution-processable polymers for electronic and photovoltaic applications.^{18–21} Meanwhile, conventional hydrogen bonds have been proposed as an ideal way

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† Electronic supplementary information (ESI) available. CCDC 2422384 and 2422385. For ESI and crystallographic data in CIF or other electronic format see DOI: <https://doi.org/10.1039/d5tc01365h>



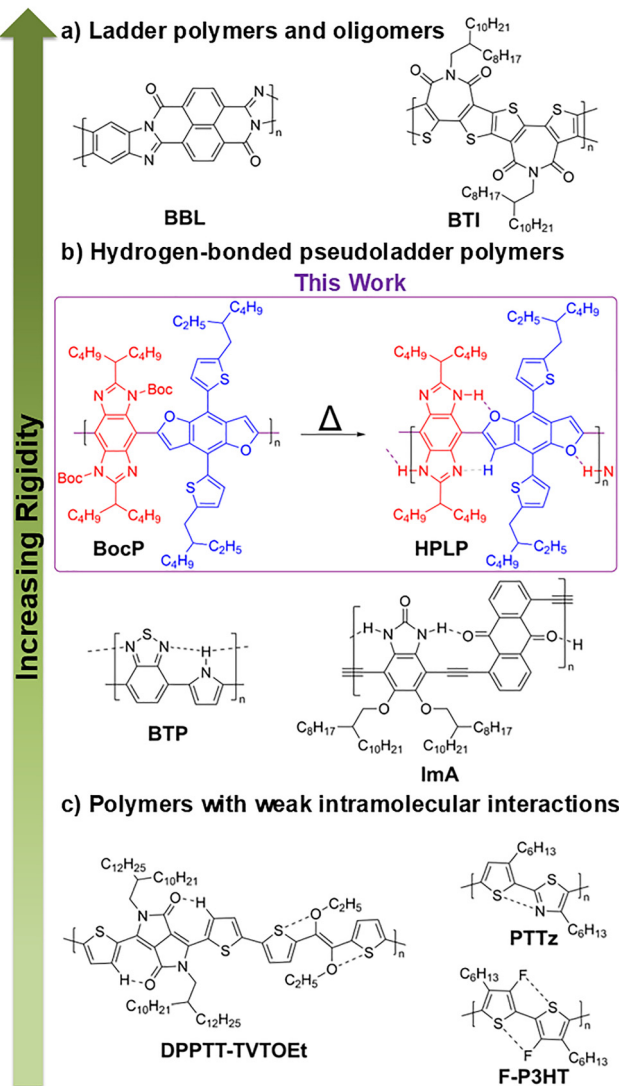


Fig. 1 Structural formulas of semiconducting polymers with coplanarity induced by (a) covalent ring-fusing, (b) hydrogen-bonds, and (c) weak non-covalent interactions.

