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Biosourced and metal-free synthesis of conjugated polymers: bPPV, bCN-PPV and **bPPTzTz**†

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The growing demand for sustainable materials has spurred interest in biosourced feedstocks and metalfree polymerization techniques for organic electronics. This study explores the synthesis of conjugated polymers using renewable succinic acid-derived monomers through environmentally benign polymerization methods. New pathways were developed to produce three biosourced polymers: poly(phenylene vinylene) (bPPV), poly(cyano-PPV) (bCN-PPV), and poly(thiazolo[5,4-d]thiazole) (bPPTzTz) using Gilch polymerization, Knoevenagel condensation, and Ketcham polymerization, respectively. Comprehensive characterization of these polymers, including molecular weight, optical, electrochemical, and thermal properties, demonstrates their potential for organic electronics. The polymers exhibit high renewable atom content, with polymer bPPV achieving 100% of its atoms coming from succinic acid. This work highlights the feasibility of integrating biosourced monomers and sustainable polymerization strategies, advancing the development of environmentally responsible materials for organic electronics.

Introduction

Organic electronics is a field that has been attracting significant attention in recent years. As society increasingly prioritizes sustainability, there is a growing drive to reduce reliance on critical minerals and replace them with more abundant organic materials. However, the majority of materials used in organic electronics are still derived from non-renewable sources, primarily petroleum¹ and they are commonly prepared using palladium-mediated coupling methods such as Stille, Suzuki and direct arylation polymerization.^{2,3} To address these challenges, it is crucial to identify sustainable sources of starting materials and to find suitable metal-free polymerization protocols.4

One promising alternative to fossil fuels is forest biomass, which consists of carbohydrates (such as cellulose and hemicellulose) and aromatics (such as lignin and tannins).5 Both of these components can be processed into added value compounds.⁶ Research into cellulosic-derived starting materials

has primarily focused on furan-substituted DPP,7 synthesized from succinic acid and furfural. Using lignin derived starting materials, significant progress has been made using vanillin for the synthesis of conjugated polymers. For instance, our group has previously reported the synthesis of heterocycle-substituted benzenes rings using vanillin as a precursor.8 In a subsequent study, we demonstrated their potential by polymerizing them through DArP with diketopyrrolopyrrole (DPP) for transistor applications.9 Additionally, the groups of Kayser10 and Brochon¹¹ have also explored vanillin-based monomers for synthesizing partially bio-sourced conjugated polymers, employing metal-free polymerization methods in both cases.

Since the last decade, only a few metal-free polymerization protocols have been reported. 12 Most of these methods rely on the same principle: the condensation of various nucleophiles with carbonyl groups to form alkene derivatives, examples include the aldol condensation,13 the Knoevenagel condensation,14,15 the Horner-Wadsworth-Emmons reaction, 16 and polyazomethine synthesis.¹⁷ However, other methods utilize a range of organic reactions to achieve conjugated polymers, such as Gilch polymerization, 18 the Ketcham reaction, 19 Yao et al.'s pyrazine construction, 20 and the multicomponent polymerization developed by Kayser et al. 10

This study seeks to explore the adaptation of metal-free polymerization techniques to biosourced monomers derived from succinic acid, a renewable feedstock readily obtainable

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from cellulose.²¹ By employing dimethyl succinate, we developed novel pathways to conjugated polymers with distinct properties and high renewable atom content. Our approach resulted in the synthesis of three biosourced polymers: poly(phenylene vinylene) (*bPPV*), cyano-PPV (*bCN-PPV*), and poly(thiazolo[5,4-*d*]thiazole) (*bPPTzTz*), *via* Gilch polymerization, Knoevenagel condensation, and Ketcham polymerization, respectively.

This paper presents the synthetic methodologies, detailed characterization, and renewable atom content analyses for these polymers. By evaluating their structural, optical, electrochemical, and thermal properties, we aim to highlight the potential of those biosourced materials in reducing reliance on petroleum-based precursors while demonstrating interesting properties required for applications in organic electronics. Through this work, we demonstrate the feasibility of integrating renewable feedstocks into advanced materials development, paving the way for more sustainable approaches to the synthesis of conjugated polymers.

