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Optimizing the vectorial component of first hyperpolarizabilities of push–pull chromophores to boost the electro-optic activities of poled polymers over broad telecom wavelength bands†

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The use of thiophene as a conjugation unit has long been studied for the design and synthesis of thermally stable donor– π –bridge–acceptor (D– π –A) chromophores for second-order nonlinear optics (NLO). Herein we report the rational molecular design of thiophene-bridge-based push–pull chromophores by alternating the sequence of π -conjugation units to boost the EO activities of poled polymers. Through facile synthesis, a thienyl group is directly attached to the 4-position of the *N,N*-dibutylaminophenyl donor to furnish a new 4-[5-(4-dibutylamino-phenyl)-thiophen-2-yl]-buta-1,3-dienyl donor-bridge component (**APTBD**) with strong tricyanovinylidihydrofuran (TCF) and CF₃-TCF acceptors. Two site-isolators of 4'-methoxyphenyl (MeOPh) and 4'-dimethylaminophenyl (DMA) groups were orthometrically attached to the middle of the primary D– π –A framework to study their role in changing the solubility and properties of chromophores. Through a systematic study of the linear and EO properties of this series of new chromophores and with the assist of DFT calculations, we find that alternating the sequence of π -conjugation units of thiophene-bridge-based chromophores can significantly affect the shape and dipole moment components of molecules, which are the key molecular parameters in determining the bulk NLO performance of this series of chromophores. We also find unexpected results of orthometric site-isolators in affecting the properties of these chromophores in terms of solubility, NLO property, and geometric isomerism. Our studies lead to remarkable performance improvement of these chromophores in their molecular hyperpolarizabilities and effective translation to large r_{33} values up to 106.1 pm V^{−1} at 1304 nm and 88.7 pm V^{−1} at 1541 nm in poled polymers.

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Introduction

Donor– π –bridge–acceptor (D– π –A) chromophores with large non-linear optical (NLO) responses are of intense research interest as the key-enabling molecular component for optical communications, information processing, sensing, and bioimaging applications.^{1–4} Over the past two decades, structure–property relationship analyses based on extensive theoretical and experimental studies have led to rational molecular design and synthesis of dipolar push–pull chromophores exhibiting very large molecular hyperpolarizabilities (β values), some of which have been used as the feed-stock of organic and polymeric electro-optic (EO) materials for high-performance photonic devices.^{5–12}

Although it is generally accepted that the molecular NLO properties of push–pull chromophores depend on the strength

of the donor and acceptor groups, as well as the nature of π -conjugation bridges, at the material level, the design of high-performance organic EO (OEO) materials requires that the molecular properties of chromophores such as dipole moment (μ), stability and solubility be optimized rather than the β values alone. For the electric field poling process as an essential step in materials processing, it remains less clearly understood how the molecular $\mu\beta$ values of chromophores can be effectively translated into large EO coefficients (r_{33} values) in poled films and devices.^{10–16}

Specifically, the first hyperpolarizability of push–pull chromophores is a third rank tensor that can be described using a $3 \times 3 \times 3$ matrix, which is usually dominated by one diagonal tensor term (β_{xxx}) along the donor–acceptor axis (denoted as the *x*-axis). However, in the experimental measurement of poled OEO films, instead of β_{xxx} , it is the β_{μ} , the magnitude of the irreducible part of the β tensor along the dipole vector of chromophores, that is responsible for the macroscopic EO activity (r_{33} values) of materials.^{17–19} This is similar to the data analysis of molecular hyperpolarizability measurement in solution through electric field induced second harmonic generation (EFISH). The recent

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development of large β chromophores is mostly based on push-pull structures with extended π -conjugation and geometry optimized 3D conformation of the donor and acceptor groups. These molecules would possess unique components of hyperpolarizability tensor and ground-state dipole moment vector in substantial magnitude, and thus they cannot be treated as a simple one-dimensional charge-transfer system for $\mu\beta$ analysis. There are a few studies calculating the β_{μ} , β_{xxx} , and total β tensor, which can be significantly different in magnitude by up to 50%.^{18,19} However, definitive experimental evidence substantiating the critical role of β_{μ} over that of the β_{xxx} term to the bulk NLO response of materials is still lacking, especially for the characterization of poled OEO films containing large β chromophores.

