A universal and facile approach for building multifunctional conjugated polymers for human-integrated electronics

Building high-performance polymer semiconductors with diverse functionalities is of great importance for developing human-integrated electronics. A universal and facile “click-to-polymer” (CLIP) synthesis strategy is established here. This strategy employs a one-step, versatile reaction to allow the covalent attachment of various functional units. This can be utilized to realize both bulk and surface functionalization in or on the deposited thin films. Using this CLIP approach, a wide range of non-conventional functional properties are and can be incorporated onto polymer semiconductors.
A universal and facile approach for building multifunctional conjugated polymers for human-integrated electronics

Nan Li, Yahao Dai, Yang Li, Shilei Dai, Joseph Strzalka, Qi Su, Nickolas De Oliveira, Qingteng Zhang, P. Blake J. St. Onge, Simon Rondeau-Gagné, Yunfei Wang, Xiaodan Gu, Jie Xu, and Sihong Wang

SUMMARY
Polymer semiconductors have shown distinct promise for the development of human-integrated electronics, owing to their solution processability and mechanical softness. However, numerous functional properties required for this application domain face synthetic challenges to being conveyed onto conjugated polymers and thus combined with efficient charge-transport properties. Here, we develop a “click-to-polymer” (CLIP) synthesis strategy for conjugated polymers, which uses a click reaction for the facile and versatile attachment of diverse types of functional units to a pre-synthesized conjugated-polymer precursor. With four types of functional groups, we show that functionalized polymers from this CLIP method can still retain good charge-carrier mobility. We take two realized polymers to showcase a photopatterning property and a biochemical sensing function, both of which advance the state of the art of realizing these two types of functions on conjugated polymers. We expect that the expanded use of this synthesis approach can largely enrich the functional properties of conjugated polymers.

INTRODUCTION
Conjugated polymers, which feature mechanical softness and solution processability, have undergone substantial development for several types of electronic devices, including organic field-effect transistors, organic photovoltaics, organic light-emitting diodes, and electrochromic devices. Their functional performances have already been shown to be on par with those of inorganic materials. More recently, conjugated polymers have been demonstrated as the only class of electronic materials that can provide skin-like stretchability, thereby rendering them prime candidates for human-integrated electronics for health monitoring, disease diagnosis, and medical treatments. However, toward the realization of such applications of collecting and delivering various types of information from and to human bodies, there have been few designs for conjugated polymers to provide a number of emerging functions, which include, but are not limited to, biochemical sensing, chemotherapeutic properties, bio/immune compatibility, micropatternability, tissue/skin adhesion, and stimulus response. Currently, the absence of these functional properties poses the major obstacle to taking advantage of the unique properties of conjugated polymers to benefit the development of human-integrated electronics.
As these functional properties are typically afforded by certain functional units, such as bioreceptors, drug motifs, antifouling groups, and photo/thermoreactive groups, the most straightforward and effective approach for building these functions into conjugated polymers should be to incorporate these functional units into the molecular design, such as on the side chains. However, significant challenges arise in the synthesis of these types of molecular structures following the conventional procedure (Figure 1A) of first incorporating the desired functional units during the monomer synthesis and then carrying out the polymerization reaction at the end.\textsuperscript{14–18} Since many functional units of the above-listed types either are highly sensitive to harsh conditions, such as high temperature, high pressure, and toxic reagents, or tend to significantly change a monomer’s solubility, their attachment onto a conjugated monomer either cannot survive through the polymerization reaction or would largely decrease the efficacy of the polymerization. For other functional units that could be compatible with this synthesis procedure, they each require a unique design and optimization of the synthesis conditions, which further elevates the technical barrier. So far, the successes along this line for conjugated polymers have been achieved on only a limited number of functional groups of the above-listed types, for example, lysine groups for enhancing cell viability\textsuperscript{14} and photochromic groups for optically tunable electrical performance.\textsuperscript{19}

Alternatively, surface modification on a pre-deposited thin film of conjugated polymers presents another option for incorporating a type of function that mainly relies on surface properties, e.g., chemical sensing, biocompatibility, and adhesion. However, with the lack of suitable reactive sites in the typical design of conjugated polymers, the viable methods are mostly limited to physical approaches such as coating or adsorption,\textsuperscript{20,21} which are applicable to only a small portion of functional groups and typically result in poor stability. For the generally preferred modification through covalent assembly, additional pre-treatments (e.g., oxygen plasma activation) of the film surface are often needed for generating reactive groups,\textsuperscript{22–25} which usually cause undesired side reactions or even degradation to conjugated polymers.