Results and discussion

Knowing the different metal-free polymerization methods, we began investigating how to adapt some of these protocols to biosourced monomers to make useful conjugated polymers. Our search stopped on poly(phenylene vinylenes) (PPVs), which are a class of conjugated polymers that have been extensively studied and continue to be investigated for their applications in polymer light-emitting diodes (PLED) devices. ²² They have also been synthesized by a few metal-free polymerization methods (Gilch, Knoevenagel condensation, Horner–Wadsworth–Emmons reaction). ²³ As shown in Scheme 1, the Gilch polymerization requires a monomer bearing two halomethyl groups, the Knoevenagel condensation involves the nucleophilic addition of a carbon atom bearing acidic hydrogen atoms to a carbonyl group and the Horner–Wadsworth–Emmons reaction is the condensation of a stabilized phosphonate carbanion on an aldehyde.

We oriented our search towards suitable starting materials that can be obtained from biomass. A publication from Short *et al.* describes that succinic esters could be dimerized and further aromatized into dimethyl 2,5-dihydroxyterephthalate, which

features two ester groups.²⁴ These ester groups could then be converted into a variety of functional groups (halomethyl groups, aldehydes groups, nitrile groups, *etc.*) suitable for different metalfree polymerization protocols. However, we decided to avoid the Horner–Wadsworth–Emmons reaction as it required more synthetic steps to reach the targeted compound than the Gilch and Knoevenagel condensation.

The detailed synthetic procedures can be seen in the ESI,† as well as the ¹H and ¹³C NMR spectra of new intermediate compounds in Fig. S16–S32 (ESI†).

Synthesis of biosourced poly(phenylene vinylene) (bPPV)

The first polymer we aimed to synthesize was polymer bPPV using the Gilch polymerization method.²⁵ The synthesis of the required dihalomethyl benzene monomer and further polymerization to bPPV is illustrated in Scheme 2. The first step involves the preparation of compound 1 (96% yield) via the Claisen condensation of succinic ester using NaH as reagent.²⁶ Compound 1 is then aromatized into compound 2 using Nchlorosuccinimide (NCS) as the oxidant, achieving a 85% vield.²⁴ The resulting yellow solid was alkylated with 2ethylhexyl bromide in the presence of K2CO3 to produce compound 3 with a 60% yield.²⁷ Next, the ester groups present in compound 3 were reduced to alcohol groups using LiAlH4, yielding compound 4 with a 96% yield.²⁸ The alcohol groups in compound 4 are subsequently converted to chlorines, thus forming the final monomer (compound 5) with an 80% yield.²⁹ Compound 5 was then polymerized via the Gilch pathway to obtain polymer bPPV.25 To prevent rapid gelation, tBuOK was added only after the solution of compound 5 in THF had been cooled to -78 °C, and the mixture was gradually heated to 50 °C. The reaction was quenched by precipitating polymer bPPV in MeOH at room temperature, followed by purification via Soxhlet extraction with MeOH, acetone, hexanes, and CHCl3. The yield of polymer bPPV in the CHCl₃ fraction was found to be 51%.

Synthesis of polymer biosourced poly(cyano-PPV) (bCN-PPV)

We then aimed for the synthesis of polymer **bCN-PPV** by the Knoevenagel condensation. CN-PPVs are also of interest for

Scheme 1 Synthesis of PPVs by Gilch polymerization, Knoevenagel condensation and Horner-Wadsorth-Emmons reaction (HWE)

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Scheme 2 Synthesis of polymer bPPV.

PLED application, as they have been shown to improve electron injection and charge captured at the interface in multilayer PLED devices which improves the overall device performance.³⁰ As illustrated in Scheme 3, we synthesized the dialdehyde compound 7 through the oxidation of compound 4 using pyridinium chlorochromate (PCC), achieving a 98% yield.²⁸ The dinitrile compound 6 was prepared by reacting dichloride

5 with NaCN, resulting in a 46% yield.³¹ With both the nucleophile and electrophile in hand, we followed the protocol described by Thompson et al. for the Knoevenagel condensation.31 Equimolar quantities of the monomers were mixed in anhydrous THF and anhydrous tBuOH, and the mixture was brought to reflux before adding 2.2 equivalents of tBuOK. Adding the base at high temperatures is crucial to

Scheme 3 Synthesis of polymer bCN-PPV

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minimize side reactions, such as the Thorpe-Ziegler reaction and Michael addition. 32,33 Upon completion of the Knoevenagel condensation, the polymer was precipitated in methanol at room temperature with the addition of two drops of AcOH and further purified via Soxhlet extraction using MeOH and acetone. The resulting polymer bCN-PPV was soluble in the acetone fraction and was obtained with a 92% yield.