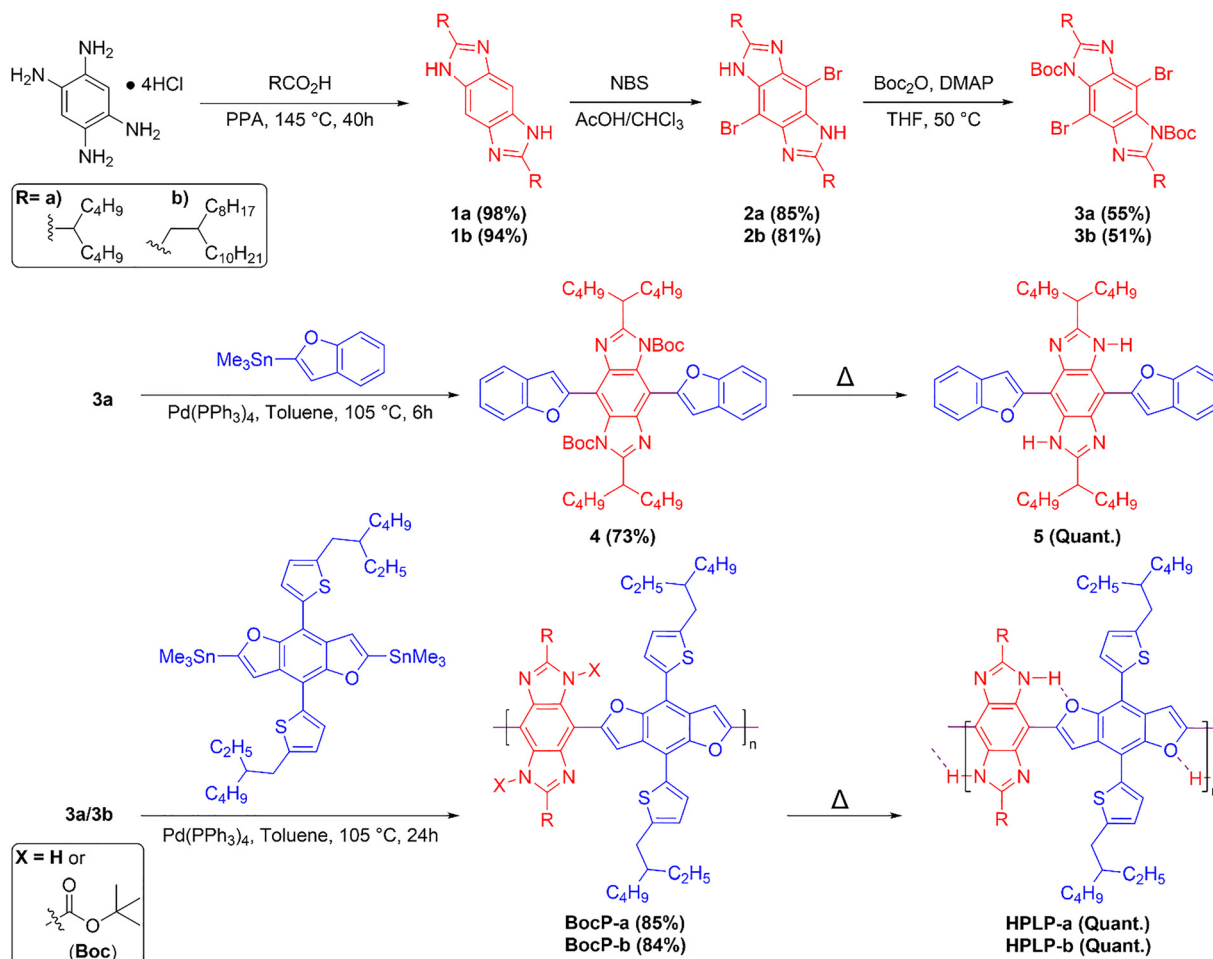
to achieve stronger noncovalent interactions ($2\text{--}10\text{ kcal mol}^{-1}$) in the pursuit of a pseudoladder polymer structure that possesses a more well-defined coplanar conformation.^{22\text{--}25} In addition, hydrogen bonds can be manipulated through masking and unmasking by simple chemical transformations, such as protection with a *tert*-butyloxycarbonyl (Boc) group and corresponding high-temperature deprotection.^{26,27} These features make it possible to achieve highly rigid, coplanar, and solvent-resistant pseudoladder polymers without sacrificing solution processability, achieved through a precursor in which hydrogen bonds are masked. Meijer and coworkers pioneered the incorporation of intramolecular hydrogen bonds in conjugated macromolecules by constructing pseudoladder structures consisting of alternating 1,4-phenylenediamide and 1,4-pyrazine repeating units,²² as well as oligomers consisting of benzothiadiazole and pyrrole repeating units (Fig. 1(b), **BTP**).²³ More recently, the Zhao²⁸ and Fang²⁹ groups have each

explored this approach on acetylene-linked conjugated backbones (Fig. 1(b), **ImA**). It has been observed that fusing hydrogen bond-donating and accepting units into the backbone is essential for establishing robust intramolecular hydrogen bonding, due to the more favorable entropy contribution in such systems. Although the semiconducting performance of these materials has not been reported, the early work of Meijer notably shows a high conductivity of 1 S cm^{-1} achieved in the iodine-doped thin films of **BTP** oligomers.²³ Recent efforts to implement this molecular design strategy into semiconducting systems have demonstrated promising performance in small molecules³⁰ and non-ladder conjugated polymers.^{24,25} However, a semiconducting pseudoladder polymer featuring its backbone entirely locked by intramolecular hydrogen bonds has yet to be reported. Herein, we present such a hydrogen-bonded pseudoladder polymer (**HPLP**) system, which is solution-processable through the manipulation of latent intramolecular hydrogen bonds. This system exhibits desired semiconducting electronic properties and solvent resistance in device tests, owing to the rigid and coplanar backbone induced by these intramolecular hydrogen bonds.

Results and discussion

The hydrogen-bonded pseudoladder polymer **HPLP** (Scheme 1) is composed of alternating benzobisimidazole (BBI) units as the hydrogen bond donors and benzodifuran (BDF) units as the hydrogen bond acceptors. This design allows the formation of a completely 'pseudoladder' constitution through a sequence of uninterrupted intramolecular hydrogen bonds between all neighboring conjugated units along the polymer backbone. The BBI unit was selected for this work for multiple reasons. First, the hydrogen-bond donating imidazole moieties are fused to the aromatic backbone, so that the resulting hydrogen bonds can form with a relatively low entropy penalty, rendering it strong and robust even at elevated temperatures. Second, BBI is feasible to synthesize *via* a simple condensation reaction and versatile for the attachment of various functional groups and solubilizing side chains.^{31\text{--}33} Third, the BBI unit can be easily masked pre-polymerization using Boc protecting groups and unmasked post-polymerization through heat-induced Boc-deprotection. Finally, BBI has been extensively utilized in the construction of conjugated small molecules and polymers,^{34\text{--}43} as well as ladder polymers^{10,11} which demonstrate unique optoelectronic properties and outstanding stability. To couple with BBI, a benzofuran-derived, hydrogen-bond accepting comonomer was chosen on account of its synthetic feasibility, and previously reported good performance in organic photovoltaics.⁴⁴ It is worth mentioning that the use of comonomers containing nitrogen centers as the hydrogen bond accepting sites, such as pyridine or pyrazine-fused derivatives, did not yield the desired synthetic outcome due to their negative impact on metal-catalyzed cross-coupling reactions and their tendency to promote undesired Boc-deprotection of neighboring hydrogen bond donors.





