Performance Comparisons of Polymer Semiconductors Synthesized by Direct (Hetero) Arylation Polymerization (DHAP) and Conventional Methods for Organic Thin Film Transistors and Organic Photovoltaics

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Abstract: C-C bond forming reactions are central to the construction of π-conjugated polymers. Classical C-C bond forming reactions such as the Stille and Suzuki coupling reactions have been widely used in the past for this purpose. More recently, direct (hetero)arylation polymerization (DHAP) has earned a place in the spotlight with an increasing number of π-conjugated polymers being produced using this atom-economic and more sustainable chemistry. As semiconductors in organic electronics, the device performances of the polymers made by DHAP are of great interest and importance. This review compares the device performances of some representative π-conjugated polymers made using the DHAP method with those made using the conventional C-C bond forming reactions when they are used as semiconductors in organic thin film transistors (OTFTs) and organic photovoltaics (OPVs).

Keywords: direct arylation; organic electronics; organic thin film transistors; organic photovoltaics; green chemistry

1. Introduction

Organic electronics enabled by polymer semiconductors is an exciting field of research that promises to pave the way for low-cost, flexible electronic devices such as organic thin film transistors (OTFTs) [1,2] and organic photovoltaics (OPVs) [3,4]. The performances of these devices have improved greatly in the past few years and has now become commercially viable, i.e., the field effect mobilities and the power conversion efficiencies for the polymer based OTFTs and OPVs have exceeded 10 cm² V⁻¹ s⁻¹ [5–8] and 10% [9–11], respectively. While studies in this field have been more performance driven, the scale-up synthesis of polymer semiconductors has been attracting increased attention. In particular, novel synthetic methodologies that can produce high performance polymer materials in a more environmentally friendly way, i.e., using “green chemistry” [12–15], with increased atom economy and reduced production costs are highly desirable.

Central to the construction of polymer semiconductors is the C-C bond forming reaction that links the monomeric units. Common methods for the C-C bond formation, such as Stille and Suzuki coupling reactions require an aryl halide and an aromatic compound with a reactive directing group, e.g., a boronic acid (or ester) for the Suzuki coupling and an organostannyl group for the Stille coupling. These reactions, while highly effective in C-C bond formation, require additional steps to install the directing groups, which increases the production cost and generates stoichiometric amounts
of by-products that are potential health and environmental hazards. Specifically, the organotin compounds formed from the Stille coupling reactions are known to be highly toxic [16], while the boronic acid derivatives used in the Suzuki coupling reactions, which were previously assumed to be less harmful, have been recently found to be potential genotoxic hazards [17].

To address these issues associated with the common synthetic methods used to prepare polymer semiconductors, a novel C-C bond forming methodology, the so-called direct (hetero) arylation polymerization (DHAP) has been explored recently (Figure 1) [12]. The DHAP method eliminates the need for adding a directing group. Instead, the carbon atom with the most “active hydrogen” in the monomer is able to couple with the halogenated carbon atom in another (or the same) monomer. However, many monomer compounds have multiple C-H bonds with close dissociation energies, which can potentially be activated and react with a C-halogen bond. Furthermore, two Pd(II) complex intermediates bearing equal (hetero)aryl groups may undergo a disproportionation reaction, resulting in a homocoupling defect [18–20]. These side reactions may impede the formation of soluble (in the case of crosslinking side reaction) or high molecular weight (in the case of homocoupling side reaction) polymer products. Even for the polymers with good solubility and high molecular weights made by DHAP, a certain amount of branching, crosslinking, and/or homocoupling defects are frequently observed [21–26]. Figure 1 shows the formation of these defects in the DHAP of 3-alkyl-2-bromo-thiophene to poly(3-alkylthiophene). In the past few years, rigorous studies have been conducted to optimize the synthetic conditions to minimize or eliminate these side reactions. With a better understanding of the DHAP mechanism, a number of high-quality polymer semiconductors with fewer structural defects have been synthesized using the DHAP method [27].

An important question or concern from the organic electronics community is: are the performances of the polymers made by DHAP comparable to those of the polymers made by the conventional synthetic methods? In this review, we will provide a preliminary answer to this question by judiciously selecting some representative polymer semiconductors that were made by both the DHAP method and the conventional methods and compare their performances in OTFTs and OPVs. For the recent and significant progress made in clarifying the mechanism and optimizing reaction conditions of DHAP to improve the yield and molecular weight and reduce/eliminate structural defects of the resulting polymers, readers are directed to several recent review articles [12,28–30].

![Figure 1. Direct (hetero)arylation polymerization (DHAP) of a 2-bromo-3-alkylthiophene, showing the potential for forming homocoupling and branching defects.](image)

2. Polymer Semiconductors for Organic Thin Film Transistors

Poly(3-hexyl-thiophene-2,5-diyl) (P1, Figure 2), also known as P3HT, is one of the most widely studied polymer semiconductors to date. Due to its widespread use in the organic electronics community, it has become a standard, to which new materials can be compared [31]. The monomer unit, 3-hexyl-thiophene-diyl, is unsymmetrical, which means that three different types of dyads, heat-to-head (HH), head-to-tail (HT), and tail-to-tail (TT) can be formed depending on the positions of the hexyl substituents.
on the two thiophene units. In an HH dyad, the two closely positioned hexyl substituents would cause severe twisting of the two thiophene rings, which would disrupt the π-conjugation pathway along the polymer backbone. The TT dyad is highly coplanar, but the formation of each TT dyad is accompanied with the formation of one HH dyad during the polymerization. On the other hand, the HT dyad is highly coplanar and a P3HT comprising of only HT dyads is a ‘regioregular’ HT P3HT or commonly rr-P3HT. The first P3HT was made using an FeCl$_3$-mediated oxidative coupling polymerization, which contained a significant amount of HH dyads with an HT mol % or regioregularity of ~70–80% [32–35] and thus had a rather twisted backbone. P3HT made in this way has poor crystallinity, which leads to low charge carrier mobilities of $\sim 10^{-4}$ cm$^2$ V$^{-1}$ s$^{-1}$ in OTFT devices [36].