Therefore, to unlock a broad range of desired functionalities for conjugated polymers, it is highly necessary to develop a different synthesis approach that can fully decouple the covalent attachment of functional units from the polymerization reaction. Herein, we report a generalizable molecular design and synthesis approach of using click chemistry as a post-polymerization step to versatility graft a variety of functional units onto the side chains of high-performance donor-acceptor-type semiconducting conjugated polymers\textsuperscript{1,18} (Figure 1B), which we name as the click-to-polymer (CLIP) method. Specifically, with the criteria of having high reaction efficacy, sufficient compatibility of the clickable site with the cross-coupling (e.g., Stille coupling) polymerization for donor-acceptor conjugated polymers, a simple reaction procedure, and minimal influence on the charge-carrier transport, we rationally chose the copper-catalyzed azide-alkyne cycloaddition (CuAAC)\textsuperscript{26,27} as the click reaction for the CLIP method. To the best of our knowledge, the CuAAC reaction so far has neither been used for the functionalization of a donor-acceptor conjugated polymer nor achieved high-performance semiconducting property through consideration of the structure-property relationships.\textsuperscript{28–31} Moreover, we hypothesize that the high efficacy and the broad tunable space of the reaction conditions of this click chemistry could also allow the versatile use of CLIP for not only bulk functionalization but also surface functionalization, by simply swapping the sequence of the click reaction and the thin-film deposition (Figure 1C), which, to the best of our knowledge, has never been reported before on a single method. This will provide the freedom of choosing the most suitable functionalization approach from the combined

\textsuperscript{1}Pritzker School of Molecular Engineering, The University of Chicago, Chicago, IL 60637, USA
\textsuperscript{2}X-Ray Science Division, Argonne National Laboratory, Lemont, IL 60439, USA
\textsuperscript{3}Department of Chemistry and Biochemistry, University of Windsor, Windsor, ON N9B3P4, Canada
\textsuperscript{4}School of Polymer Science and Engineering, University of Southern Mississippi, Hattiesburg, MS 39406, USA
\textsuperscript{5}Nanoscience and Technology Division, Argonne National Laboratory, Lemont, IL 60439, USA
\textsuperscript{6}Lead contact
*Correspondence: sihongwang@uchicago.edu
https://doi.org/10.1016/j.matt.2021.07.013
considerations of both the nature of the targeted function and the preservation of electrical performance.

In this work, we demonstrate the versatility of this approach on four types of functional groups (Figure 1B, bottom) with photo-cross-linkable,\textsuperscript{32} bioconjugation,\textsuperscript{33} immune-modulatory,\textsuperscript{34} and ionic conducting\textsuperscript{35} properties, respectively. From the synthetic perspective, they cover the challenging characteristics of thermal reactivity, solvent incompatibility, and sized volume, as well as serving to solve the challenge from less stable biochemical units. The obtained conjugated polymers with the bulk functionalization of these units still give largely preserved charge-carrier mobility (above 0.1 cm$^2$ V$^{-1}$ s$^{-1}$ from three of the four functionalized polymers), while the
surface functionalization does not have any influence on the mobility. To demonstrate the usefulness of this CLIP strategy, we realized two highly desired functions on conjugated polymers: direct photopatterning with sub-10-μm resolution enabled by the incorporation of benzophenone (BP) groups, and amplified biomolecular sensing enabled by the incorporation of N-hydroxysuccinimide (NHS) ester, which can conjugate with enzymatic bioreceptors. Moving forward, we envision that the further expansion of our CLIP synthesis approach to the incorporation of a large variety of functional units, such as therapeutic motifs and diverse groups of biomolecules, can greatly enrich the functions and applications of conjugated polymers for human-integrated wearable and implantable electronics.

RESULTS AND DISCUSSION

In our proposed CLIP method, the first key step is to attach the azide group onto the side chains of conjugated monomers (e.g., the acceptor units for the synthesis of donor-acceptor polymers, as shown in Figure 1B, top) during the monomer synthesis process, which serves as the click site for the later CuAAC reaction with a functional group modified with the alkyne group. For most types of functional groups, such alkyne modification can be achieved under compatible, mild conditions. To reduce potentially unfavorable influences of bulky functional groups on the packing structure of conjugated polymers, we introduced a linear alkyl linker between the clickable site and the backbone. Since the azide group will neither be activated (Figures S1 and S2) under the typical conditions of the polymerization reaction (e.g., the Stille-coupling reaction with the temperature range of 180°C–200°C) nor influence the monomer’s solubility, the subsequent polymerization can still be carried out as usual.