Synthesis of polymer biosourced poly(thiazolo[5,4-d]thiazole) (bPPTzTz)

The synthesis of the dialdehyde compound 7, used in the synthesis of polymer bCN-PPV, also enabled the preparation of other conjugated polymers, such as thiazolothiazoles (TzTz).

TzTz units have shown great potential in organic electronic. Indeed, they are more electron deficient than their thiophene counterpart which improve the oxidation stability, and they also show strong π - π interaction at the solid state which generally improve charge carrier mobility. 34,35 As illustrated in Scheme 4, according to the protocol of Lolaeva et al., the dialdehyde was polymerized into polymer bPPTzTz using dithiooxamide in DMF under microwave heating.¹⁹ Initially, we used ethylhexyl side chains, but due to the poor solubility of the resulting material, we opted to increase their length to decyltetradecyl side chains. To achieve this, we modified the previously used procedure for synthesizing compound 7 to incorporate the longer side chains, successfully obtaining monomer 8 with a 75% yield over the three steps from compound 2. Dialdehyde 8 was then polymerized using the Ketcham polymerization method. The resulting polymer was precipitated in methanol and purified via Soxhlet extraction using methanol, acetone, and hexanes. The polymer dissolved in the hexanes fraction and after evaporation of the solvent, polymer bPPTzTz was obtained with a 90% yield.

Renewable atoms content

To evaluate the renewability of the biobased polymer obtained in this study, we performed calculations based on the method described by Dupont et al. 9 Specifically, we calculated both the %BioT, which represents the number of biosourced atoms divided by the total number of atoms in the polymer structure, and the %BioC, which excludes the alkyl side chains and considers only the atoms in the polymer core. Both values provide valuable insights: %BioT considers the entire polymer structure, including the side chains, which may not always be

Table 1 Renewable atom content of polymers bPPV, bCN-PPV and **b**PPTzTz

Polymer	$\%_{ m BioT}$	$\%_{\mathrm{BioC}}$		
bPPV	100	100		
bCN-PPV	96.9	86.7		
bPPTzTz	96.3	66.7		

biosourced, while $\%_{\text{BioC}}$ excludes the side chains to avoid an overestimation of renewability caused by simply increasing the length of the side chains.

In our case, the side chains were also biosourced. Both 2-ethylhexyl bromide and 11-(bromomethyl)tricosane were synthesized from their respective alcohols.³⁶ These alcohols, in turn, are produced via the Guerbet reaction using linear alcohols, which are commonly biosourced either from oils and fats (e.g., palm oil, coconut oil)^{37–39} or by fermentation.⁴⁰

As shown in Table 1, polymer bPPV consists entirely of atoms (100%) derived from a renewable source (succinic acid). For polymer bCN-PPV, slightly lower but still high percentages of 96.9% and 86.7% were achieved, with and without including the side chains, respectively. The decrease is due to the inclusion of the nitrile group necessary for the Knoevenagel polymerization, which is not biosourced. For polymer bPPTzTz, the percentages were 96.3% and 66.7%, with and without the side chains, respectively. This result is expected, as the dithiooxamide required for forming the thiazolothiazole ring in the Ketcham reaction is not biosourced. These findings highlight the importance of considering both metrics. While the %_{BioT} values for polymer bPPTzTz and polymer bCN-PPV are similar due to the longer side chains present in polymer bPPTzTz, the % metric reveals a significant difference. Polymer bPPTzTz contains 20% fewer biosourced atoms than polymer bCN-PPV when the side chains are excluded.

Polymer characterization and thermal properties

The polymers were first characterized using ¹H NMR, FTIR spectroscopy, and SEC analysis. The yields and SEC data are summarized in Table 2. Polymer bPPV was obtained from the CHCl₃ fraction with a yield of 51%, polymer bCN-PPV from the acetone fraction with a yield of 92%, and polymer bPPTzTz from the hexanes fraction with a yield of 90%. For polymer bPPV, the lower yield of 51% can be attributed to discarding both the hexanes fraction and the insoluble polymer remaining in the Soxhlet thimble after CHCl₃ washing.