To better understand how the β_{μ} values of push-pull chromophores are related to the r_{33} values of materials, we hereby report the rational molecular design of thiophene-bridge-based push-pull chromophores by alternating the sequence of π -conjugation units to boost the EO activities of poled polymers (Chart 1). Using thiophene as a conjugation unit has long been studied for the design and synthesis of thermally stable push-pull chromophores for second-order NLO.^{20–29} A representative example is the dialkylamino-substituted thiophene stilbenes with strong TCF (tricyanovinyl) dihydrofuran and CF_3 -TCF acceptors, previously denoted as FTC chromophores.³⁰ Compared to the push-pull polyenes, FTC-type chromophores showed blue-shifted absorption

in the near-infrared (NIR) region and excellent optical transparency around 1300 nm and 1550 nm, respectively, applicable to both the second and third telecom windows for low loss waveguides. However, the reported certified r_{33} values of poled polymer waveguides containing these chromophores are mostly in the range of 40–60 pm V^{−1} at the telecom wavelengths, which were suboptimal leaving room for improvement.^{31–33}

Through a systematic study of the linear and EO properties of this series of new chromophores, the relationship between molecular and bulk NLO responses has been analyzed and compared with thiophene-stilbene-based chromophores as one of the benchmarks for EO devices. Through density functional theory (DFT) calculations, we find that alternating the sequence of π -conjugation units of thiophene-bridge-based chromophores can significantly affect the shape and dipole moment components of molecules, which are the key molecular parameters in improving the $\mu\beta_{\mu}$ product of chromophores for higher r_{33} values of poled polymers. We also find unexpected results of orthometric site-isolators in affecting the properties of these chromophores in terms of solubility, NLO property, and geometric isomerism, which is highly dependent on the strength and structural asymmetry of TCF and CF_3 -TCF acceptors. Our study leads to remarkable performance improvement of these chromophores in their molecular hyperpolarizabilities and very effective translation to large r_{33} values up to 106 pm V^{−1} at 1304 nm in poled polymers. Furthermore, it demonstrates that optimizing the $\mu\beta_{\mu}$ product (instead of $\mu\beta_{xxx}$) as the chromophore figure of merit and increasing the loading density of chromophores are critically important in realizing the large bulk NLO response of OEO materials for performance breakthrough.

Results and discussion

Synthesis and structural characterization

Scheme 1 shows the synthetic routes for new push-pull chromophores based on the 4-[5-(4-dibutylamino-phenyl)-thiophen-2-yl]-buta-1,3-dienyl donor-bridge component (APTBD) and strong TCF or CF_3 -TCF acceptors. A generic forward scheme has been devised to synthesize the key intermediates of donor- π -bridge aldehydes, optionally being incorporated with the site-isolation groups, which were further condensed with TCF and CF_3 -TCF acceptors, respectively, to afford six new chromophores (APTBD-1 to APTBD-6) in good yields.

The donor precursor dibutyl-(4-thiophen-2-yl-phenyl)-amine (N-3) was prepared by a simple, two-step synthesis of bromination and Stille coupling. Compound N-3 was then converted to the aldehyde intermediate N-4 through Vilsmeier formylation, followed by base-catalyzed Knoevenagel condensation with 4-substituted phenylacetonitriles³⁴ and reduction with diisobutylaluminum hydride (DIBAL-H) to produce the APTBD donor-bridge aldehydes N-7 and N-8 bearing the site-isolators of 4'-methoxyphenyl (MeOPh) and 4'-dimethylaminophenyl (DMA) groups, respectively. Alternately, the lithiation of N-3, followed by electrophilic quenching with 3-dimethylaminoacrolein, produced the pristine APTBD bridge aldehyde N-9 without the isolation

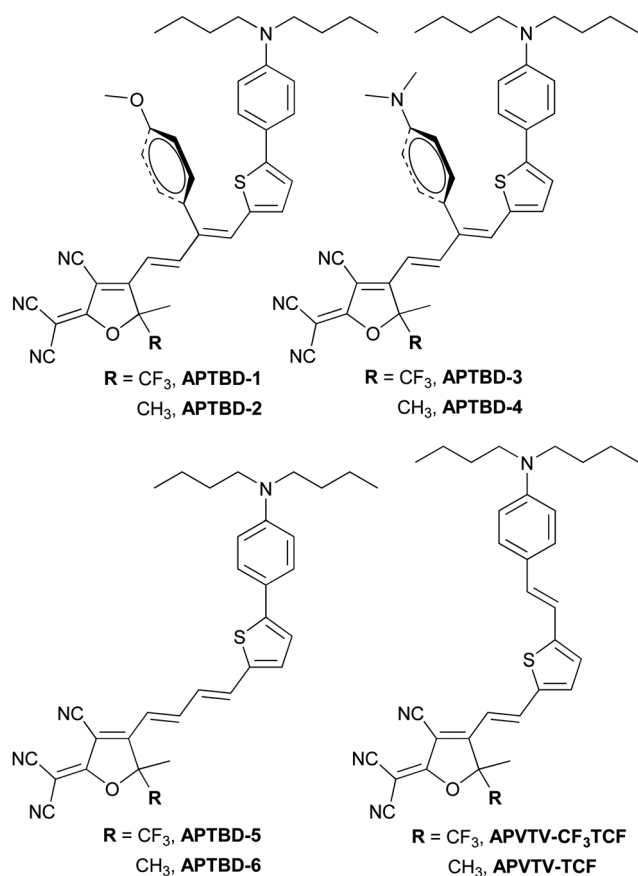
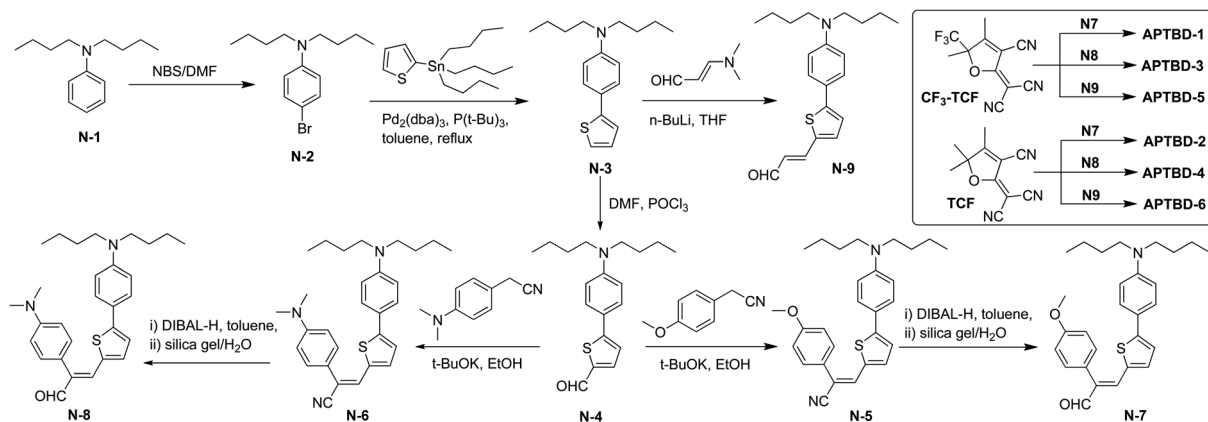