Bulk functionalization on conjugated polymers through the CLIP method

With the donor-acceptor backbone of diketopyrrolopyrrole (DPP)-thieno[3,2-b]thiophene (DPPTT) used as the model system, we first demonstrate the bulk functionalization process and characterize the polymers obtained. We synthesized DPP monomers (Figure 2A) with azide-substituted side chains in two different alkyl-linker lengths, 6 and 10 carbons, which were further taken to copolymerize with DPP units of conventional branched alkyl side chains in two ratios of 1:9 and 2:8. The reason for copolymerization with branched alkyl chains is to balance the desired functionality and the good solution processability of the modified polymers. It should be noted that by adding some solubilizing side chains onto the donor (TT) units, the fraction of the functionalized DPP units could be further increased while keeping sufficient solubility. Through the same polymerization condition, the four types of azide-substituted polymers, namely C6-10a (i.e., six-carbon alkyl linker and 10% azide-substituted DPP units), C6-20a, C10-10a, and C10-20a, all have lower molecular weights than the DPPTT polymer only with branched alkyl side chains (Table S1), which should come from the slight decrease in the solubility caused by the partial replacement of branched side chains with linear side chains. Subsequently, the four types of functional units as shown in Figure 1B, i.e., tetrahydropyran (PY), poly(ethylene glycol) 5-NHS (PEG5NHS), BP, and poly(ethylene glycol)-2000-kDa (PEG2000), were grafted onto the polymers via the CuAAC click reaction. The success of our CLIP method was verified by nuclear magnetic resonance (NMR; Figure 2B) and Fourier transform infrared (FTIR; Figure S3) spectroscopy results, which all clearly show the existence of the azide groups after the polymerization reaction and the subsequent disappearance of the azide after the click reaction.

To evaluate the influence of side-chain functionalization done by the CLIP method on the polymers’ electrical performance, we fabricated thin-film transistor (TFT)
Figure 2. Characterization of conjugated polymers with bulk functionalization through the CLIP method

(A) Reaction scheme for the use of the CuAAC click reaction for functionalizing four types of units on azide-functionalized DPPTT-conjugated polymer, with two different alkyl linker lengths ($n = 4$ or $8$, respectively) and two copolymerization ratios ($y = 0.1$ or $0.2$, respectively) to the number of repeating units.

(B) High-temperature $^1$H NMR spectra for the representative polymers before and after the click reaction. The green line labeled with a circle ($\bullet$) stands for the proton peak adjacent to the azide group in the starting polymer as indicated in (A). The green line labeled with a triangle ($\triangle$) stands for the proton peak adjacent to the triazole group in the functionalized polymer as indicated in (A).

(C) OTFT device structure for charge-carrier mobility measurement.

(D) Typical transfer characteristics obtained from the functionalized polymer C10-10PY. $I_d$, drain current, represented by solid lines; $I_g$, gate current, $V_{gs}$, the gate-source voltage.

(E) Typical output characteristics from C10-10PY.

(F–H) Saturation and linear charge-carrier mobilities (F), relative degrees of crystallinity and $\pi-\pi$ spacings (extracted from GIXD measurements) (G), and normalized UV-vis absorption spectra (H), from the unfunctionalized DPPTT polymer, azide-modified polymers (C6-10a, C10-10a, C10-20a), and further obtained PY-functionalized polymers (C6-10PY, C10-10PY, C10-20PY). C6 and C10 stand for the alkyl-linker lengths of the 6-carbon and 10-carbon, respectively; 10a/PY and 20a/PY stand for the side-chain modification with the azide/PY group in the molar ratio of 10% and 20%, respectively.
devices to characterize the charge-carrier mobility. These are in the bottom-contact/ 
bottom-gate device structure with SiO₂ and n⁺⁺-Si as the gate dielectric and gate 
electrodes and gold as the source and drain electrodes (Figure 2C). In general, all 
the obtained polymers give ideal transfer and output characteristics (Figures S4 
and S5), as represented by the polymer with PY functionalization on C10-10a (Fig-
ures 2D and 2E). Compared with the unfunctionalized DPPTT polymer with 100% 
branched-alkyl side chains, there is only some minor decrease in mobility (Figures 
2F–2I and S6, Tables S2 and S3) resulting from the side-chain modifications with 
the azide group alone and the four types of functional units. Such changes in mobility 
could be the combined effects of the changes in the packing structure, the molecular 
weight, and the deposited film morphology. The extent of such influences is dictated 
by both the side-chain structure (i.e., the alkyl-breaker length) and the molar ratio in 
the copolymerization. As shown in Figure 2F, the CLIP of the functional group PY on 
C10-10a results in slightly less decrease in mobility than on C6-10a. From Figures 2G 
and 2H, the longer alkyl linker (C10 versus C6) reduces the steric hindrance from the 
functional group to the polymers’ packing ordering, as shown by the crystallinities 
measured from grazing-incidence X-ray diffraction (GIXD) and aggregation level 
characterized by ultraviolet-visible (UV-vis) light (Figures 2G, 2H, and S7). For the 
C10 polymers, on the other hand, the increase in the functionalization ratio from 
10% to 20% (i.e., the resulting polymer of C10-20PY) leads to lower mobility (Fig-
ure 2F), which should also come from the elevated disruption to the packing struc-
ture. In addition, the decreased molecular weights of the functionalized polymers 
could also be a reason for the decrease in mobility,⁴¹ which in the future could be 
improved by enhancing the solubility of the monomers and resulting polymers by 
introducing additional solubilizing groups to the backbone.⁴²