In 1992, McCullough et al. reported the use of Grignard reagents to produce an rr-P3HT with a high regioregularity of up to 98% [37,38]. Also reported in 1992 was the discovery of a synthetic methodology based on zinc reagents by Rieke et al. [39,40], which could produce rr-P3HT with regioregularity of >98%. In subsequent studies, regioregular poly(3-alkyl-thiophene-2,5-diyl) samples were also made using Suzuki coupling (HT mol % = 96–97%) [41], Stille coupling polymerization (HT mol % > 96%) [42], or Grignard metathesis (GRIM, HT mol % > 98%) [43,44]. The synthetic schemes for these methods are depicted in Figure 2. P3HT with a high HT mol % of 96% made by the Rieke method demonstrated a high hole mobility of up to $0.10$ cm$^2$ V$^{-1}$ s$^{-1}$ in OTFTs, which is several orders of magnitude higher than that for the P3HT with lower regioregularity prepared using FeCl$_3$ and Yamamoto couplings [36]. Thus, it would be no exaggeration to say that the birth of rr-P3HT has brought the study of organic electronics into a new era.

![Figure 2. Synthesis of rr-P3HT by (a) McCullough’s [38], (b) Rieke’s [40], (c) Grignard metathesis (GRIM) [43], (d) Stille coupling [42], (e) Suzuki coupling [41], and (f) DHAP [45] methods.](image-url)
The aforementioned synthetic routes to rr-P3HT require additional synthetic steps to add a directing group and/or produce stoichiometric harmful by-products. Thus, it was of interest to researchers to explore alternative synthetic pathways for forming highly regioregular P3HT without the above drawbacks.

In 1998, Lemaire and co-workers reported a new method to synthesize polythiophenes using 2-iodothiophenes as the monomers and Pd(OAc)$_2$ as the catalyst [46]. The authors initially thought the polymerization was a Heck-type reaction, but it was later found to be a direct (hetero) arylation reaction, which involves a direct coupling of an active aromatic C-H bond with an aromatic C-X bond (X is halide) to form a C-C bonding motif [47]. In 2010, Wang et al. conducted a DHAP of 2-bromo-3-hexylthiophene with Herrmann’s catalyst to make an rr-P3HT [45]. High number average molecular weights ($M_n$) of up to 31 kDa were achieved, which are much higher than those of the polythiophenes (10$^3$ Da) through DHAP reported previously by Lemaire and coworkers [46]. A very high degree of regioregularity (>98%) of this rr-P3HT was confirmed by the $^1$H NMR spectroscopic analysis, indicating that electronic grade P3HT was attainable using DHAP synthetic methods.

Pouliot et al. [48] prepared a sample of P1 with a very high regioregularity of >99.5% and an $M_n$ of 33 kDa by optimizing the synthesis of the 2-bromo-3-hexylthiophene monomer and DHAP conditions (P1$_{DHAP}$, Table 1). They compared the OTFT performance of this polymer with other rr-P3HT polymers made by GRIM and Rieke’s methods using the same device configuration and testing conditions (P1$_{GRIM}$ and P1$_{Rieke}$, Table 1). It was found that high hole mobilities of up to 0.19 cm$^2$ V$^{-1}$ s$^{-1}$ were achieved by the polymer made by DHAP, which are much higher than those of P1$_{GRIM}$ (0.11 cm$^2$ V$^{-1}$ s$^{-1}$) and P1$_{Rieke}$ (up to 0.02 cm$^2$ V$^{-1}$ s$^{-1}$). The mobility trend can be explained by their regioregularity order: P1$_{DHAP}$ (HT mol % > 99.5%) > P1$_{GRIM}$ (HT mol % = 98.0%) > P1$_{Rieke}$ (HT mol % = 95.5%). The results from this study show that the performance of rr-P3HT made by DHAP in OTFTs is improved compared to those of the rr-P3HT polymers made by the Rieke and GRIM methods.

A donor-acceptor (D-A) type polymer, poly [[N,N$'$-bis(2-octyldodecyl)-naphthalene-1,4,5,8-bis(dicarboximide)-2,6-diyl]-alt-5,5$'$-(2,2$'$-bithiophene)], (P2, Figure 3) also known as N2200, is considered to be the most extensively investigated n-type polymer for OTFTs [49–51] and OPVs [52–57]. P2 has commonly been synthesized using the Stille coupling polymerization [58], so the ability to synthesize it using atom-economical methods is of interest. In 2015, the DHAP synthesis of N2200 was reported by Sommer et al. [50] (P2$_{DHAP}$, Table 1). Optimized DHAP conditions gave the polymer with an $M_n$ of up to 31 kDa and a ‘perfect’ NDI-bithiophene alternating structure, with no homocoupling, branching or cross-linking, according to $^1$H NMR spectroscopic analysis. Importantly, the molecular weight of the polymer was readily controlled via in situ solvent (toluene) end capping by adjusting the monomer concentration. Another important observation was that the optical and thermal properties of the DHAP polymers reach their maxima at $M_n$ ~20 kDa. This is in excellent agreement with the sample produced via the Stille coupling (P2$_{Stille}$, Table 1). In OTFTs, the mobilities of P2$_{DHAP}$ (2.9 cm$^2$ V$^{-1}$ s$^{-1}$) and P2$_{Stille}$ (3.2 cm$^2$ V$^{-1}$ s$^{-1}$) with similar $M_n$ were very close, which are among the highest values reported for n-type polymers to date (OTFT transfer curves for P2$_{DHAP}$ and P2$_{Stille}$ are shown in Figure 4).