Chart 1 Molecular structures of thiophene-bridge-based push-pull chromophores in this study.





Scheme 1 Synthesis of thiophene-bridge-based push-pull chromophores in this study.

groups. Two benchmark chromophores based on the traditional π -conjugation structure of dialkylamino-substituted thiophene stilbenes, namely **APVIV- CF_3TCF** and **APVIV-TCF**, were also included for the comparison study.

All of the prepared chromophores were fully characterized by ^1H NMR, ^{13}C NMR, HRMS, UV-vis-NIR spectroscopy, and TGA as reported in the ESI.† A careful analysis of ^1H NMR spectra for these chromophores revealed an all-*trans* configuration of the protons of the olefinic backbones, except that **APTBD-6** with the TCF acceptor contains a *cis/trans* isomer mixture with a percentage ratio of 13%:87%. It suggests that the introduction of rigid isolation groups and the use of a stronger $\text{CF}_3\text{-TCF}$ acceptor would facilitate the formation of the all-*trans* rigid configuration of this series of D- π -A chromophores.

From TGA, these new **APTBD**-series chromophores show very high decomposition temperatures (T_d) greater than 270 $^\circ\text{C}$, which is nearly 30 $^\circ\text{C}$ higher than those of **APTPV** chromophores.³⁵ The comparison suggests that the relocation of π -conjugation units can improve the intrinsic stability of chromophores by impeding the potential pathways for bimolecular reaction mechanisms of thermal decomposition.

Solvatochromic properties of chromophores

In order to study the effect of π -bridge alternation on the intra-molecular charge-transfer (ICT) of chromophores, the UV-vis-NIR absorption spectra of these chromophores were measured in a series of solvents with different dielectric constants. As shown in