The further expanded comparison of the other three types of functional groups, i.e., 
PEG5NHS, BP, and PEG2000, together with PY, functionalized on C10-10a polymer 
all achieved mobility above 0.1 cm² V⁻¹ s⁻¹, except PEG2000 (Figure 2I). In general, 
the primary influence from the type of functional group should be the different steric 
hindrance effects in disrupting the packing structure (Figures 2J and S8). As ex-
pected, PEG2000, as the bulkiest group, results in the most substantial decrease in 
 mobility. Moreover, it is observed that the functionalization overall leads to 
increased surface roughness of the deposited thin film compared with C10-10a (Fig-
ures 2K and S9), which could be another factor causing the slight decrease in 
 mobility. Overall, benefiting from the minimal influence on the polymerization effi-
cacy and the tunability of the side-chain structure, the functionalization to conju-
gated polymers using the CLIP method can largely preserve the electrical 
performance.

To demonstrate the unprecedented synthesis capability given by the CLIP strategy, 
we made an experimental comparison to the use of the conventional pre-polymer-
ization functionalization approach for the synthesis of the 10PEG2000 polymer, in 
which the PEG2000 unit is first grafted onto the DPP monomers prior to polymeri-
ization. With such a functionalized monomer, the polymer obtained from the same 
Stille-coupling reaction indeed has a much lower molecular weight, together with 
a very low mobility of around 0.001 cm² V⁻¹ s⁻¹ (Figure S10), which should come
from the substantial decrease in the reactivity of the functionalized DPP monomer as a result of the substitution of such a bulky and polar group. This comparison clearly shows that the CLIP method provides unparalleled capability in functionalizing conjugated polymers without affecting the synthesis outcomes.

**Surface functionalization on conjugated polymers through the CLIP method**

Compared with bulk functionalization, surface functionalization on a pre-deposited thin film can avoid the disruption to the polymer’s packing structures by sizable functional groups. To enable the CuAAC click reaction on such surfaces, the azide groups as the clickable sites need to be effectively exposed on the surfaces, which is indeed satisfied by the edge-on packing orientation (Figure S7) of the deposited C10-10a and C10-20a films. However, with the much lower amount of azide groups available on the surface compared with the fully solubilized polymers, sufficiently high reaction efficiency is highly needed. For this, we chose N,N-dimethylformamide (DMF) as the solvent for the click reaction, as it offers good wettability to these conjugated polymer films, thus facilitating the transport of the reactant molecules to the surface. Under this condition, we carried out the surface functionalization on a C10-10a film with the unit of PEG2000. After the reaction at room temperature overnight followed by a thorough rinse to fully remove the excess reagents, we observed a decrease in the water contact angle from 98° to 67°, which suggests the successful grafting of the PEG2000 chain (Figures 3B and S11). This was further verified through the X-ray photoelectron spectroscopy (XPS) characterization, which shows the appearance of the C 1s peak (Figure 3C) and the increased intensity of the O 1s peak (Figure S12), both corresponding to the formed C-O bond. To further exclude the possibility of physical absorption, a control experiment was carried out with the same reaction condition but without the catalyst for the click reaction, which gave little change in the contact angle or the XPS spectrum. The measurement of its charge-carrier mobility in organic thin-film transistors (OTFTs) indeed proves that such surface functionalization causes negligible influence on the electrical performance (Figures 3D and S13), so that the mobility is almost 1 order of magnitude higher than that from its bulk-functionalized counterpart.