Scheme 4 Synthesis of polymer bPPTzTz.

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Table 2 Polymerization yield, number-average molecular weights (M_n) ,

mass-average molecular weight (M_w) , polydispersity index (PDI), temperature of decomposition (T_d) and glass transition temperature (T_d) of all the polymers

Polymer	Yield (%)	$M_{\rm n}$ (kDa)	$M_{\rm w}$ (kDa)	PDI	T _d (°C)	T _g (°C)
bPPV	51	370	1300	3.5	395	80
bCN-PPV	92	7	21	3.0	400	35
bPPTzTz	90	12	16	1.3	370	—

The molecular weights of the polymers were determined using SEC analysis in 1,2,4-trichlorobenzene (TCB). Polymer **bPPV** shows a high $M_{\rm p}$ of 370 kDa and an $M_{\rm w}$ of 1300 kDa, resulting in a PDI of 3.5. Achieving a high molecular weight is often a problem in Gilch polymerization due to lack of solubility, 41 but the polymer obtained remained soluble in THF, 2-MeTHF, CHCl₃, and TCB.

For polymer bCN-PPV, synthesized via Knoevenagel condensation, the $M_{\rm p}$ was measured to be 7 kDa, and the $M_{\rm w}$ was 21 kDa, with a PDI of 3.0. These lower molecular weights align with literature reports and are likely due to side reactions such as the Thorpe-Ziegler condensation and Michael addition, which alter the monomer ratios and hinder polymer chain growth. 32,33 Lastly, for polymer bPPTzTz, synthesized using the Ketcham polymerization, a M_n of 12 kDa and a M_w of 16 kDa were obtained, giving a PDI of 1.3. These results are consistent with previous results in the literature.¹⁹ However, the molecular weight was likely limited by poor solubility in N,N'-dimethylformamide (DMF), as the polymer precipitated during the reaction.

Thermogravimetric analysis (TGA) and differential scanning calorimetry (DSC) were performed on each of the polymers (see Table 2 and Fig. S1-S4, ESI†). Polymers bPPV, bCN-PPV, and

bPPTzTz exhibited high thermal stability, with a 5% weight loss occurring at 395 °C, 400 °C, and 370 °C, respectively. DSC measurements were conducted on each sample at heating rates of 20 °C min⁻¹ and 5 °C min⁻¹ to scan for thermal transitions such as the glass transition temperature (T_g) and the temperature of fusion (T_f) , respectively. Polymer **bPPV** showed a T_{σ} of 80 °C, which is slightly higher than what is reported for similar polymer MEH-PPV that possesses a T_g of around 70 °C. 42,43 This increase may be due to the high molecular weight of bPPV. Polymer **bCN-PPV** displayed a T_g of 35 °C, which is consistent with literature.44 Lastly, no thermal transitions were observed for polymer bPPTzTz. it is worth noting that, at slower scan rate $(5 \, {}^{\circ}\text{C min}^{-1})$, no thermal transition corresponding to the $T_{\rm f}$ was observed for the three samples.

¹H NMR spectra of polymers **bPPV**, ⁴⁵ **bCN-PPV**, ⁴⁶ and bPPTzTz⁴⁷ (see Fig. 1 and Fig. S30-S32 in the ESI†) were analyzed and compared with similar reports in the literature. 45-47 For polymer bPPV, the signal at 7.55 ppm corresponds to the vinylene protons, while the signal at 7.23 ppm corresponds to the benzylic protons. Signals at 4.00 ppm and in the range of 1.85 to 0.92 ppm are attributed to the ethylhexyl side chains. For polymer bCN-PPV, the multiplet at 8.1-7.7 ppm, which integrates for three protons, corresponds to two protons on the benzene ring and one proton from the alkene group. The multiplet at 7.1-6.8 ppm corresponds to two protons on the other benzene ring. Signals between 4.2-3.6 ppm and 1.9-0.6 ppm are assigned to the alkyl side chains. Notably, signals at 6.56 ppm and 3.24 ppm have been attributed to the cis-conformation of polymer bCN-PPV in previous reports. 46 By integrating the signal at 3.24 ppm and comparing it to the signal at 4.2-3.6 ppm, the cis-conformation content