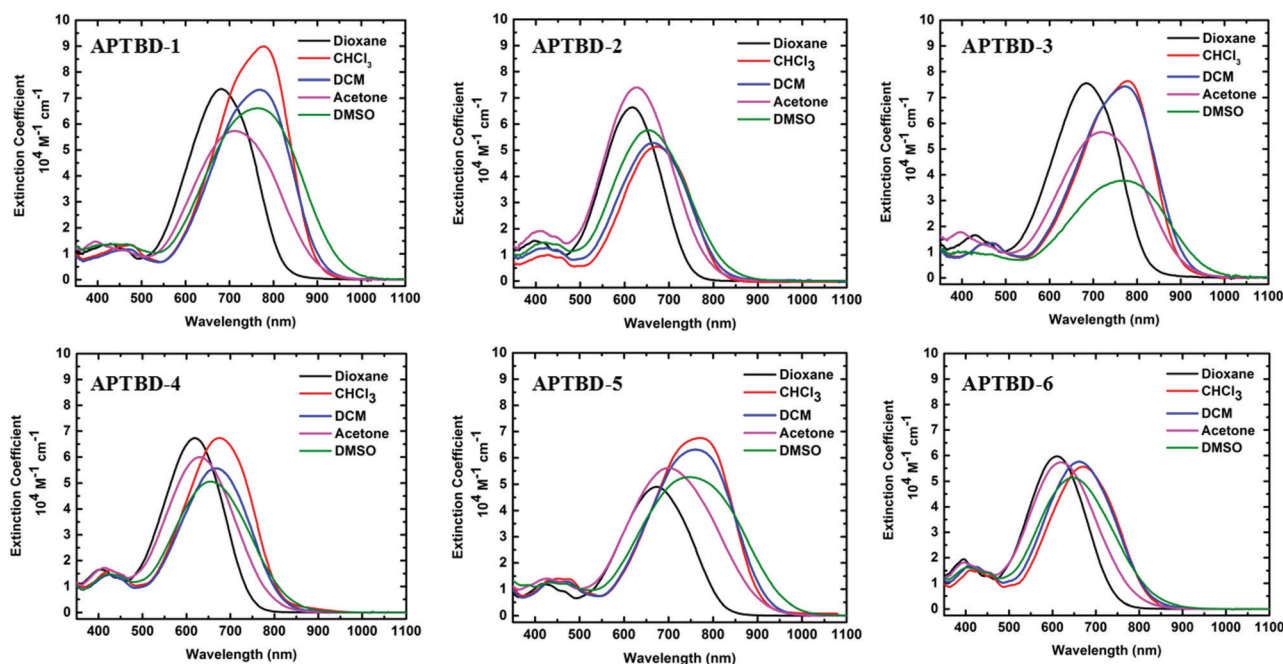


Fig. 1 UV-vis-NIR spectra of **APTBD**-series chromophores recorded in different solvents of variable dielectric constants at 25 $^\circ\text{C}$.



Fig. 1, all these chromophores exhibit highly solvatochromic ICT absorption bands in the visible to NIR region. Chromophores **APTBD-1**, **APTBD-3**, and **APTBD-5** have the maximum absorption (λ_{max}) around 775 nm in chloroform, red-shifted by 100 nm than those of **APTBD-2**, **APTBD-4**, and **APTBD-6**, respectively. It reflects a much stronger electron-withdrawing power of $\text{CF}_3\text{-TCF}$ than that of simple TCF. It is worth noting that the solvatochromism of **APTBD-5** is very similar to its constitutional isomer **APVTV-CF₃TCF** (Fig. S50, ESI[†]), suggesting a minor effect of π -bridge alternation on the ICT of chromophores.

The introduction of the site-isolating MeOPh group for **APTBD-1** and **APTBD-2**, or the DMA group for **APTBD-3** and **APTBD-4** contributed to minor red-shift of their λ_{max} values in the solution, as compared to the unsubstituted chromophores with the same donor and acceptor groups. The preliminary analysis of solvatochromism data suggests that the substitution of isolation groups to the π -conjugation bridge does not interfere with the efficient charge-transfer property of **APTBD**-series chromophores. Instead, the site-isolation groups can increase the rigidity of the π -conjugation bridge, leading to increased molar extinction coefficients of chromophores, especially for **APTBD-1** and **APTBD-2**.

Electrochemical properties and density functional theory (DFT) study of chromophores

As shown in Fig. 2, all chromophores exhibited one reversible oxidative wave with a half-wave potential, $E_{1/2} = 0.5(E_{\text{ox}} + E_{\text{red}})$, versus ferrocene/ferrocenium (Fc/Fc^+) at about 0.26 V for **APTBD-1**, 0.23 V for **APTBD-2** and **APTBD-3**, 0.20 V for **APTBD-4**, 0.27 V for **APTBD-5** and 0.24 V for **APTBD-6**. The gradual decrease of onset oxidation potential in the series indicates slightly increased electron-donating strength from two unsubstituted chromophores (**APTBD-5** and **APTBD-6**), to two substituted with the MeOPh group (**APTBD-1** and **APTBD-2**), and further to two substituted with the DMA group (**APTBD-3** and **APTBD-4**), for two series of chromophores containing $\text{CF}_3\text{-TCF}$ and TCF acceptors, respectively.

A quasi-reversible reduction wave was also recorded for chromophores, and their HOMO and LUMO levels were then calculated from their corresponding oxidation and reduction potentials. We found that the HOMO levels of these chromophores were at around -5.0 eV to -5.07 eV, typical for NIR absorbing D- π -A chromophores containing the 4-*N,N*-dialkylaminophenyl donors. Meanwhile, the corresponding LUMO levels of these chromophores were located at around -4.0 eV for **APTBD-1**, **APTBD-3**, and **APTBD-5**, and at around -3.8 eV for **APTBD-2**, **APTBD-4**, and **APTBD-6**. The results are consistent with the solvatochromic study, reiterating the stronger electron-withdrawing power of $\text{CF}_3\text{-TCF}$ than that of TCF.