**Direct photopatterning of conjugated polymers enabled by functionalization of the benzophenone group**

Among the four types of functional units incorporated in this work, the BP group can enable interchain cross-linking (Figure 4A) for the functionalized conjugated polymer through the photoinitiated reaction with an adjacent C-H bond (Figure 4B) under mild UV (365 nm) exposure. This will impart solvent resistance to pre-annealed (at 140°C for 30 min) 10BP films when the UV exposure time is longer than 5 min (under the 365-nm wavelength and 2.6-mW/cm² intensity), as verified by the immersion test in chloroform (Figure S14). This could thus enable one-step direct photolithography for the patterning of the conjugated polymer film, by using a photomask to limit the UV exposure to selected areas. As conjugated polymers are generally not compatible with the solvents of normal photoresists and developers, their patterning into small feature sizes has been one of the major challenges to using them to fabricate functional circuits. Among all the potential options, the direct photopatterning allowed by imparting photo-cross-linkable properties to a conjugated polymer is the most ideal choice for simplicity in fabrication. So far, there are very few reports about the direct photopatterning of a semiconducting polymer through the use of an azide cross-linking reaction, by either blending an azide-containing cross-linker (e.g., bis(fluorophenyl azide)) or attaching the azide group onto the polymer side chain. However, since the azide cross-linking chemistry requires the use of deep-UV light (i.e., with a wavelength of 254 nm) and a relatively
high exposure dose, detrimental effects on the conjugated polymers could occur, as
verified by the UV-exposure experiment on our C10-10a polymer that also has the
azide group (Figure S15). Alternatively, the BP photo-cross-linking chemistry that
is incorporated into conjugated polymers for the first time should be more benign
for preserving electrical performance after the direct photopatterning.

We then carried out a systematic study of the cross-linking conditions to realize the
full preservation of electrical performance from 10BP films after the direct photol-
ithography process. By testing varied UV exposure durations, all followed by a
post-bake step (at 170°C for 30 min) to fully remove the development solvent (i.e.,
chloroform) from the film, we found that a UV exposure time equal to or longer
than 10 min can ensure the full preservation of the charge-carrier mobility (Figures
4Ca and S16) after the solvent removal of the unexposed areas. Consequently, the
mobility of ~0.2 cm² V⁻¹ s⁻¹ is obtained from the patterned films of conjugated
polymers. Under this photopatterning condition, we further demonstrated the trans-
fer of pre-designed patterns from masks to 10BP films through the one-step photo-
lithographic process, which can indeed work well for complicated geometries and
achieve a resolution below 10 μm (Figures 4D, 4E, and S17). It can be envisaged
that this CLIP-based incorporation of a BP group can be adopted to other designs
of conjugated polymers, so as to obtain photopatternable polymer semiconductors
with even higher mobility.

Biochemical sensing enabled by the immobilization of an enzymatic
bioreceptor
The NHS ester, another group that gets functionalized onto conjugated polymers using
the CLIP method, can serve to endow conjugated polymers with selective sensing to tar-
targeted biomolecules through the bioconjugation reaction with the primary amine group

Figure 3. Characterizations of conjugated polymers with surface functionalization through the CLIP method
(A) Schematic images showing the process for performing surface CLIP on a deposited film from an azide-modified conjugated polymer (i.e., C10-10a in
the demonstrated experiment).
(B) Water contact angles on C10-10a film, the film modified with PEG2000 by the surface CLIP process, and the control film obtained from a C10-10a film going
through the same surface CLIP process but without the Cu²⁺ catalyst. The error bars represent the standard deviations obtained from at least four measurements.
(C) XPS spectra showing the changes in the C 1s peaks from the three films in (B).
(D) Saturation and linear charge-carrier mobilities from C10-10a and the functionalized polymers with PEG2000 through surface-CLIP and bulk-CLIP.
The error bars represent the standard deviations obtained from at least six measurements.
in protein/polypeptide-type bioreceptors. Since biochemical sensing is one of the major approaches for health monitoring and disease diagnosis, imparting such functions to conjugated polymers is highly desirable for their use in human-integrated electronics.

So far, although there have been a few examples of using conjugated polymers as the channel layer in transistor-type sensors that can uniquely offer built-in amplification for achieving higher sensitivity, the specificity was imparted through either a non-cova-
lent physical absorption method,20 which lacks long-term robustness, or complicated surface treatment methods,47,48 such as templating and plasma deposition, which are applicable to only a narrow range of bioreceptors.