Another naphthalene-1,4,5,8-bis(dicarboximide) (NDI) based polymer P3 (Figure 3), which is composed of NDI, thiophene and tetrafluorobenzene, was synthesized by DHAP (P3$_{DHAP}$, Table 1) [59]. A rather low $M_n$ of 7.8 kDa was obtained. A detailed $^1$H NMR spectroscopic analysis showed the absence of β-arylation, indicating that this polymer has a linear structure. Despite its rather low molecular weight, P3$_{DHAP}$ showed high electron mobilities of up to 1.3 cm$^2$ V$^{-1}$ s$^{-1}$ when being used as an n-channel semiconductor in OTFTs. The same polymer synthesized by other methods has not been reported.
Diketopyrrolopyrrole (DPP) or pyrrolo[3,4-c]pyrrole-1,4(2H,5H)-dione is a strong acceptor building block, which has been used extensively for the construction of a large number of ultrahigh mobility donor-acceptor (D-A) type polymer semiconductors for OTFTs [60]. However, the most high-performing DPP-based polymer materials for OTFTs such as PDBT-co-TT [5,61], PDQT [62], PDPP-TVT [63], and P-29-DPPDTSE [64] have not been successfully synthesized by DHAP yet. For example, an attempt to prepare PDPP-TVT by DHAP of 3,6-bis(5-bromothiophen-2-yl)-2,5-bis(2-decytritetradecyl) pyrrolo[3,4-c]pyrrole-1,4-(2H,5H)-dione (DPP) and (E)-1,2-bis(thiophen-2-yl)ethane (TVT) under typical conditions failed to produce the target polymer [65]. Interestingly, when a fluorinated TVT, (E)-1,2-bis(3,4-difluorothiophen-2-yl)ethane, 4FTVT), was used, a polymer PDPP-4FTVT (P4) with a high $M_n$ of up to 60 kDa was successfully obtained (P4_DHAP, Table 1). This new polymer showed an ambipolar charge transport behavior with high hole and electron mobilities of up to 3.4 cm$^2$ V$^{-1}$ s$^{-1}$ and 5.9 cm$^2$ V$^{-1}$ s$^{-1}$, respectively, in OTFTs. This study suggests that fluorine substitution might be able to suppress the β-arylation side reactions and/or activate the α-H of the 4FTVT monomer.

Another copolymer of DPP with tetrafluorobenzene, PDPPTh2F4 (P5), was synthesized by DHAP [66], which has not been approached by other synthetic methods (P5_DHAP, Table 1). Due to the strong electron withdrawing effect of the tetrafluorobenzene units, this polymer showed n-type electron transport semiconductor behavior, opposed to the majority of other DPP polymers that
showed either p-type or ambipolar charge transport performance. High electron mobility of up to 0.60 cm² V⁻¹ s⁻¹ was achieved in OTFTs.

A copolymer of DPP and bithiazole, PDBTz-24 (P6), was synthesized by Guo et al. using DHAP of dibrominated bithienyl DPP and 2,2′-bithiazole monomers (P6DHAP, Table 1) [67]. ¹H NMR spectra showed some unidentified structural defects possibly resulting from homocoupling and/or branching. Nonetheless, this polymer exhibited an electron transport dominant ambipolar performance with hole and electron mobilities of up to 0.06 cm² V⁻¹ s⁻¹ and 0.53 cm² V⁻¹ s⁻¹ in OTFTs, outperforming a similar polymer PDBTz-27 (P6Stille) with 5-decylheptadecyl side chains made by the Stille coupling previously reported by Reichmanis et al. [68], which showed electron mobilities of up to 0.31 cm² V⁻¹ s⁻¹.

A DPP-DPP type polymer (P7) was synthesized via DHAP between 3,6-bis(thiophen-2-y1)-2,5-bis(dodecyl)pyrrolo[3,4-c]pyrrole-1,4(2H,5H)-dione and 2,5-bis(2-octyldodecyl)-3,6-bis(5-bromo-4-methylthiophene-2-yl)pyrrolo[3,4-c]pyrrole-1,4(2H,5H)-dione by Pouliot et al. (P7DHAP, Table 1) [24]. A high M₀ of 46 kDa along with a narrow dispersity (Mₚ/M₀) of 2.5 was obtained, which suggested that negligible side reactions occurred. P-type semiconductor performance with high hole mobilities of up to 1.2 cm² V⁻¹ s⁻¹ was achieved in OTFTs. Interestingly, when the reactive C-H and C-Br exchanged between the two monomers, that is, 2,5-bis(2-octyldodecyl)-3,6-bis(4-methylthiophene-2-yl)-pyrrolo[3,4-c]pyrrole-1,4(2H,5H)-dione and 3,6-bis-(5-bromo thiophen-2-yl)-2,5-bis(2-dodecyl)pyrrolo[3,4-c]pyrrole-1,4(2H,5H)-dione were polymerized, the resulting polymer had a lower M₂ (15 kDa) and showed lower hole mobility (0.26 cm² V⁻¹ s⁻¹), which was ascribed to some ill-defined couplings such as homocoupling and branching.

Table 1. Comparison of Polymeric Materials Used in Thin Film Transistors.

| ID  | Name          | Synthesis | OTFT Performance |
|-----|---------------|-----------|------------------|
|     |               | M₀, kDa (HT mol %) | Mₚ/M₀ | Yield, % | Device | uᵣ/uₑ | cm² V⁻¹ s⁻¹ | Iₒ/Iₘ off | Vₒₜ, V | Ref.   |
| P1DHAP | P3HT        | 33 (>99.5)       | 1.8 | 96      | BGBC   | 0.19/- | 1000/-       | -/-          | -/-      | 2016 [48] |
| P1DHAP | P3HT        | 33 (98.0)        | 1.5 | NA      | BGBC   | 0.31/- | 1000/-       | -/-          | -/-      | 2016 [48] |
| P1Stille | P3HT      | 25 (95.5)        | 1.9 | NA      | BGBC   | 0.02/- | 1000/-       | -/-          | -/-      | 2016 [48] |
| P1DHAP | N2200       | 31              | 2.9 | 99      | TGBP   | -/2.9 | <1000/-      | -/-          | -/-      | 2015 [50] |
| P1DHAP | N2200       | 32              | 5.4 | 100     | TGBP   | -/3.2 | <1000/-      | -/-          | -/-      | 2015 [50] |
| P1DHAP | N2200       | 7.8             | 1.7 | -       | TGBP   | -/1.3 | <10⁻¹/-       | -/-          | -/-      | 2014 [59] |
| P1DHAP | P(ThNDThF₄) | 7.8             | 1.7 | -       | TGBP   | -/1.3 | <10⁻¹/-       | -/-          | -/-      | 2014 [59] |
| P3DHAP | PDDP        | 18              | 3.8 | 66      | TGBP   | 0.06/- | 10⁻³/-        | 10⁻⁵/-       | -/-      | 2015 [67] |
| P3DHAP | PDDP        | 24              | 3.8 | 66      | TGBP   | 0.06/- | 10⁻³/-        | 10⁻⁵/-       | -/-      | 2015 [67] |
| P3DHAP | PDDP        | 46              | 2.5 | 84      | BGBC   | 1.2/- | 10⁻⁵/-        | 0/-          | -/-      | 2015 [24] |