To obtain a better understanding of these molecules, the geometrical optimization and orbital distribution of ground state for these molecules were conducted using the Gaussian 09 package at the level of B3LYP/6-31G(d,p) (Fig. 3). All molecules were assumed to be in a *trans* configuration, and their geometries were optimized in the gas phase and given solvents. It is shown that the electronic distribution of HOMOs is delocalized

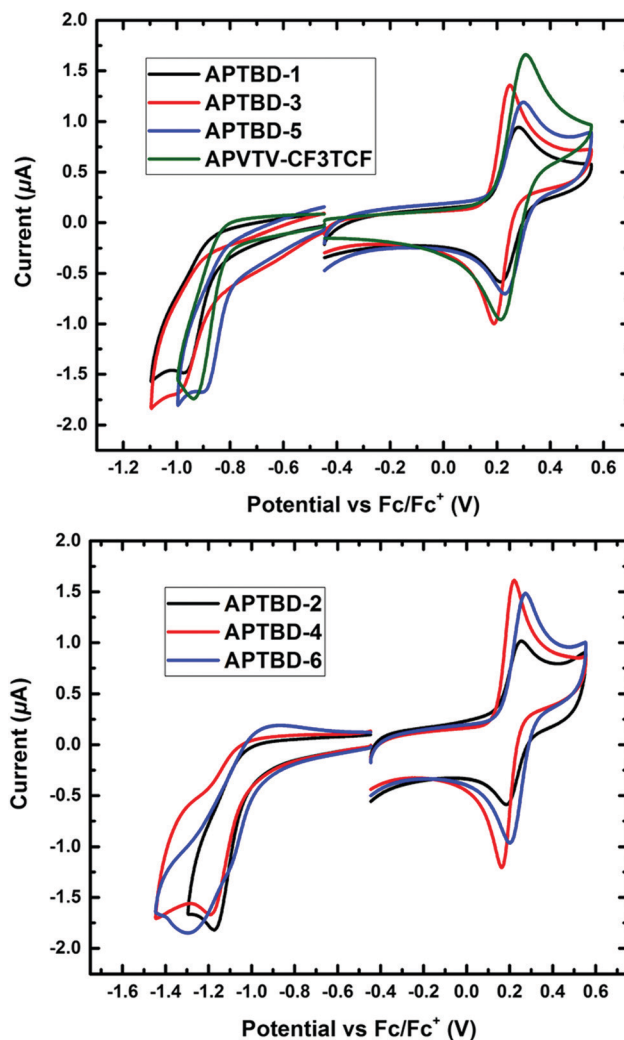


Fig. 2 Cyclic voltammetry (CV) measurements of chromophores recorded in dichloromethane solutions containing 0.1 M Bu_4NPF_6 supporting electrolyte.

over the (4-dibutylamino-phenyl)-thiophen-2-yl moiety, whereas the LUMOs are mainly constituted by the dienyI acceptor moieties. The primary D- π -A frameworks of all chromophores are highly coplanar, while the **APTBD**-series chromophores give smaller bending angles from the (4-dibutylamino-phenyl)-thiophen-2-yl moiety to the dienyI acceptors than the **APVTV**-based chromophores. For chromophores (**APTBD-1** to **APTBD-4**) with isolation groups, the optimized ground-state geometries of chromophores show that the MeOPh or DMA groups be orthometrically attached to the middle of D- π -A structures (Fig. S54, ESI[†]). The dihedral angles between the phenyl ring of MeOPh or DMA groups and the D- π -A plane are around 85° , and interestingly minor participation could be seen from the substitution groups to their HOMOs.

Overall, the trend in the changes of HOMO and LUMO energy levels for these chromophores is consistent from DFT analysis to CV measurements and agrees well with the data of optical bandgap (λ_{onset} and E_g) from the optical absorption of chromophores. Such analyses can be used as a guide to explain



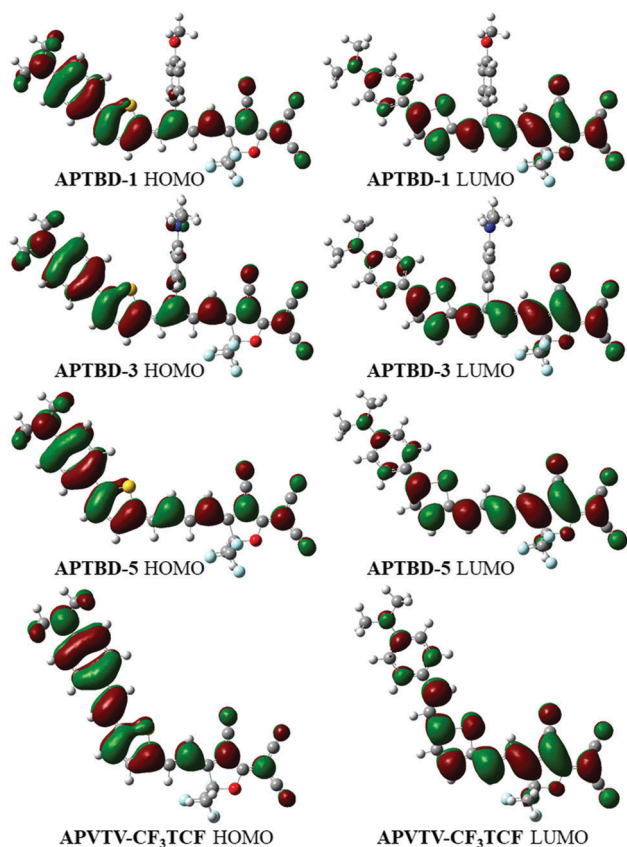


Fig. 3 Optimized structures and frontier molecular orbital diagrams of chromophores with the CF₃-TCF acceptor by DFT with the Gaussian basis set of B3LYP/6-31G(d,p) in gas phase.

the results in the study of solvatochromic and electrochemical properties of chromophores (Table 1).