Scheme 1 Synthesis of benzobisimidazole-derived monomers **3a** and **3b**, small molecular models **4** and **5**, polymer precursors **BocP-a** and **BocP-b**, and pseudoladder polymers **HPLP-a** and **HPLP-b**.

In order to optimize the synthetic conditions and to instruct the characterization of the desired polymeric product, a small molecular model compound **5** was designed and synthesized prior to the synthesis of **HPLP** (Scheme 1). First, benzobisimidazole **1a** and **1b** are prepared in over 94% yields through simple condensation reactions of 1,2,4,5-tetraminobenzene hydrochloride salt with a carboxylic acid of choice (2-butylhexanoic acid for **1a** and 3-octyltridecanoic acid for **1b**). Bromination of **1a** and **1b** afforded the corresponding intermediates **2a** and **2b**. A mixed solvent of chloroform and acetic acid is needed here to fully dissolve **1a** and **1b** due to their poor solubility in most organic solvents. Although **2a** and **2b** seem suitable for cross-coupling, their poor solubility and the presence of N-H hydrogens prevented efficient conversion during our initial attempts of direct cross-coupling reactions. Therefore, **2a** and **2b** were subjected to Boc-protection, which proceeds quickly at 50 °C and converts the insoluble starting material suspended in tetrahydrofuran into a homogenous solution of the desired product. Note that two isomers are possible as the result of *syn*- and *anti*-protection of the two imidazole moieties. The more favorable *anti*-protected product **3a** or **3b** was isolated in good yields (55% and 51%, respectively).

Subsequently, trimethyltin-functionalized benzofuran was used as the coupling partner to react with **3a** *via* palladium catalyzed Stille reaction to afford small molecule model **4**, in which the hydrogen bonds are masked by Boc groups. In parallel, to synthesize the target polymer, Stille coupling reactions of **3a** or **3b** with a benzodifuran comonomer functionalized with two trimethylstannyl groups was conducted to give polymer **BocP-a** or **BocP-b**, respectively, in high efficiency. It is noteworthy that the Boc groups were partially removed during the Stille reaction, although both the resulting products **BocP-a** and **BocP-b** remained soluble in common organic solvents. The number average molar mass (M_n) and polydispersity (D) of both polymers were measured on an analytical gel permeation chromatography that was calibrated by polystyrene standards (Fig. S17 and Table S1, ESI[†]). **BocP-a** showed a measured M_n of 11.5 kg mol⁻¹ and D = 2.15; while **BocP-b** had a higher M_n of 28.6 kg mol⁻¹ and D = 1.98. Note that **BocP-a** is decorated with much smaller 1-butylpentyl side chains compared to the 2-octyldodecyl side chains of **BocP-b**, resulting in the molar mass obtained for **BocP-a** being considerably lower than that of **BocP-b** due to solubility limitations for longer polymer chains during the reaction. **BocP** polymers were purified and separated



into fractions of different molar masses (Table S1, ESI[†]) on a preparative gel permeation chromatography to be used for various analyses such as nuclear magnetic resonance (NMR) and UV-Vis spectroscopy, cyclic voltammetry (CV), and scanning tunneling microscopy (STM). This was done to ensure lower polydispersity, and to accommodate the different solubility requirements for each mode of analysis.

Thermogravimetric analysis of compound **4** and **BocP** are marked by a significant loss in weight percent between 180–200 °C corresponding to Boc cleavage with no more significant decomposition occurring until the temperature exceeded 350 °C (Fig. S18 and S19, ESI[†]). These observations assure that thermal treatment at 180 °C under an inert atmosphere does not result in significant decomposition of the molecular backbone other than Boc cleavage. Indeed, synthetically, the Boc groups in **4** can be removed simply by heating at 180 °C for 1 h under a nitrogen atmosphere to afford the unmasked product **5** in quantitative yield. Quantitative conversion of **BocP** to the desired hydrogen-bonded pseudo ladder polymer **HPLP** was also achieved under the same conditions. The ¹H NMR (Fig. 2(a)) and IR (Fig. S15 and S16, ESI[†]) spectra clearly illustrate the presence of Boc groups in **4** and **BocP-a**, and the cleavage of Boc groups in **5** and **HPLP-a**. Due to the partial cleavage of Boc groups in **BocP-a** during Stille polymerization, there is a weak peak at ~10.7 ppm in the ¹H NMR spectrum and a weak peak at 3417 cm⁻¹ in the IR spectrum, respectively, which correspond to the unmasked N-H group. After thermal Boc cleavage, a higher intensity peak of the unmasked

intramolecular hydrogen-bonded N–H group is observed in both ¹H NMR and IR spectra.