3. Polymer Semiconductors for Organic Photovoltaics

The use of rr-P3HT (P1) synthesized by DHAP in solar cells was first reported by Thompson et al. [25]. A sample of rr-P3HT (P1DHAP) with M₀ = 19 kDa, Mₚ/M₀ = 2.0, and HT mol % = 90% and a control polymer P1Stille with M₀ = 19 kDa, Mₚ/M₀ = 2.7, and HT mol % = 93% were synthesized by DHAP and Stille coupling methods, respectively (Table 2). Interestingly, it was shown that P1DHAP had ~0.75% β-defects, but still showed even better solar cell performance compared to P1Stille, which had no β-defects (PCE: 2.70% for P1DHAP vs. 2.30% for P1Stille). The authors made another sample using DHAP, having 1.41% β-defect concentration, and found that the performance was poorer due to formation of unfavorable morphology when blended with fullerene and the low space-charge-limited current (SCLC) mobility, which resulted in low Jₜsc. The authors concluded that rr-P3HT synthesized by DHAP with β-defect concentrations below 1% were able to achieve similar or better performance in solar cell devices when compared with their counterparts made by the Stille coupling method.

Another recent study by Thompson et al. compared the properties of rr-P3HT synthesized using a “green solvent assisted” DHAP (P1DHAP) with a sample prepared by the traditional Stille coupling method (P1Stille, Table 2) [69]. P1DHAP had M₀ = 20 kDa, Mₚ/M₀ = 2.1, and HT mol % = 96.2%,
while $P_{1\text{Stille}}$ had $M_n = 18$ kDa, $M_w/M_n = 2.4$, and HT mol % = 92.9%. These authors found that the use of a bulky carboxylic acid additive, neodecanoic acid, and a “green” solvent, 2-methyl-tetrahydrofuran, was critical in completely avoiding the formation of $\beta$-defects. In UV-Visible absorption spectra, the $\beta$-defect free $P_{1\text{DHAP}}$ displayed a higher extinction coefficient at the wavelength of maximum absorbance ($\lambda_{\text{max}}$) and stronger vibrational fine structure, indicating more structural ordering in the solid state. $P_{1\text{DHAP}}$ also had larger crystalline coherence lengths, as confirmed by X-ray diffraction, which were correlated to the increased SCLC hole mobility for this polymer. When evaluated as a donor with PC$_{61}$BM as the acceptor in solar cells, $P_{1\text{DHAP}}$ outperformed $P_{1\text{Stille}}$ (PCE: 3.28% vs. 2.86%, Table 2). This is in agreement with the previous results by the same group, implying that a $\beta$-P3HT sample with negligible $\beta$-defects produced by DHAP can outperform its counterpart made by the Stille coupling [25].

Kanbara et al. synthesized copolymers of fluorene and EDOT ($P_8$, Figure 5) using both DHAP and Suzuki coupling methods ($P_{8\text{DHAP}}$ and $P_{8\text{Suzuki}}$, Table 2) [70]. $P_{8\text{DHAP}}$ made by DHAP with a microwave reactor had a much higher $M_n$ of 150 kDa than that of $P_{8\text{Suzuki}}$ made by Suzuki coupling, which had an $M_n$ of 17 kDa. As a donor, $P_{8\text{DHAP}}$ demonstrated a high PCE of 4.08%, while $P_{8\text{Suzuki}}$ exhibited a very low PCE of 0.480%. The large discrepancy in the OPV performance of these two polymers is ascribed mainly to their different hole mobilities. In OTFTs, $P_{8\text{DHAP}}$ showed an average hole mobility of $1.2 \times 10^{-3}$ cm$^2$/V·s, while the average hole mobility of $P_{8\text{Suzuki}}$ is about two orders of magnitude lower at $3.2 \times 10^{-5}$ cm$^2$/V·s. Another $P_{8\text{DHAP}}$ with a lower $M_n$ of 48 kDa showed a lower PCE of 2.55%, which was believed to be due to the presence of terminal Br groups and residual Pd impurities in the polymers, causing a decrease in the hole mobility ($7.7 \times 10^{-4}$ cm$^2$/V·s$^{-1}$).

In a later study by the same group, $P_{8\text{DHAP}}$ showed further improved PCE of up to 4.60% after the terminal Br groups were removed through a post-polymerization treatment of the polymer with sodium $N,N$-diethylidithiocarbamate [71].