Study of the poling-induced birefringence, EO properties and rigid oriented gas model analysis for poled films

Following the standard formulation and processing protocols of EO polymers, the optical quality thin films of guest–host polymers containing a loading level of 20 wt% of chromophores were prepared, except only 10 wt% for **APTBD-6** due to its poor solubility. Poly(styrene-*co*-methyl methacrylate) (P(*S-co*-MMA)) is selected as the host polymer due to its excellent optical and

dielectric properties and good compatibility with TCF-based chromophores. The modest number density is used in the study, and the films were poled under an electric field of 100 V μm^{−1}. This poling protocol is applied such that all films can be poled efficiently to the optimal temperature around the glass transition temperature (*T*_g) of polymers, which is essential for systematic structure–property relationship studies.

The *r*₃₃ values and poling-induced optical birefringence of all poled films were measured in slab waveguide geometry on a widely used prism-coupler system Metricon 2010/M. Without any encapsulation, the poled films showed excellent temporal stability in the handling of slab waveguide measurements under ambient conditions. The *r*₃₃ values of poled films were calculated by measuring the change of refractive indices under the application of modulation field *E* according to the Pockels effect:^{36,37}

$$\Delta\left(\frac{1}{n_{\text{TM}}^2}\right) = r_{33}E \quad (1)$$

The order parameters of poled films were calculated from the refractive indices of *n*_{TE} and *n*_{TM}, and used as a measure of chromophore alignment in the poled films. A Sellmeier fit of refractive indices was conducted on the values of refractive indices at 1304 nm and 1541 nm using a simple model of absorption and dispersion, in which the refractive index (*n*) is assumed to have contributions from both a non-resonant background *n*₀ and the two-level charge transfer resonance at frequency *ν*₀ in cm^{−1}.³⁸ Thus, at an operating frequency of *ν*, we have

$$n = n_0 + \frac{A}{\nu_0^2 - \nu^2} \quad (2)$$

The order parameter *Φ* for poled EO films is given by

$$\Phi = \frac{\delta_{\text{TM}} - \delta_{\text{TE}}}{\delta_{\text{TM}} + 2\delta_{\text{TE}}} \quad (3)$$

where *δ*_{TM} = *n*_{TM} − *n*₀ and *δ*_{TE} = *n*_{TE} − *n*₀. The obtained results for order parameters and *r*₃₃ values are summarized in Table 2. Since the refractive indices are measured at two wavelengths away from the main ICT absorption peak of chromophores, this characterization technique is more reliable than the absorbance measurement in evaluating the poling efficiency of high vis-NIR absorbing poled EO films.

With the modest loading level of chromophores, it is reasonable to use a rigid oriented gas model (ROGM) to analyze the orientational order of dipolar chromophores from poling-induced refractive index changes of poled EO films. This is a simplified semi-empirical model that laid a solid foundation for the field of poled polymers. It encompasses a variety of molecular and material parameters for reliable analyses without overly involving extensive computational and experimental analyses. It has been used in our research to achieve consistent results. According to ROGM, we correlate the measured *Φ* values as the second-rank axial order parameter to the non-centrosymmetric order parameter ⟨cos³θ⟩ of poled films, where θ is the angle between the permanent dipole moment of chromophores and applied electric

Table 1 The analysis of the HOMO and LUMO levels and bandgap by DFT and CV

	DFT (gas phase)			CV in CH ₂ Cl ₂		
	HOMO (eV)	LUMO (eV)	<i>E</i> _g (eV)	HOMO (eV)	LUMO (eV)	<i>E</i> _g (eV)
APTBD-1	−5.26	−3.20	2.06	−5.06	−3.97	1.09
APTBD-2	−5.21	−3.05	2.16	−5.03	−3.79	1.24
APTBD-3	−5.21	−3.14	2.07	−5.03	−3.97	1.06
APTBD-4	−5.14	−2.97	2.16	−5.00	−3.78	1.22
APTBD-5	−5.38	−3.35	2.04	−5.07	−4.02	1.05
APTBD-6	−5.32	−3.19	2.13	−5.04	−3.83	1.21
APVTV-CF₃TCF	−5.32	−3.32	2.00	−5.06	−4.02	1.04



Table 2 Summary of the poling-induced birefringence, EO properties and ROGM analysis for poled films