Overall, Boc-protected polymers **BocP-a** and **BocP-b** exhibited good solubility in common organic solvents such as toluene, chlorobenzene, chloroform, and tetrahydrofuran. Upon Boc-deprotection, however, the solubility of **HPLP** polymers was greatly decreased (Table S2, ESI[†]). It should be noted that the well resolved ¹H NMR peaks observed for **HPLP-a** in Fig. 2(a) were obtained using a sample of low molar mass (**BocP-a**: $M_n = 7.3 \text{ kg mol}^{-1}$, $D = 1.83$) due to the poorer solubility of higher polymer chains. For **HPLP-a** and -**b**, this low solubility was taken as an advantage to fabricate solvent-resistant thin films, by solution-processing the soluble precursor **BocP-a** or -**b** followed by *in situ* thermal Boc-cleavage, yielding thin films of **HPLP** exhibiting excellent resistance against a variety of organic solvents (Table S2 and Fig. S21, ESI[†]). This unique property renders this system highly promising for fabricating solvent-resistant functional polymer thin films for devices that are durable under strong organic solvents and suitable for multi-layer solution processing techniques.

Boc-cleavage also results in significant changes to the optical properties and electronic structures of the designed conjugated systems (Table 1). Conversion of **4** to **5** resulted in a 45 nm redshift in λ_{max} accompanied by an astounding change in absorption features from one broad peak in **4** to well-defined vibrational peaks for **5** (Fig. 2(b)). The redshift, *i.e.* decrease of optical bandgap, indicates better π -conjugation between the repeating units due to the more coplanar conformation, while

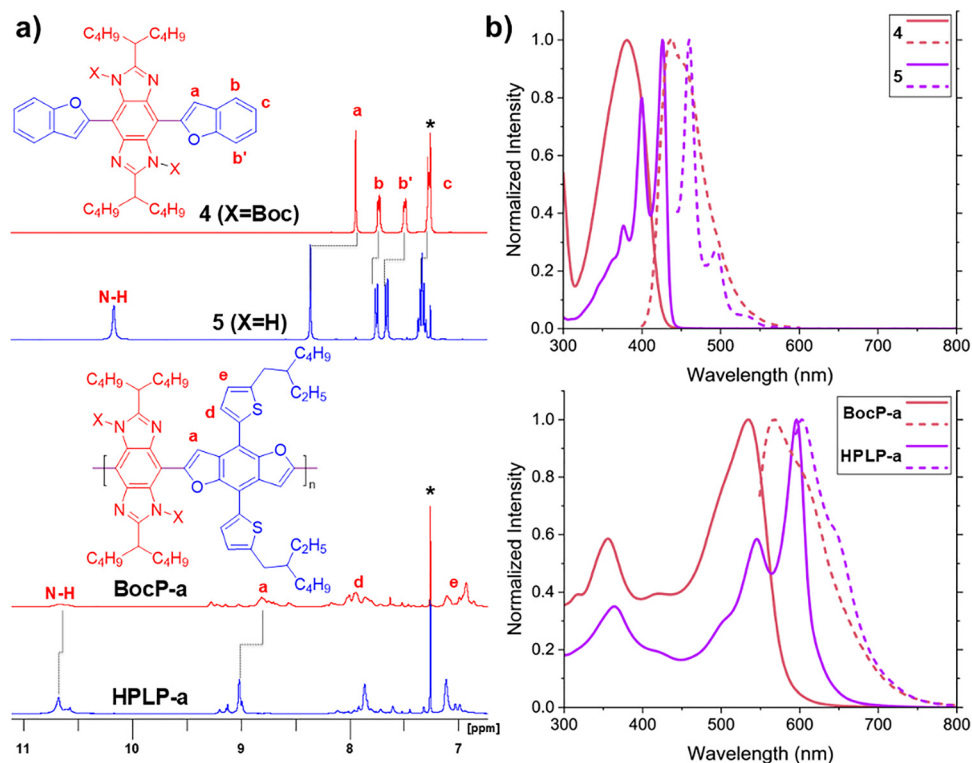


Fig. 2 (a) ¹H NMR spectra and of the Boc-masked species **4** and **BocP-a** ($M_n = 7.3 \text{ kg mol}^{-1}$, $D = 1.83$) and (b) UV-Vis (solid line) and fluorescence (dashed line) spectra of **4** and **BocP-a** ($M_n = 16.1 \text{ kg mol}^{-1}$, $D = 1.42$) in comparison with the corresponding unmasked derivatives.