To demonstrate this biochemical sensing function offered by the functionalized NHS ester group, we chose to immobilize the glucose oxidase (GO) enzyme as the bio-
receptor for the sensing of glucose. Recently, based on the mixed electronic-ionic conducting property of conjugated polymers, organic electrochemical transistors (OECTs) have been developed as a very promising device platform for biosensing, which can provide even higher amplification to sensing signals than normal field-eff-
tector transistors.49–51 To adopt the functionalization of the NHS ester group to conju-
gated polymers for OECT operations, we copolymerized the azide-attached DPP monomer with the glycolated 2,2'-bithiophene (g2T) monomer,52,53 at a ratio of

1:9 (Figure S18), so as to achieve the desired ionization potential and aqueous-
swelling property for OECT operation. Using this polymer, namely g2T-NHS, the glucose sensor was fabricated (Figure 5B) by first depositing the semiconducting film onto patterned gold source/drain electrodes on a glass substrate, and then treating the channel area with a solution of GO enzyme for 2 h to achieve the covalent immobilization of this bioreceptor. The gating is made with the extended configuration of having another piece of gold film on the side of the channel, which gets bridged to the channel by the added analyte solution (Figure S19). When there is glucose in the solution, its oxidation under the catalysis of the GO enzyme will transfer electrons to the channel layer, thus dedoping the p-type conjugated polymer20 (Figure 5B, right).

As shown in Figure 5C (see also Table S3), the enzyme immobilization on the semiconducting channel causes only a small shift in the threshold voltage of the OECT.

Figure 5. Biochemical sensing to glucose from conjugated polymers enabled by covalent grafting of enzymatic bioreceptors

(A) Reaction scheme of the covalent bioconjugation of glucose oxidase (GO) enzyme on g2T-NHS polymer.

(B) Fabrication process for the enzymatic sensor based on OECT device design, and the recognition mechanism to glucose at the surface of the GO enzyme-modified channel layer (right). Glc, glucose; Glc-ox, glucono-1,5 lactone.

(C) Transfer curves from the OECT sensor devices with GO-enzyme-modified g2T-NHS (labeled as g2T-NHS-enzyme) or pristine g2T-NHS as the channel layer.

(D) Real-time current response ($V_{gs} = -0.2$ V, $V_{ds} = -0.05$ V) of an as-fabricated glucose sensor with g2T-NHS-enzyme to the stepwise addition of glucose in different concentrations.

(E) Drain current ($|\Delta I_d|$) response to glucose concentrations in the range of 1–1,000 μM.
operation in phosphate-buffered saline (PBS) solution. The sensing of glucose was characterized by adding glucose in controlled concentrations into the PBS solution. When the concentration of the added glucose was equal to or greater than 1 μM, a stepwise decrease in the current was observed (Figures 5D and S20). The extent of the current decrease exhibits good correlation with the added glucose concentration (Figure 5E). In contrast, when only the PBS solution was added, there was no change in the current, suggesting the specificity for glucose as afforded by the enzyme. As a control, the device made from the pristine NHS-functionalized semiconducting film showed no response to glucose (Figure S21), which verifies that the sensing function is indeed afforded by the GO enzyme. Since the conjugation chemistry to the NHS ester group has broad applicability to other types of protein- or polypeptide-based bioreceptors (including both enzymes and antibodies), this material design and synthesis concept based on the CLIP method opens up an avenue to covalently functionalizing bioreceptors on mechanically soft, potentially stretchable, conjugated polymers for realizing biochemical sensing at biointerfaces with specificity and biocompatibility.

DISCUSSION

In summary, we established a new synthesis strategy, the CLIP method, for largely expanding the synthesizable molecular design of conjugated polymers, toward incorporating a wide range of emerging functions. By using a click reaction as a post-polymerization step for the functionalization of side chains, this CLIP method is shown to be facile and highly versatile for different functional units and for both bulk and surface functionalization. This method was successfully utilized to covalently graft four different types of functional groups onto the side chains of conjugated polymers, which, otherwise, are all highly challenging to incorporate. Two of the developed polymers were further taken to realize two highly desirable functions on conjugated polymers: direct photopatternability and biochemical sensing with specificity. Additional functionalities that can be incorporated by the approach can become highly valuable topics for future investigations on the expanded uses of conjugated polymers. Through innovative incorporation of click chemistry into the design and synthesis paradigm of high-performance conjugated polymers, we anticipate that this synthesis strategy will greatly enlarge the molecular design space and thus enrich the functional properties of conjugated polymers. This will satisfy the need for their development toward the far-reaching application area of human-integrated electronics.

EXPERIMENTAL PROCEDURES

Resource availability

Lead contact

Further information and requests for resources and materials should be directed to and will be fulfilled by the lead contact, Sihong Wang (sihongwang@uchicago.edu).