Chromophore	N (10^{20} cm^{-3})	$\lambda_{\text{max}}/\text{nm}$ in films	$n_{\text{TE}}/n_{\text{TM}}$ at 1304 nm	r_{33} at 1304 nm (pm V^{-1})	r_{33}/N	Φ	$\beta_{\mu}(-\omega; \omega, 0)$ (10^{-30} esu)
APTBD-1	1.94	725	1.5854/1.6291	53.8	27.7	0.160	2284
APTBD-3	1.90	737	1.5783/1.6107	53.4	28.1	0.107	2346
APTBD-5	2.30	716	1.6034/1.6632	78.9	34.3	0.146	2629
APTBD-5	2.87	719	1.6149/1.7092	106.1	36.9	0.190	2646
APVTV-CF ₃ TCF	2.30	722	1.5962/1.6419	60.8	26.4	0.098	2405
APTBD-2	2.11	648	1.5790/1.5937	26.5	12.6	0.066	1289
APTBD-4	2.06	654	1.5767/1.6079	28.3	13.7	0.142	1056
APTBD-6	1.27	639	1.5494/1.5632	18.1	14.3	0.072	1365

field.^{39–41} Overall, the poled films of these chromophores give a high poling-induced polar order, comparatively large but highly variable r_{33} values at 1304 nm.

According to ROGM, the r_{33} value of poled films is given by

$$r_{33} = 2Nf_0f_{\lambda}^2\beta_{\mu}(-\omega; \omega, 0)(\cos^3 \theta)/n_{\text{TM}}^4 \quad (4)$$

where $\beta_{\mu}(-\omega; \omega, 0)$ is the vector part of β tensors along the dipole moment direction of chromophores, f_0 is the zero-frequency local field factor for the electric field (or Onsager local field correction), and f_{λ} is the local field factor for the lightwave field (or Lorentz–Lorentz local field correction), respectively.^{39–41} We follow the convention of local field factors according to

$$f_0 = \varepsilon(n^2 + 2)/(n^2 + 2\varepsilon) \quad (5)$$

$$f_{\lambda} = (n^2 + 2)/3 \quad (6)$$

where ε is the dielectric constant of guest–host polymers in P(S-co-MMA) from the impedance analyzer.

After quantifying all the key parameters including r_{33} values, refractive indices, loading density, local field factors, and polar order parameters ($\langle \cos^3 \theta \rangle$), our studies indicate that APTBD-based chromophores exhibit large $\beta_{\mu}(-\omega; \omega, 0)$ values at 1304 nm.‡ For three chromophores with TCF acceptors, the values are $1056 \times 10^{-30} \text{ esu}$ for APTBD-4, $1289 \times 10^{-30} \text{ esu}$ for APTBD-2, and $1365 \times 10^{-30} \text{ esu}$ for APTBD-6.³⁹ The values are nearly doubled for chromophores with a stronger CF₃-TCF acceptor, around $2300 \times 10^{-30} \text{ esu}$ for APTBD-1 and APTBD-2, and $2629 \times 10^{-30} \text{ esu}$ for APTBD-5. The $\beta_{\mu}(-\omega; \omega, 0)$ values of APTBD-series chromophores are among the highest non-resonant values for dipolar chromophores with the same conjugation length and donor/acceptor structures. At the given loading level of 20 wt% and poling condition, their r_{33} values, up to 53.4 pm V^{-1} to 78.9 pm V^{-1} for chromophores with the CF₃-TCF acceptor, are significantly higher than those from FTC-type chromophores in poled slab waveguide geometry by the EO measurement of the prism coupling system.

We found that our analyses of β values from poled films agree reasonably with reported results for similar compounds in the solutions by electric-field-induced second-harmonic generation (EFISH) or hyper-Rayleigh scattering (HRS). It reiterates that our EO characterization and quantitative structure–property relationship analysis in guest–host solid–solid solutions is highly

applicable to evaluate the molecular and materials' NLO properties at the off-resonance wavelengths.^{39–42}

It has been suggested that the relative merits of NLO chromophores are consistently predictable with common DFT methods.^{17–19} To investigate the critical role of β_{μ} over that of the β_{xxx} term to the bulk NLO response of materials, we select two chromophores that are constitutional isomers, namely APTBD-5 and APVTV-CF₃TCF, for further investigation. We noticed that, at a loading density of $2.30 \times 10^{20} \text{ cm}^{-3}$, the r_{33} value of poled films with APTBD-5 was up to 78.9 pm V^{-1} at 1304 nm and improved by 30% over that of APVTV-CF₃TCF. Based on the fully optimized geometry of chromophores in the gas phase, we calculated their static first hyperpolarizabilities at the cam-B3LYP level by employing the split valence 6-31G(d,p) basis set using the “polar” keyword in the Gaussian 09 program package, which gives ten of the unique components of hyperpolarizability tensor. For comparison with experiments, various scalar measures of the tensors need to be computed according to

$$\beta_{\text{total}} = (\beta_x^2 + \beta_y^2 + \beta_z^2)^{1/2} \quad (7)$$

where the x , y and z components of the hyperpolarizability tensor are

$$\beta_i = \beta_{iii} + \frac{1}{3} \sum_{j \neq i} (\beta_{ijj} + \beta_{jji} + \beta_{jii}), \quad i, j \in (x, y, z)$$