D-A polymers are among the most popular materials for use in OPVs and synthesis of high-quality D-A polymers by DHAP was therefore explored. Russel et al. reported on two D-A polymers based on DPP, DPP-co-phenylene (DPDP-TPT, $P_9$, Figure 5) and DPP-co-thiophene, (DPDP3T, $P_{10}$), which were synthesized by DHAP ($P_{9\text{DHAP}}$ and $P_{10\text{DHAP}}$, Table 2) [22]. Optimized DHAP conditions provided a sample of $P_{9\text{DHAP}}$ with an $M_n$ of 14 kDa and an $M_w/M_n$ of 1.8. $P_{9\text{DHAP}}$ produced via DHAP in this work was compared with one produced by Janssen’s group using a Suzuki coupling protocol ($P_9\text{Suzuki}$, Table 2) [72], which showed a bimodal GPC trace with two peak molecular weights of 65 kDa and 10 kDa due to the gelation and aggregation. As a donor, $P_{9\text{DHAP}}$ showed slightly lower PCE in OPVs than $P_9\text{Suzuki}$ when PC$_{61}$BM was used as the acceptor (4.37 vs. 5.50%, Table 2). Later, optimization of the Suzuki coupling reaction gave a sample of $P_9\text{Suzuki}$ with a higher $M_n$ reaching 72 kDa and an $M_w/M_n$ of 1.98. Solar cells using $P_9\text{Suzuki}$ gave PCE of up to 7.40% [73]. For the second polymer reported in this work [22], DPDP3T, optimized DHAP conditions gave a sample of $P_{10\text{DHAP}}$ with $M_n$ 29 kDa and an $M_w/M_n$ of 3.8. A previous paper by Janssen et al. in 2009 [74] reported on the synthesis of DPDP3T using a Suzuki coupling protocol, which gave $P_{10\text{Suzuki}}$ with an $M_n$ of 54 kDa, nearly twice that of $P_{10\text{DHAP}}$ reported in this work. In OPVs, $P_{10\text{DHAP}}$ was also outperformed by $P_{10\text{Suzuki}}$ (4.01% vs. 4.69%, Table 2). A later work by Janssen et al. improved the synthesis of $P_{10}$ via Stille coupling [73] ($P_{10\text{Stille}}$, Table 2), achieving an improved $M_n$ of 150 kDa and an $M_w/M_n$ of 2.72. Solar cells made using $P_{10\text{Stille}}$ gave much better PCE of up to 7.10% when used as the donor material in OPVs. $^1$H NMR spectroscopy was unable to determine the presence of homocoupling defects in $P_{9\text{DHAP}}$ and $P_{10\text{DHAP}}$ due to the tendency to aggregate in solution; however, the authors suggested that some homocouplings (<5%) may be present in these samples because a broadening of the low energy absorption shoulders was seen, a feature which is often attributed to minor homocoupling defects [75]. Both the presence of the homocoupling defects as well as the lower molecular weights might account for the lower mobilities observed for $P_{9\text{DHAP}}$ and $P_{10\text{DHAP}}$ compared with their counterparts made by the Suzuki and Stille couplings, $P_9\text{Suzuki}$, $P_9\text{Stille}$, $P_{10\text{Suzuki}}$ and $P_{10\text{Stille}}$ which contained lesser amounts of defects and had higher molecular weights.
Figure 5. Materials made using DHAP and conventional methods, evaluated in organic photovoltaic (OPV) devices.

Poly[4,4-bis(2-ethylhexyl)-4H-cyclopenta[2,1-b;3,4-b’]dithiophene-2,6-diyl-alt-2,1,3-benzothiadiazole-4,7-diyl] (PCPDTBT, P11) is a high performance polymer semiconductor, used commonly as a donor in OPVs [76–78], and a p-type semiconductor in OTFTs [79]. In 2012, Horie et al. synthesized PCPDTBT using both DHAP and Suzuki methods (P11_DHAP, P11_Suzuki, Table 2) for a direct comparison [80]. P11_DHAP had an $M_n$ of 72 kDa with an $M_w/M_n$ of 4.52, while P11_Suzuki had a much lower $M_n$ of 15 kDa with an $M_w/M_n$ of 2.1. By using MALDI-TOF
and ^1^H NMR spectroscopic analysis, it was found that P11\textsubscript{DHAP} contained both homocoupling and branching defects, while P11\textsubscript{Suzuki} did not. The \( \beta \)-defect concentration was found to be 12–13% for P11\textsubscript{DHAP}, which is well above the “threshold” value of ~1% for rr-P3HT for achieving good OPV performance observed by Thompson et al. \cite{25}. Despite such high \( \beta \)-defect concentration, P11\textsubscript{DHAP} exhibited higher PCE in solar cells using PC\textsubscript{71}BM as the acceptor under the same processing and testing conditions compared to P11\textsubscript{Suzuki} (PCE: 3.98% vs. 3.74%, respectively) \cite{80}. P1\textsubscript{1}\textsubscript{Suzuki} synthesized by the Stille coupling previously, which had an \( M_n \) of 28 kDa, achieved a lower PCE of 3.50% \cite{77} (P1\textsubscript{1}\textsubscript{Suzuki}, Table 2). These results suggest that the PCPDITBT system has a high tolerance for \( \beta \)-defects, which might be due to the beneficial effect of the branching structure on the morphology of the P1\textsubscript{1}\textsubscript{DHAP}-PC\textsubscript{71}BM blend films \cite{80} as well as the much higher intrinsic hole mobility of this polymer \cite{79} compared to P3HT. The fact that P1\textsubscript{1}\textsubscript{DHAP} showed the best PCE is most likely due to its highest molecular weight, which is a more dominant factor than the \( \beta \)-defect concentration in the OPV performance. Given that the rigorous optimization of device fabrication for commercial samples of P1\textsubscript{1}\textsubscript{DHAP} may also exhibit a higher PCE when the active layer formation is further optimized.

Thompson et al. synthesized benzothiadiazole-co-benzodithiophene polymers (P12) using DHAP and Stille methods (P12\textsubscript{DHAP}, P12\textsubscript{Suzuki-LMW}, P12\textsubscript{Suzuki-HMW}, Table 2) \cite{81}. The optimized DHAP protocol afforded a ‘defect-free’ PPDITBT sample P12\textsubscript{DHAP} with an \( M_n \) of 15 kDa and an \( M_w/M_n \) of 2.1 (P12\textsubscript{DHAP}, Table 2). The use of THF as a solvent and the bulky neodecanoic acid as an additive was found to be essential in minimizing structural defects during DHAP based on the \(^1^H\) NMR spectroscopic analysis \cite{69}. Two PPDITBT samples having \( M_n \) of 59 and 16 kDa and \( M_w/M_n \) of 3.3 and 2.1, respectively, were synthesized by the Stille method for comparison (P12\textsubscript{Suzuki-HMW} vs. P12\textsubscript{Suzuki-LMW}, Table 2). P12\textsubscript{DHAP} showed a better PCE (3.40%) compared with the counterpart P12\textsubscript{Suzuki-LMW} made by Stille (PCE = 2.90%), which has a similar \( M_n \) and \( M_w/M_n \). On the other hand, polymer P12\textsubscript{Suzuki-HMW} with a high molecular weight made by the Stille method was able to achieve a higher PCE of 3.80%. These results illustrate, once again, the importance of the polymer molecular weight on device performance. It can be concluded that the quality of the polymer P12 made by DHAP is similar to that made from Stille coupling if the molecular weight can be matched.