Table 1 Optoelectronic data of Boc-protected compounds **4** and **BocP**, in comparison with their hydrogen-bonded counterparts **5** and **HPLP**, respectively

Compound	λ_{\max} (nm)	Em_{\max} (nm)	Δ (nm)	HOMO_{CV} (eV)	LUMO_{CV} (eV)	E_{gCV} (eV)
4	381	436	55	-5.40	-3.58	1.82
5	426	460	34	-5.15	-3.56	1.59
BocP	534 ^a	569 ^a	35	-5.71 ^b	-3.54 ^b	2.17
HPLP	596	602	6	-4.96	-3.58	1.38

^a **BocP-a:** ($M_n = 16.1 \text{ kg mol}^{-1}$, $D = 1.42$). ^b **BocP-b:** ($M_n = 13.5 \text{ kg mol}^{-1}$, $D = 1.30$).

the narrowed vibrational peak indicates the local rigidity of the π -system has increased drastically. Similar changes in the UV-Vis and fluorescence spectra peak patterns between **BocP-a** and **HPLP-a** were also observed along with an increased redshift of 62 nm and a decreasing stoke shift (Δ) from 35 to 6 nm, respectively (Fig. 2(b) and Table 1). Although boc-protected precursors show similar absorption onsets to their hydrogen-bonded counterparts indicating similar band gaps, cyclic voltammetry showed significant changes to HOMO energy levels and small changes to LUMO levels. The similar absorption onset is likely a result of relatively coplanar conformations in boc-protected precursors being present in low concentrations, whereas their hydrogen bonded counterparts exist in mostly coplanar conformations as indicated by their sharp vibration peak pattern and λ_{\max} lying nearer to their respective onset absorptions. To obtain accurate CV measurements in solution, a lower molar mass sample of 2-octyldodecyl-decorated **BocP-b** ($M_n = 13.5 \text{ kg mol}^{-1}$, $D = 1.30$) and **HPLP-b** were investigated as the model polymers due to the improved solubility of **HPLP-b** in tetrahydrofuran compared to that of **HPLP-a**. CV measurements (Fig. S29 and S30, ESI[†]) demonstrate a significantly decreased bandgap of **5** (1.59 eV) and **HPLP-b** (1.38 eV) compared to **4** (1.82 eV) and **BocP-b** (2.17 eV), respectively (Table 1). The decreased bandgaps in both **5** and **HPLP** is primarily attributed to the increase of the HOMO level after removal of the electron-withdrawing Boc groups and the coplanar conformation

induced by the intramolecular hydrogen bonds, with little change to the LUMO being observed.

To gauge the strength of these intramolecular hydrogen bonds and the coplanarity of the designed conjugated system, torsional analysis was performed using density functional theory (DFT) on a BBI-benzofuran model molecule by rotating about the bridging single bond in 15° intervals and recording free energy at B3LYP/6-31G(d) level of theory (Fig. 3(a)). Results showed rotational barriers of 9.6 kcal mol⁻¹ and a calculated coplanarity parameter⁴⁵ of $\langle \cos^2(\varphi) \rangle = 0.93$. This result suggested that intramolecular hydrogen bonding interactions lead to energetically favorable coplanar conformations in the gas phase. Single crystals of compounds **4** and **5** were grown for X-ray diffraction analysis (Fig. 3(b) and (c)). Despite its sterically demanding Boc groups, **4** demonstrates a fairly small dihedral angle of 21° between the BBI unit and the benzofuran unit (Fig. 3(a), bottom). This dihedral angle is likely decreased by the weak C-H···N hydrogen bonding interactions between 2-position of the furan ring and the basic nitrogen center on the BBI unit, evidenced by the short C···N distance of 3.15 Å falling within traditional hydrogen bonding distance. In the unmasked molecule **5**, a smaller dihedral angle of 13° was observed between the BBI unit and the benzofuran. The short distances of O-N (2.85 Å) and C-N (3.04 Å) confirm the formation of two hydrogen bonds N-H···O and C-H···N on both sides of the covalent C-C bond. It is interesting to observe that **5** exhibits a non-zero dihedral angle in the solid-state structure, while gas-phase DFT calculations suggest that the dihedral angle should be zero. This discrepancy is likely due to strain induced by intermolecular interactions in the solid state, primarily the intermolecular interaction between the furan oxygen and the BBI's N-H unit that competes against intramolecular hydrogen bonding. This interaction is evident from the relatively short intermolecular O-N distance of 3.3 Å in the crystal structure (Fig. 3(d)). Further evidence for this interaction is provided by an endothermic peak observed at ~200 °C during differential scanning calorimetry (DSC) of single crystals of **5** (Fig. S19b, ESI[†]). This peak does not appear in the second