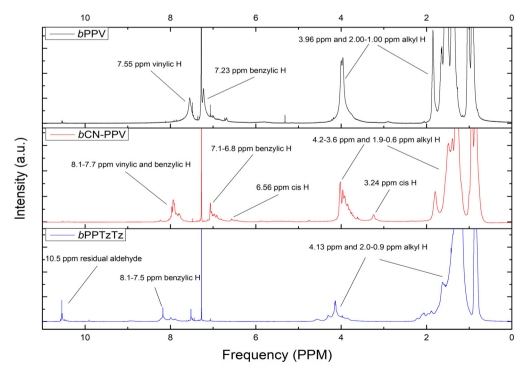


Fig. 1 NMR spectra of bPPV, bCN-PPV and bPPTzTz.

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was estimated to be approximately 8%. For polymer **bPPTzTz**, the signal at 10.5 ppm corresponds to residual aldehyde end groups. Signals in the 8.1-7.5 ppm region correspond to protons on the benzene ring, while signals at 4.13 ppm and 2.0-0.9 ppm are assigned to the alkyl side chains.

The analysis by FTIR spectroscopy provided key insights into the structural features of polymers bPPV, 48 bCN-PPV31 and **bPPTzTz.** 47,49 As shown in Fig. 2, for polymer **bPPV**, signals in the region of 2800-3100 cm⁻¹ correspond to sp² and sp³ C-H bond stretching, with the signal at 3060 cm⁻¹ attributed to the trans-vinyl C-H stretch. Signals around 1500-1350 cm⁻¹ correspond to semicircular phenyl stretching and alkyl CH2 stretching. The signals at 1033 cm⁻¹and 1195 cm⁻¹ are assigned to C-O stretching of the alkyl-O bond and the phenyl-O bond, respectively. Signals at 856 cm⁻¹ and 969 cm⁻¹ are attributed to out-of-plane wagging of the benzylic C-H and trans-vinyl C-H, respectively. For polymer bCN-PPV, signals in the 2800-3000 cm⁻¹ region correspond to sp³ and sp² C-H stretching. The signal at 2012 cm⁻¹ corresponds to nitrile stretching, which is shifted to a lower wavenumber compared to the monomer (2250 cm⁻¹) due to conjugation with the vinylene groups formed during polymerization. Signals around 1500-1350 cm⁻¹ correspond to semicircular phenyl stretching and alkyl CH₂ stretching. Signals at 1029 cm⁻¹ and 1205 cm⁻¹ are assigned to C-O stretching of the alkyl-O bond and the phenyl-O bond, respectively. For polymer **bPPTzTz**, sp³ and sp² C-H stretching signals were observed between 2800-3000 cm⁻¹. A signal at 1684 cm⁻¹ likely corresponds to residual aldehyde end groups, which are expected under the polymerization

conditions used. The signal at 1604 cm⁻¹ could correspond to the C=N stretching. Signals around 1400 cm⁻¹ are assigned to C=C stretching vibrations within the aromatic ring. The signal at 1211 cm⁻¹ corresponds to C-O stretching of the phenyl-O bond. Lastly, the signal at 1022 cm⁻¹ is assigned to a combination of both the C-O alkyl stretching and the C-S stretching in the thiazolothiazole ring.

Optical and electrochemical properties

The optical absorption spectra were measured for each sample, both in solution in CHCl3 and in thin films, which were spincoated from a chloroform solution. The optical band gap $(E_{g,optic})$ was calculated from the absorption onset in the thin film spectra. The $\varepsilon_{\mathrm{max}}$ was determined from a solution of the polymers in a known concentration following Beer-Lambert's law. The α_{max} was obtained from a spin-coated film of the polymers at a known thickness. The photoluminescence spectra were measured in solution in CHCl3. To obtain the fluorescence quantum yield ($\varphi_{\rm fl}$), we used the comparative method with a solution of rhodamine B in ethanol as our reference standard.⁵⁰ The electrochemical properties of the polymers were measured by cyclic voltammetry. The HOMO energy level was estimated from the onset of the oxidation wave, and the LUMO was determined from the onset of the reduction wave.⁵¹