Materials availability

All the chemicals used in the study were purchased from Sigma-Aldrich and used without further purification. Anhydrous solvents were purchased from either Sigma-Aldrich or Fisher Scientific. 5,5'-Dibromo-3,3'-bis(2-(2-(methoxyethoxy)ethoxy)ethoxy)ethoxy)ethoxy)-2,2'-bithiophene was purchased from SunaTech (cat.no. IN1273) and was used as received. Column chromatography was carried out with silica gel for flash chromatography from VWR Scientific. The other compounds reported in the paper can be produced following the procedures in the supplemental information.
Data and code availability
The published article includes all data generated or analyzed during this study.

Characterizations
Microwave polymerization was conducted using a Biotage Initiator+. NMR spectra were recorded on a Bruker Avance III HD console spectrometer (T H 400 MHz, T C 100 MHz) at 293 K. Chemical shifts are given in parts per million with respect to tetramethylsilane as an internal standard, and coupling constants (J) are given in hertz. Polymer NMR spectra were recorded at 393 K in deuterated 1,1,2,2-tetrachloroethane. High-resolution mass spectra were recorded on an Agilent 6530 LC Q-TOF mass spectrometer using electrospray ionization with fragmentation voltage set at 70 V and processed with an Agilent MassHunter operating system. Number average molecular weight, weight average molecular weight, and polydispersity index were evaluated by high-temperature size-exclusion chromatography using 1,2,4-trichlorobenzene and performed on an EcoSEC HLC-8321GPC/HT (Tosoh Bioscience) equipped with a single TSKgel GPC column (GMHHR-H; 300 mm × 7.8 mm) calibrated with monodisperse polystyrene standards. The samples were prepared using 1 mg/mL of sample in trichlorobenzene, and were allowed to stir at 80 °C for 12 h prior to injection. The analysis of the samples was performed at 180 °C with a flow rate of 1.0 mL/min with injection quantities of 300 μL. The data were collected and integrated using the EcoSEC 8321GPC HT software suite. UV-vis absorption spectra were recorded on the Shimadzu UV-3600 Plus UV-VIS-NIR spectrophotometer. FTIR spectra were recorded on the Shimadzu IRTracer-100 FTIR spectrometer. Differential scanning calorimetry experiments were performed with a TA Instruments Discovery 2500 differential scanning calorimeter. Thermal gravimetric analysis plots were recorded with a TA Instruments Discovery thermogravimetric analyzer. The water contact angle measurement was done with a KRÜSS DSA100 drop shape analyzer. The XPS was done with a Kratos AXIS Nova with a monochromatic Al Kα X-ray source and a delay line detector system. Optical microscope images were captured with a Zeiss Axioscope 5/7/Vario microscope. GIXD was performed at the Advanced Photon Source at Argonne National Laboratory on beamline 8-ID-E. The X-ray energy was 10.92 keV, the incident angle was 0.14°, and the exposure time was 10 s. The area detector (Pilatus 1M, Dectris) was translated vertically for a second exposure. The two images were combined to eliminate gaps due to rows of inactive pixels at the borders between modules using the GIIXSGUI package55 for MATLAB and demonstrating that the samples were not damaged by the exposure.

Device fabrication and characterization
The TFTs were based on bottom-contact/bottom-gate structure. The n-octadecyltriethoxysilane (OTS)-treated Si/SiO2 (300 nm) substrates56 were cleaned with toluene. After that, the source/drain gold electrodes (50 nm) were patterned via thermal evaporation with a metal shadow mask. The channel length (L) and width (W) were 200 μm and 4 mm, respectively. The gold surface was then modified by submerging the substrates in 30 mL isopropyl alcohol (IPA) with 10-μL 2,3,4,5,6-pentafluorothiophenol to form a self-assembled monolayer. The substrates were gently rinsed with IPA and blow-dried with nitrogen gas. The polymer solution (5 mg/mL in chlorobenzene) was then spin-coated at 1,000 rpm for 60 s, followed by annealing at 150°C for 1 h in a nitrogen atmosphere. All of the electrical characteristics of the semiconducting layer were measured using a Keithley 4200 (Keithley Instruments, Cleveland, OH, USA) under an ambient environment. The details on mobility evaluation in OTFTs are discussed in Figure S6.
The OECTs were fabricated similar to the methods previously reported. Briefly, the glass substrates were cleaned with acetone, IPA, and water. After that, the source/drain gold electrodes (50 nm) were patterned via thermal evaporation with a metal shadow mask. The channel $L$ and $W$ were 200 $\mu$m and 4 mm, respectively. Then the polymer solution (5 mg/mL in chloroform) was spin-coated at 1,000 rpm, followed by annealing at 110 $^\circ$C for 30 min in nitrogen atmosphere. The electrolyte was a PBS solution dropped on top of the transistors. The gate electrode was gold, which was immersed in the electrolyte. The electrical characteristics of the semiconducting layer were measured using a Keithley 4200 (Keithley Instruments, Cleveland, OH, USA) under an ambient environment.