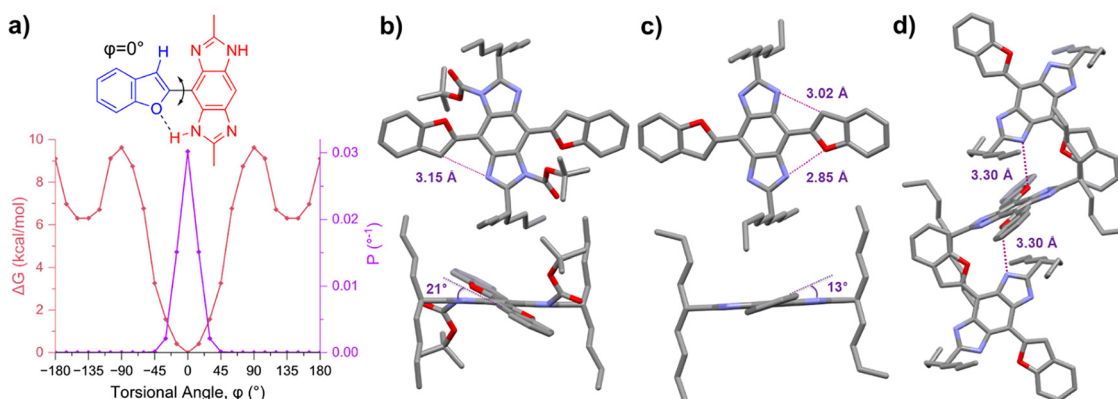


Fig. 3 (a) Relative energy (ΔG) and probability density (P) obtained through torsional analysis by rotating 15° across the bridging C-C bond of a benzobisimidazole-benzofuran system using density functional theory at B3LYP/6-31G(d) level of theory; (b)–(d) front view (top) and side view (bottom) of single molecule crystal structures of **4** and **5**, as well as crystal packing of **5** showing intermolecular hydrogen-bonding interactions.



scan of the same sample following a melt-solidify cycle or in a powder sample of 5 (Fig. S19a, ESI[†]). Considering the solution-phase conformation of 5 and the polymers **HPLP**, it is believed that such intermolecular interactions are not dominant, allowing the intramolecularly hydrogen-bonded molecule to adopt a near-zero dihedral angle in solution as suggested by DFT calculations.

Supporting the DFT calculations, variable temperature-NMR spectra of **HPLP-a** show little change from 20 to 100 °C (Fig. 4(a)), and variable temperature UV-Vis spectra of its thin films show only a slight blue shift when taken to 250 °C (Fig. 4(b)). Similar behavior was observed in the solution-phase UV-Vis analyses of **HPLP-a** and **HPLP-b** (Fig. S27 and S28, ESI[†]), as well as NMR and UV-vis spectra of small molecule analog **5a** (Fig. S13 and S26, ESI[†]). The high-temperature robustness of intramolecular hydrogen bonds in 5 and **HPLP-a** can be attributed to the unfavorable entropy change for breaking these hydrogen bonds, due to the additional solvation of the hydrogen-bond donor and acceptor sites once they are dissociated. These results suggest that, although not as strong as the covalent bonds used in traditional ladder polymers,

these intramolecular hydrogen bonds can still be used for the same purpose of locking conjugated polymers into a highly robust coplanar rigid conformation.

Both **BocP-b** ($M_n = 12.8$, $D = 1.5$) and **HPLP-b** polymers were analyzed by ultrahigh vacuum STM after having been deposited as monolayer films in vacuum by electrospray deposition onto an atomically clean, flat Au(111) surface (Fig. S32 and S33, ESI[†]).^{46–48} In both cases, the polymers appear as extended linear structures with few bends. No substantial difference could be observed between **BocP-b** and **HPLP-b**, likely due to the planarizing effect of face-on surface adsorption on the backbone conformation. In areas of higher coverage (Fig. 5), the polymers tend to self-assemble into locally compact arrangements by aligning their backbones parallel to each other. The average distance between polymers, $d = 3.5 \pm 0.5$ nm, is compatible with a packing model in which the side chains of adjacent polymers interlock with a minimal interdigitation (Fig. S33, ESI[†]), similar to what has been observed in other conjugated polymers with a high density of side chains attached to the backbones.⁴⁶ Polymer crossings are occasionally observed at near-perpendicular angles (Fig. 5 and Fig. S32a, ESI[†]), a feature that was recently highlighted as beneficial for promoting a highly interconnected charge carrier transport network, thereby enabling high charge transport mobility.^{47,48} In addition, the molar mass of the **BocP-b** sample was evaluated by measuring the lengths of 607 individual chains identified in the STM images. The statistical average molar mass and its distribution closely align with the results obtained from GPC (Fig. S34, ESI[†]). This finding demonstrates the powerful capability of STM in accurately measuring and calibrating the molar mass of rigid conjugated polymers.