The optical and electrochemical properties are shown in Table 3, Fig. 3 and in Fig. S5-S15 (ESI†). For polymer bPPV, the UV-Vis analysis and cyclic voltametric analysis showed an $\lambda_{\text{max sol}}$ at 510 nm and a slightly red-shifted $\lambda_{\text{max film}}$ at 520 nm, with an $E_{\rm g\,optic}$ of 2.1 eV and an $E_{\rm g\,elec}$ of 2.3 eV. At an excitation

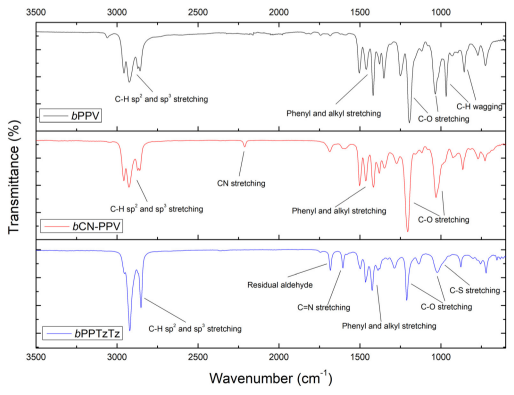


Fig. 2 FTIR spectra of bPPV, bCN-PPV and bPPTzTz.

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Table 3 Optical and electronical properties of the polymers

Polymer	λ _{max film} ^a nm	$\lambda_{\max sol}^b$ nm	$\lambda_{\mathrm{F}\max}^{c}$ nm	$E_{ m g optic}^{d}$ eV	E_{gelec}^{e} eV	${\varepsilon_{ m max}}^f \ 10^4~{ m M}^{-1}~{ m cm}^{-1}$	$\begin{array}{c} \alpha_{\max}^{\ g} \\ 10^4 \ cm^{-1} \end{array}$	HOMO eV	LUMO eV	$\frac{{{{arphi}_{ m{fl}}}^h}}{-}$
bPPV	520	510	555	2.1	2.3	2.6	15	-5.0	-2.7	0.49
b CN-PPV	450	450	555	2.3	2.3	1.7	9	-5.4	-3.1	0.20
b PPTzTz	510	500	510	2.0	2.1	1.8	8	-5.5	-3.4	0.31

^a Absorption maxima in thin films. ^b Absorption maxima in solution in CHCl₃. ^c Photoluminescence maxima in solution in CHCl₃. ^d Optical band gap from the thin film spectra. ^e Electronical band gap. ^f Molar absorption coefficient in CHCl₃. ^g Absorption coefficient in thin films. ^h Relative fluorescence quantum yield.

wavelength of 450 nm, polymer **bPPV** exhibited an $\lambda_{F \, max}$ of 555 nm and a $\varphi_{\rm fl}$ of 0.49. The HOMO and LUMO energy levels were estimated to be -5.0 eV and -2.7 eV, respectively. We also determined the ϵ_{max} to be 2.6 \times 104 $M^{-1}~cm^{-1}$ and the α_{max} to be 15 \times 104 cm $^{-1}$. This is consistent with data found in literature. 52-55 For polymer bCN-PPV, no shift in absorption maxima was observed between the solution and thin film spectra, both showing a λ_{max} of 450 nm, with an E_{goptic} of 2.3 eV and an $E_{\rm g\,elec}$ of 2.3 eV. At an excitation wavelength of 425 nm, an $\lambda_{F\,max}$ of 555 nm was obtained, resulting in a large Stokes shift of 105 nm, and a $\varphi_{\rm fl}$ of 0.20. The HOMO and LUMO energy levels were -5.4 eV and -3.1 eV, respectively. We also determined the ε_{max} to be 1.7 \times 104 M⁻¹ cm⁻¹ and the α_{max} to be 9 × 104 cm⁻¹. This is mostly consistent with data found in