In 2015, Farinola et al. reported on the DHAP synthesis of a random copolymer (P13), which used benzo[\{1,2,5\}thiadiazole or benzo[d\{1,2,3\}triazole as the accepting units and benzo[1,2-b:4,5-b’]dithiophene as the donor component \cite{23}. Samples of P13 were prepared via DHAP and also by Stille coupling polymerization for direct comparison. DHAP yielded a sample (P13\textsubscript{DHAP}) with a low \( M_n \) of only 10 kDa compared to the \( M_n \) of 20 kDa obtained for P13\textsubscript{Suzuki} synthesized via Stille coupling. In addition, the \( M_w/M_n \) was much larger for the DHAP synthesized sample compared to the one made using a Stille coupling (\( M_w/M_n \): 7.6 for P13\textsubscript{DHAP} vs. 3.1 for P13\textsubscript{Suzuki}, Table 2). A hypsochromic shift in the absorption spectra is noticed for the DHAP synthesized polymer, indicative of a lower effective conjugation length. \(^1^H\) NMR analysis revealed the presence of \( \beta \)-defects for P13\textsubscript{DHAP}. When the two polymers were tested in solar cells with PC\textsubscript{71}BM as the acceptor, P13\textsubscript{DHAP} gave a lower PCE of 2.80% compared to a value of 4.80% for P13\textsubscript{Suzuki}. Such a large difference in performance between P13\textsubscript{DHAP} and P13\textsubscript{Suzuki} can be directly ascribed to the lower molecular weight and higher defect concentration in the former. Optimization of the catalytic systems for DHAP such as the use of bulky carboxylic acid additives and specialized catalysts \cite{27,69,82} may provide a sample of P13 with a lower concentration of defects and thus a higher effective conjugation length, which would improve its performance to be on par with P13\textsubscript{Suzuki}.

In 2016, Marks et al. reported a comprehensive study to compare the physical and electronic properties of a series of D-A polymers made using both DHAP and Stille methods \cite{83}. The first polymer is PBDTT-FTTE (P14), a well-known high performance p-type polymer that has been used in record setting OPVs \cite{84,85}. It was found that increasing the steric bulkiness of the carboxylic acid additive, i.e., the use of 2,2-diethylhexanoic acid instead of pivalic acid, in the DHAP system was critical in forming high molecular weight, defect-free polymers, which is in agreement with other
recent reports [69,81]. Optimization of DHAP conditions gave P14$_{\text{DHAP}}$ with $M_n = 25$ and $M_w/M_n = 2.2$, a very close match with P14$_{\text{Stille}}$ made by the Stille coupling ($M_n = 25$ kDa and $M_w/M_n = 2.2$). When evaluated in solar cell devices with PC$_{71}$BM as the acceptor, P14$_{\text{DHAP}}$ gave an almost same PCE compared with P14$_{\text{Stille}}$ (PCE: 8.36% vs. 8.40%, Table 2 and Figure 6a). In addition, the two polymers showed very similar film morphologies when blended with PC$_{71}$BM (Figure 6b). Two other pairs of D-A polymers based on thieno[3,4-c]pyrrole-4,6-dione (TPD), namely PBDTT-TPD (P15$_{\text{DHAP}}$ and P15$_{\text{Stille}}$, Table 2) and PTTPD3T (P16$_{\text{DHAP}}$ and P16$_{\text{Stille}}$, Table 2), were also made using both DHAP and Stille methods [83]. It was found that P15$_{\text{DHAP}}$ has a higher $M_n$ than that of P15$_{\text{Stille}}$ ($M_n$: 30 kDa vs. 15 kDa), while P16$_{\text{DHAP}}$ had a lower $M_n$ than that of P16$_{\text{Stille}}$ ($M_n$: 19 kDa vs. 30 kDa). $^1$H NMR spectroscopy confirmed that the structural regularity of both polymers made by DHAP was similar to that of the ones made by the Stille method. In addition, the absorption spectra of each pair of polymers are nearly identical. In OPVs, P15$_{\text{DHAP}}$ was found to outperform P15$_{\text{Stille}}$ (PCE: 5.84% vs. 5.20%), while P16$_{\text{DHAP}}$ performed slightly poorer than P16$_{\text{Stille}}$ (PCE: 7.20% vs. 7.38%). Since P15$_{\text{DHAP}}$ and P16$_{\text{DHAP}}$ did not contain noticeable structural defects, the differences in their OPV performance with their counterparts P15$_{\text{Stille}}$ and P16$_{\text{Stille}}$ are likely related to their different molecular weights, with the higher molecular weight ones showing higher OPV performance.

PBDTTPD (P17) is another high performance donor polymer for OPVs [86]. Farinola et al. conducted a study comparing the optoelectronic properties and device performance of PBDTTPD made by both DHAP and Stille methods (P17$_{\text{DHAP}}$ and P17$_{\text{Stille}}$, Table 2) [87]. Optimized DHAP conditions produced P17$_{\text{DHAP}}$ with a low $M_n$ of 12 kDa, while P17$_{\text{Stille}}$ synthesized via the Stille coupling protocol had an even lower $M_n$ of 10 kDa. The slightly higher $M_n$ of P17$_{\text{DHAP}}$ resulted in a red-shifted UV-Visible absorption profile as well a clearer vibronic structure in the low-energy region compared to P17$_{\text{Stille}}$. In addition, the aggregation propensity of P17$_{\text{DHAP}}$ in solution was noted to be higher. Furthermore, I I was found that P17$_{\text{Stille}}$ allowed the formation of large micro-sized domains of PC$_{71}$BM in the bulk-heterojunction blend, while P17$_{\text{DHAP}}$ did not. OPV devices using these two polymers as the donors and PC$_{71}$BM as the acceptor were fabricated and compared. PCEs of up to 5.31% and 4.82% were obtained for P17$_{\text{DHAP}}$ and P17$_{\text{Stille}}$, respectively, which is in good agreement with the disparity in their molecular weights.