**Surface CLIP protocol**
The surface CLIP follows the general procedure. C10-10a and C10-20a films were first spin-coated on OTS-treated silicon substrate and annealed at 150 $^\circ$C for 1 h. The solution of alkyne-PEG2000 (4 mg, 2 $\mu$mol) was dissolved in DMF (0.5 mL) and mixed with a solution of CuSO$_4$ (10 $\mu$L, 0.1 M, 1 $\mu$mol) and sodium ascorbate (20 $\mu$L, 0.1 M, 2 $\mu$mol) in water. The reaction mixture was placed on the film surface surrounded by a polydimethylsiloxane well and incubated at room temperature overnight. The surface was rinsed with H$_2$O and DMF and finally dried with nitrogen gas.

**Direct photopatterning for 10BP conjugated polymer**
The C10-10BP polymer (5 mg/mL in chlorobenzene) was spin-coated on octadecyltrichlorosilane-modified silicon oxide substrate at 1,000 rpm for 60 s. The polymer film was first annealed at 140 $^\circ$C for 30 min to remove solvent and boost the relative crystallinity. The film thickness obtained was around 30–40 nm. Then the conjugated polymer film was exposed to UV light (365 nm, 2.6 mW/cm$^2$, Spectrolinker XL-1000) with selected areas for a given period. After the film was immersed in chloroform (also a good solvent for dissolving C10-10BP polymer) for 30 s with gentle shaking, the film was blow-dried with nitrogen gas and then baked at 170 $^\circ$C for 30 min inside a glove box to fully remove the trapped solvent in the polymer network and further increase the relative crystallinity.

**OECT glucose sensing**
For the sensing experiment, GO dissolved in PBS (pH 7.4) (3 mg/mL) was incubated on the device channel area for 2 h and then rinsed with DI water. Glucose was dissolved as stock solutions in PBS. Current-voltage characteristics of the devices were recorded using a Keithley 4200 under an ambient environment. After a steady baseline was obtained for the drain current, the glucose stock solution (4 $\mu$L each time) was injected into the electrolyte gently. For all the experiments, the volume of the electrolyte solution was kept at 0.4 mL and drain current changes in response to subsequent additions of increasing concentrations of glucose solutions into the electrolyte were monitored as a function of time.

**SUPPLEMENTAL INFORMATION**
Supplemental information can be found online at https://doi.org/10.1016/j.matt.2021.07.013.

**ACKNOWLEDGMENTS**
This work is supported by a start-up fund from the University of Chicago and the US Office of Naval Research (N00014-21-1-2266). J.X. acknowledges the Center for Nanoscale Materials, a US Department of Energy Office of Science User Facility
and supported by the US Department of Energy Office of Science, under contract DE-AC02-06CH11357. Y.W. and X.G. thanks the U.S. Department of Energy, Office of Science, Office of Basic Energy Sciences under award number of DE-SC0019361 for providing resources related to the scattering experiments in this work. This research used resources of the Advanced Photon Source, a US Department of Energy Office of Science User Facility, operated for the Department of Energy Office of Science by Argonne National Laboratory under contract DE-AC02-06CH11357. S.R.-G. would like to thank the Natural Sciences and Engineering Research Council of Canada (NSERC) for financial support through a Discovery Grant (RGPIN-2017-06611) and the Canadian Foundation for Innovation (CFI).

**AUTHOR CONTRIBUTIONS**

N.L. and S.W. conceived and designed the experiments. N.L. synthesized and characterized the monomers and polymers. N.L., Y.D., and Y.L. fabricated the transistor devices and made the measurements. J.S., Q.Z., Y.W., and X.G. helped with the GIXD characterizations. S.D., Q.S., and N.O. participated in the discussion of the results. B.O. and S.R.-G. performed the polymer molecular weight analysis. J.X. gave help on materials characterization and discussion. N.L. and S.W. cowrote the paper. All authors reviewed and commented on the manuscript.

**DECLARATION OF INTERESTS**

The authors declare no competing interests.

Received: April 26, 2021
Revised: June 25, 2021
Accepted: July 8, 2021
Published: August 4, 2021

**REFERENCES**