literature, however, we did not observe a reversibility peak, so this result should be interpreted with caution. 52,53 It was noted that a lower $\varphi_{\rm fl}$ at 0.20 was obtained (0.35–0.4 in literature^{52,53}), however it is hard to point to a specific explanation for this result as results found in literature typically have different side chains, molecular weight and/or methodology to obtain the $\varphi_{\rm fl}$. For polymer **bPPTzTz**, an $\lambda_{\rm max\,sol}$ of 500 nm and a slightly red-shifted $\lambda_{max film}$ at 510 nm were observed, with an $E_{\rm g\,optic}$ of 2.0 eV and an $E_{\rm g\,elec}$ of 2.1 eV. Notable shoulders appeared at 460 nm and 540 nm on either side of the absorption maximum in the UV-Vis spectra of polymer bPPTzTz, which can be attributed to transitions at different vibrational levels from the ground state to the excited state. This effect is more pronounced in the photoluminescence spectra, where an

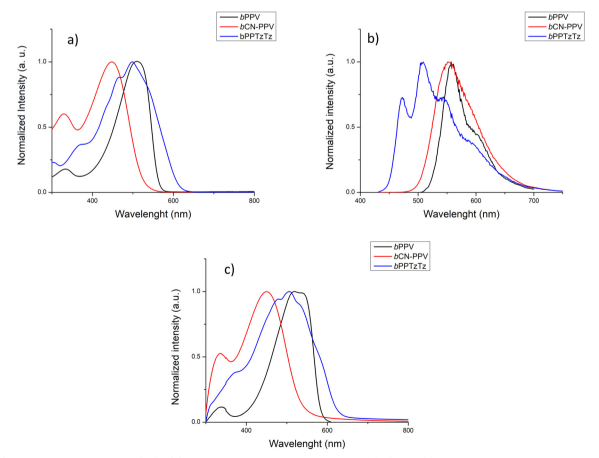


Fig. 3 (a) UV-Vis spectra in solution in CHCl₃, (b) photoluminescence spectra in solution in CHCl₃ and (c) UV-Vis spectra in thin films of polymers bPPV, bCN-PPV and bPPTzTz.

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 $\lambda_{\rm Fmax}$ of 510 nm was obtained, with additional peaks at 470 nm and 550 nm, resulting from transitions from the excited state to the ground state at different vibrational levels. A $\varphi_{\rm fl}$ of 0.31 was obtained, and the HOMO and LUMO energy levels were -5.5 eV and -3.4 eV, respectively. We also determined the ϵ_{max} to be 1.8 \times 104 M⁻¹ cm⁻¹ and the α_{max} to be 8 \times 104 cm⁻¹. There aren't precedents in the literature to compare the data obtained for bPPTzTz. Overall, the data highlight distinct optical and electrochemical characteristics for each polymer and show their potential for PLED applications.

Conclusions

In conclusion, this study demonstrates the viability of using forest biomass-derived materials, specifically succinic acid from cellulose, a renewable building block for the synthesis of conjugated polymers through three different metal-free polymerization methods. Three distinct polymers (bPPV, bCN-PPV, and bPPTzTz) were successfully synthesized, characterized, and evaluated in terms of renewable content, optical, electrochemical, and structural properties.

The results highlight the potential of biosourced materials in reducing reliance on petroleum-based precursors while maintaining relevant properties for applications such as transistors and PLEDs. Notably, polymer bPPV achieved 100% biosourced atom content and suitable optical properties, while polymers bCN-PPV and bPPTzTz demonstrated high renewable atom percentages with distinct electronic and structural characteristics. These findings underline the feasibility of incorporating renewable resources and sustainable synthesis approaches in the development of advanced materials for organic electronics. Future efforts should focus on improving the renewability aspect of the reaction condition of the monomer synthesis (NaH, SOCl2, PCC, DMF, halogenated solvents...), as the chemical industry reduces its dependence on petroleum-based, toxic and/or dangerous chemicals, but it is still marking a significant step toward the creation of fully renewable organic electronic devices.

Conflicts of interest

The authors declare no conflicts of interest.

Data availability

The data supporting the findings of this study (relative quantum yield study, synthesis of the monomers and polymers, UV-vis spectra of the polymer, photoluminescence spectra of the polymers, cyclic voltammograms of the polymers, thermogravimetric analysis and differential scanning calorimetry of the polymers as well as the ¹H and ¹³C NMR of the intermediates and the ¹H NMR of the polymers) can be found in the ESI.†

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