To test the effect of hydrogen bond-induced coplanarity on electronic performance, organic thin-film transistors composed of **BocP-a** ($M_n = 16.1 \text{ kg mol}^{-1}$, $D = 1.42$) and **HPLP-a** generated *in situ* from **BocP-a** are tested (Fig. 6). A bottom-gate/top-contact device architecture is employed. To fabricate the devices, **BocP-a** polymers were spin-coated onto *n*-octadecyltrichlorosilane (OTS)-treated SiO₂/Si substrates, while gold was deposited as the source and drain electrodes, detailed in the ESI.[†] The as fabricated device of **BocP-a** shows a low hole mobility of

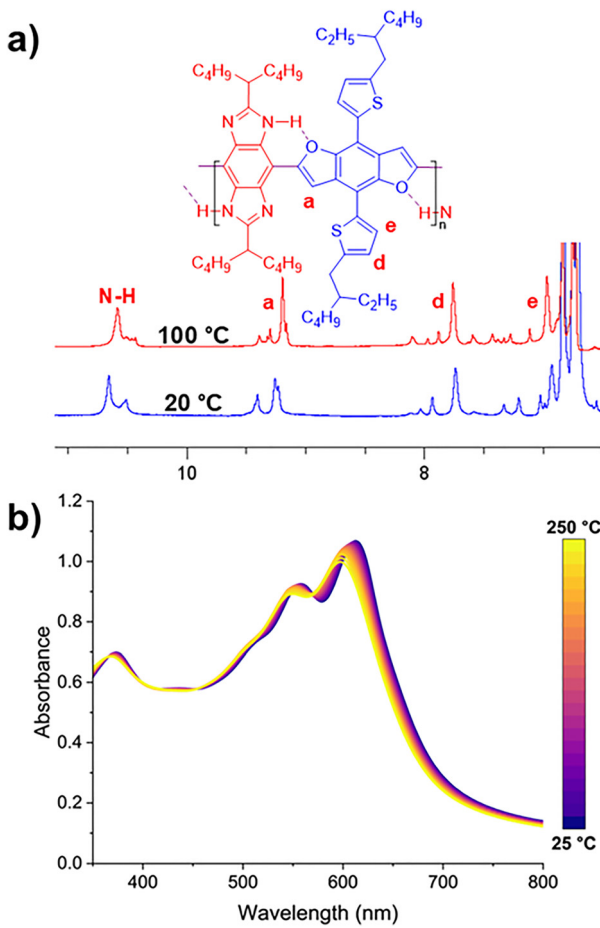


Fig. 4 (a) Variable-temperature NMR of **HPLP-a** (derived from **BocP-a**: $M_n = 7.3 \text{ kg mol}^{-1}$, $D = 1.83$) in toluene-d₆, and (b) variable-temperature UV-Vis spectra of **HPLP-a** (from **BocP-a**: $M_n = 16.1 \text{ kg mol}^{-1}$, $D = 1.42$) in thin film.

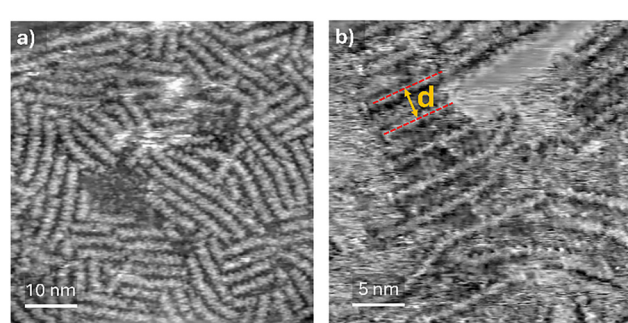


Fig. 5 (a) STM image of a high coverage area of **BocP-b** ($M_n = 19.3 \text{ kg mol}^{-1}$, $D = 1.50$) deposited by electrospray deposition on Au(111). (b) Zoomed-in image showing sections of the polymers align parallel to each other, with a distance d between them.



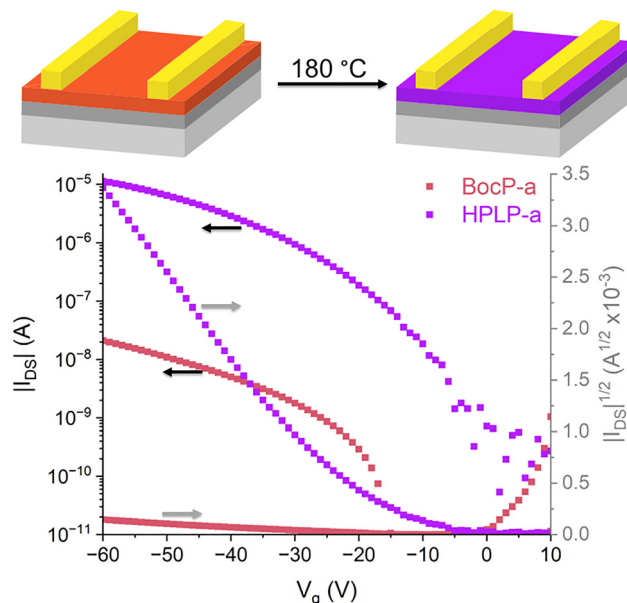


Fig. 6 (top) Schematic illustration of the thermal treatment process on the OFET device of **BocP-a**, enabling the *in situ* transformation of the active layer from **BocP-a** to **HPLP-a**; (bottom) transfer characteristics of the OFET devices before (**BocP-a**) and after (**HPLP-a**) thermal treatment at $V_{DS} = 60$ V.