Figure 6. (a) J-V curves for Stille and DHAP produced PBDTT-FTTE with PC$_{71}$BM; (b) AFM Images comparing the morphology of Stille and DHAP samples of P6, blended with PC$_{71}$BM. Reproduced with permission from (J. Am. Chem. Soc. 2016, 138, 15699–15709). Copyright (2016) American Chemical Society.

A copolymer of TPD and cyclopenta[2,1-b:3,4-b’]dithiophene (P18) was synthesized by Kanbara et al. using an improved DHAP in toluene, which had an $M_n$ of 25 kDa with an $M_w/M_n$ of 1.9 (P18$_{\text{DHAP}}$, Table 2) [19]. The molecular weight is much higher than that ($M_n = 2.5$ kDa) of the same polymer made by DHAP by Harris et al. using DMAC as the solvent, where severe homocoupling occurred, resulting in the lower $M_n$ [80]. On the other hand, the $^1$H NMR and MALDI-TOF-MS analysis of P18$_{\text{DHAP}}$ made in toluene in this work showed no detectable homocoupling defects. A high
PCE of up to 6.80% was achieved when \textbf{P18DHAP} was used as a donor in OPVs, which is higher than that (PCE = 5.20%) of \textbf{P18Stille} prepared by the Stille coupling, which had a lower $M_n$ of 15 kDa \cite{88}.

Ahmed et al. reported on another series of polymers based on [3,4-c]pyrrole-4,6-dione derivatives with sulfur, oxygen or selenium heteroatom substitution, made by the DHAP method \cite{89}. The champion polymer from their study, \textbf{P19DHAP}, was based on a seleno[3,4-c]pyrrole-4,6-dione acceptor and a dithienosilole donor and was obtained with an $M_n$ of 29 kDa and an $M_w/M_n$ of 1.6. When paired with PC$_{61}$BM in inverted organic solar cells, a high PCE of 7.13% was achieved. No equivalent polymer to \textbf{P19DHAP} made by a conventional coupling method was reported for comparison.

A recent report by Li et al. compared the properties of indacenodithiophene (IDT) and thiophene-quinoxaline-thiophene (TQ) derived polymers (\textbf{P20}) made using DHAP and Stille coupling methods (\textbf{P20DHAP} and \textbf{P20Stille}, Table 2) \cite{21}. \textbf{P20DHAP} had an $M_n$ of 27 kDa with an $M_w/M_n$ of 1.6, while \textbf{P20Stille} had an $M_n$ of 23 kDa and an $M_w/M_n$ of 1.5. The close number average molecular weights of the two samples allow for a fair comparison between the two polymers. $^1$H NMR spectroscopy, MALDI-TOF MS, and elemental analysis revealed that \textbf{P20Stille} contained a certain amount of TQ−TQ homocoupling defects, while \textbf{P20DHAP} had a very well-defined IDT-TQ alternating structure with no detectable structural defects. Interestingly, SCLC experiments showed a weak correlation between their hole mobility and structural regularity. Furthermore, grazing incidence X-ray diffraction (GIXRD) studies indicated no significant differences in the film crystallinity between two polymers. \textbf{P20DHAP} and \textbf{P20Stille} achieved PCE of 5.10% and 4.82%, respectively. It was found that the larger number of defects in the backbone of \textbf{P20Stille} had influenced the morphological properties and SCLC mobilities, which caused severe recombination and thus the lower PCE.
weights at 65 kDa and 10 kDa, respectively. Different monomers were used: eq. K P12

Different synthetic conditions were used. APO23

C /40 min for DHAP3a

Suzuki

DHAP Stille1b

ID Name Mα, kDa (HT %) Mα/Mω Yield, % Acceptor PCE (Best Reported) Jsc, mA cm⁻² Voc, V FF, % Ref.
P1DHAP a P3HT 19 (90.0) 2.0 - PC61 BM 2.70 8.70 0.62 50.0 2013 [25]
P1Silile b P3HT 19 (93.0) 2.7 - PC61 BM 2.81 8.37 0.61 45.0 2013 [25]
P1DHAP a P3HT 20 (98.2) 2.1 74 PC61 BM 3.28 9.40 0.59 59.1 2017 [69]
P1Silile b P3HT 18 (92.9) 2.4 68 PC61 BM 2.86 8.34 0.59 58.1 2017 [69]
P8DHAP PEDOTF 150 2.89 89 PC71 BM 4.08 9.41 0.83 52.0 2014 [70]
P8sazaki PEDOTF 17 2.08 85 PC71 BM 0.48 2.58 0.59 31.0 2014 [70]
P9DHAP PDPP-TPT 14 1.8 29 PC71 BM 4.37 13.3 0.77 41.5 2015 [22]
P8sazaki c PDPP-TPT 65 1.98 93 PC71 BM 4.50 14.0 0.80 67.0 2013 [73]
P10DHAP PDPP-3T 29 3.8 45 PC71 BM 4.01 10.3 0.71 56.2 2015 [22]
P10Silile PDPP-3T 54 3.15 84 PC71 BM 4.69 11.8 0.66 60.0 2009 [74]
P10Silile PDPP-3T 150 2.72 85 PC71 BM 7.50 15.4 0.67 69.0 2013 [73]
P11DHAP PCPDTBT 72 4.52 76 PC71 BM 3.98 13.9 0.63 45.5 2012 [80]
P11sazaki PCPDTBT 15 2.1 83 PC71 BM 3.74 12.7 0.64 43.8 2012 [80]
P11Silile PCPDTBT 28 1.5 61 PC61 BM 3.50 11.8 0.65 46.0 2007 [77]
P12DHAP P3DTBT 15 2.1 78 PC61 BM 3.40 10.5 0.72 45.0 2016 [81]
P12Silile PCPDTBT 59 3.3 79 PC61 BM 3.80 11.5 0.73 45.0 2016 [81]
P12Silile-5,5'-diketomethyleneidene d P3DTBT 16 2.1 70 PC61 BM 2.99 8.88 0.72 46.0 2016 [81]
P13DHAP - 10 7.6 70 PC71 BM 2.80 5.58 0.89 56.0 2015 [23]
P13Silile - 20 3.1 85 PC71 BM 4.80 9.89 0.81 60.0 2015 [23]
P14DHAP PDDBT-FFTTE 25 2.2 98 PC71 BM 8.36 15.5 0.78 68.8 2016 [83]
P14Silile PDDBT-FTTE 25 2.2 - PC71 BM 8.40 14.9 0.78 72.2 2016 [83]
P15DHAP PDDBT-TPD 30 2.7 76 PC71 BM 5.84 10.0 0.99 57.9 2016 [83]
P15Silile PDDBT-TPD 15 2.4 69 PC71 BM 5.20 9.10 0.99 58.7 2016 [83]
P16DHAP PTDPDT 19 2.0 83 PC71 BM 7.20 13.3 0.82 66.0 2016 [83]
P16Silile PTDPDT 30 1.8 94 PC71 BM 7.38 13.2 0.78 71.1 2016 [83]
P17DHAP BDTTPPD 12 - 80 PC71 BM 5.31 10.41 0.92 56.0 2016 [87]
P17Silile BDTTPPD 10 - 90 PC71 BM 4.82 9.17 0.93 57.0 2016 [87]
P18DHAP - 25 1.8 82 PC71 BM 6.80 10.13 0.89 53.5 2016 [87]
P18Silile - 15 1.2 52 PC61 BM 5.20 10.0 0.88 59.0 2013 [88]
P19DHAP PscPDST 29 1.6 57 PC61 BM 7.13 13.2 0.85 64.0 2015 [89]
P20DHAP IDT-TQ 27 1.6 71 PC61 BM 5.10 10.8 0.89 53.4 2016 [21]
P20Silile IDT-TQ 23 1.5 64 PC61 BM 4.82 10.4 0.89 52.1 2016 [21]