$5.2 \times 10^{-5} \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$. Such device was directly heated in N_2 atmosphere at 180°C for 1 h to convert **BocP-a** in the active layer into **HPLP-a** *in situ*. The resulting devices show a hole mobility of $4.6 \times 10^{-2} \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$, marking a significant increase for three orders of magnitude. An improvement was also observed when comparing **BocP-b** to **HPLP-b** with films exhibiting hole mobilities of $8.5 \times 10^{-9} \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$ and $3.4 \times 10^{-6} \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$, respectively (Fig. S36, ESI†). The drastic enhancement in charge carrier mobility can be attributed to change in electronic band gap, lower reorganization energies, and improved packing of the pseudoladder polymer chains containing intramolecular hydrogen-bonding interactions. To investigate this, atomic force microscopy (AFM) was performed on hydrogen-bonded polymers and their Boc-protected precursors (Fig. S37, ESI†). AFM revealed a reduction in film thickness from **BocP-a** (40 nm) to **HPLP-a** (26 nm), while the root-mean-square (rms) roughness remained similar (0.5 nm). A less pronounced thickness change was observed between **BocP-b** (42 nm) and **HPLP-b** (35 nm), with the rms roughness again unchanged (1.7 nm). The decreased film thickness in the hydrogen-bonded polymers correlates with their enhanced charge transport, while the higher rms roughness of **HPLP-b** compared to **HPLP-a** may contribute to its lower charge-carrier mobilities.

Along with the extraordinary improvement in electronic performance upon hydrogen bond-induced coplanarity, **HPLP-a** lies among some of the highest mobilities achieved for benzodifuran based conjugated polymers.^{49–55} As benzobisimidazole is unexplored in the field of organic semiconducting materials, its effect on coplanarity and rigidity of polymer

chains may serve to maximize performance of other high-performing monomers as it has done with benzodifuran.

Conclusion

In conclusion, we demonstrate a feasible approach to achieve conjugated pseudoladder polymers locked into a coplanar conformation by intramolecular hydrogen bonds, and the unique changes that result in the properties of the polymers. The incorporation of Boc protecting groups also allows for processability of the polymer precursors for solution-based device fabrication, while allowing access to hydrogen bond-induced coplanar conformation through simple *in situ* thermal treatment. The transition from a flexible, noncoplanar polymer chain to a rigid coplanar conformation was accompanied by a significant decrease in solubility and in the bandgap. The rigidity of the hydrogen bonded polymers was confirmed by the sharp vibrational peaks in the UV-vis spectra, and demonstrated to be highly robust as evidenced by a series of variable temperature analyses. OFET devices were fabricated by solution processing the soluble Boc-protected precursor followed by *in situ* Boc-deprotection. The device performance of the hydrogen bonded pseudoladder polymer, **HPLP-a**, showed an improvement in hole mobility by almost 3 orders of magnitude when compared to its Boc-protected counterpart. Solvent resistance of the semiconducting **HPLP** films also shows potential for multilayer film processing in organic electronic devices. This work demonstrates the potential of hydrogen-bond manipulation as a versatile approach for developing solution-processable, high-performance semiconducting polymers, effectively addressing the trade-off between electronic performance and processability in organic electronic materials.

Author contributions

The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript.

Conflicts of interest

The authors declare no competing financial interests.

Data availability

The data supporting this article have been included as part of the ESI.† Crystallographic data for compound 4 has been deposited at the CCDC under 2422384 and can be obtained from <https://www.ccdc.cam.ac.uk/structures/Search?Ccdeid=2422384&DatabaseToSearch=Published>. Crystallographic data for compound 5 has been deposited at the CCDC under 2422385 and can be obtained from <https://www.ccdc.cam.ac.uk/structures/Search?Ccdeid=2422385&DatabaseToSearch=Published>.



Acknowledgements

O. M. acknowledges the National Science Foundation Graduate Research Fellowship Program under grant no. DGE-2139772. The authors acknowledge the Robert A. Welch Foundation (A-1898) for financial support of this work. Portions of this research were conducted with the advanced computing resources provided by Texas A&M High Performance Research Computing. Work at the Molecular Foundry was supported by the Office of Science, Office of Basic Energy Sciences, of the U.S. Department of Energy under contract no. DE-AC02-05CH11231.

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