Table 2. Comparison of Polymeric Materials Used in Solar Cells.

a Different synthetic conditions were used. P1DHAP: 1 mol % Pd(OAc)₂/pivalic acid/K₂CO₃/DMAC/45 °C, 72 h; P1Silile: 1 mol % Pd₂(dbta)₂/P(o-MeOPh)₃/neodecanoic acid/C₆H₄CO₂/MethyTHF/120 °C, 12 h; b Same synthetic conditions (Pd(PPh₃)$_3$/DMF, 95 °C, 48 h) were used. Similar device fabrication procedures, except for the annealing conditions (110 °C /40 min for P1Silile vs. 150 °C/30 min for P1Silile2), were used; c Different synthetic conditions: P9sazaki: 5 mol % Pd₂(dbta)₂/11 mol % PPh₃/4 eq. K₂PO₄/Aliquat 336, toluene-water (8:1)/115 °C, 72 h; P9sazaki: 3 mol % Pd₂(dbta)₂/12 mol % PPh₃/5 eq. K₂PO₄/Aliquat 336, toluene-water (6:1), 115 °C, 16 h; d Different monomers were used: P12silile-5,5'-diketomethyleneideneidene: (5,5'-2,5-bis(2-phenylcdecyl)oxy)-1,4-phehylene)bis(thiophene-5,2-diyl)bis(trimethylstannane) and 4,7-dibromobenzimidazol[1,2,5]thiazadiazole; P12silile-5,5'-diketomethyleneideneidene: (5,5'-2,5-bis(2-phenylcdecyl)oxy)-1,4-phehylene)bis(thiophene-5,2-diyl)bis(trimethylstannane) and 4,7-dibromobenzimidazol[1,2,5]thiazadiazole; A bimodal GPC trace shows two peak molecular weights at 65 kDa and 10 kDa, respectively.
4. Conclusions

Recently, direct (hetero)arylation polymerization (DHAP) of a halogenated (hetero)aryl compound with another (hetero)aryl compound possessing activated C-H bonds has emerged as an alternative approach to the conventional C-C bond formation methods such as Stille and Suzuki coupling polymerization for the synthesis of linear π-conjugated polymers. Since DHAP does not require the use of a (hetero)aryl compound possessing a reactive directing group such as an organostannyl group in the case of a Stille coupling polymerization, fewer steps are needed for DHAP, which can improve the atom economy and reduce the cost for the synthesis of conjugated polymers. Furthermore, the by-product of a DHAP is a hydrogen halide (HX), which is much easier to treat compared with the toxic wastes generated from the Stille and Suzuki coupling polymerization. Therefore, DHAP has been considered a more cost-effective and “greener” technique for the mass production of conjugated polymers. However, since many (hetero)aryl compounds have more activated C-H bonds than required for the formation of a linear polymer, side reactions at the unwanted C-H bonds result in branching or cross-linking structural defects, often leading to degradation of the optoelectronic properties or even the formation of unprocessable insoluble products. In addition, homocoupling between the halogenated monomer molecules produces homocoupling defects in the polymer backbone and also results in the formation of lower molecular weight polymers. The presence of these structural defects in the polymers made by DHAP potentially have impacts on the performances of these polymers in organic electronic devices.

This review presented some exemplary conjugated polymer semiconductors made by DHAP and compared their performances in OTFTs and OPVs with their counterparts synthesized by conventional synthetic methods, such as the Stille and Suzuki coupling polymerization. It can be seen that by minimizing the structural defects and improving the molecular weight of the polymers through optimization of the DHAP conditions, similar or sometimes better OTFT and OPV performances could be achieved for the polymers made by DHAP in comparison to the ones prepared by the conventional methods. With the rapid progress made in the development of novel catalyst systems and optimization of reaction conditions (solvent, temperature, reaction time, etc.), it is expected that more and more high-performance polymer semiconductors, which have been synthesized and cannot be approached by conventional methods, will be developed. Further study of the atom-economic, cost-effective, and “greener” DHAP to enable the mass production of top-performing polymer semiconductors will propel the commercialization and wide-spread applications of organic electronics.

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