The static β_{μ} as the vector part of β tensors along the dipole moment direction of chromophores was then calculated by

$$\beta_{\mu} = \sum_{i=x,y,z} (\mu_i \beta_i) / \left(\sum_{i=x,y,z} \mu_i^2 \right)^{1/2} \quad (8)$$

$$\mu_{\text{total}} = (\mu_x^2 + \mu_y^2 + \mu_z^2)^{1/2} \quad (9)$$

where μ_x , μ_y , and μ_z are three components of the ground-state dipole moment vector of chromophores from the geometry optimization.^{17–19} These results are summarized in Table 3.

From the DFT analysis, APTBD-5 possesses a μ_{total} of 21.9 D and a β_{μ} of $617.0 \times 10^{-30} \text{ esu}$. Its nearly identical β_{μ} and β_{xxx} values suggest an excellent overlap between the dominant β tensor term and the dipole moment of chromophores, mainly due to the relatively small bending angle along with the majority of its D– π –A framework. In comparison, APVTV-CF₃TCF has a μ_{total} of 20.6 D and a β_{μ} of $572.6 \times 10^{-30} \text{ esu}$, the latter being considerably smaller than its β_{xxx} . The angular shape of APVTV-CF₃TCF reduces

‡ The values of hyperpolarizabilities (β) in MKS unit are converted to the values in electrostatic units (esu) according to the unit conversion: $1 \text{ statvolt}^{-1} \text{ cm}^4 = 4.18888 \times 10^{-10} \text{ m}^4 \text{ V}^{-1}$.



Table 3 Summary and figure-of-merit analysis of components of the ground-state dipole moment vector and static first hyperpolarizability tensors of chromophores by DFT

Chromophore	μ (D)				β ($\times 10^{-30}$ esu)					$\mu_{\text{total}}\beta_{\mu}$ ($\times 10^{-48}$ esu)	r_{33} at N of $2.30 \times 10^{20} \text{ cm}^{-3}$
	μ_x	μ_y	μ_z	μ_{total}	β_x	β_y	β_z	β_{total}	β_{μ}		
APTBD-5	−20.8	−6.9	−1.5	21.9	652.7	−0.2	−14.5	652.9	−617.0	13 523	78.9
APVTV-CF₃TCF	−18.6	−8.7	−1.8	20.6	633.0	5.7	−12.2	633.2	−572.6	11 774	60.8

the overlap between the principle β term and μ . The subtle differences in the components of ground-state dipole moment vector of chromophores lead to a substantial improvement of the $\mu_{\text{total}}\beta_{\mu}$ value of $13\,523 \times 10^{-48}$ esu for **APTBD-5** over that of $11\,774 \times 10^{-48}$ esu for **APVTV-CF₃TCF**. The DFT analysis follows the trend of experimental results reasonably well and clearly verifies the $\mu_{\text{total}}\beta_{\mu}$ product as the figure of merit in determining the NLO activity of chromophores in poled polymers.

One of our initial motivations in the design of chromophores **APTBD-1** to **APTBD-4** is to follow our earlier study in a shorter D- π -A structure to test the feasibility of such site-isolation groups in large D- π -A chromophores for performance improvement.^{15,26} However, we obtained mixed results for **APTBD**-series chromophores with isolation groups. For chromophores with the TCF acceptor, we found that **APTBD-2** and **APTBD-4** chromophores showed higher solubility and better compatibility with the host polymer for higher r_{33} values than that of **APTBD-6**. In sharp contrast, all chromophores with the CF₃-TCF acceptor, namely **APTBD-1**, **APTBD-3**, and **APTBD-5**, are highly soluble in organic solvents and compatible with the host polymer up to a high loading level. The comparison underscores the critical role of the trifluoromethyl group in chromophores containing the CF₃-TCF acceptor over the TCF-based analogs, not only increasing the electron-withdrawing power of the acceptor for higher β but also creating a small z-component (μ_z) of dipole moment. It can effectively prevent close packing of D- π -A chromophores with predominant μ_x and μ_y components.^{43–47}

As shown earlier, the introduction of the site-isolating MeOPh group for **APTBD-1** and **APTBD-2**, or the DMA group for **APTBD-3** and **APTBD-4** contributed to a slight red-shift of their λ_{max} values and higher molar extinction coefficients in the solutions, as compared to the unsubstituted chromophores **APTBD-5** and **APTBD-6**, respectively. However, in the study of poled films, the unsubstituted chromophores gave better EO properties than their substituted ones. We speculate that the performance of these chromophores may be affected by some factors such as condensed phase effects and vibrational contributions. This is beyond the scope of this work and may deserve further study.

Finally, we found that **APTBD-5** is highly compatible with the host polymer P(S-co-MMA) through solution processing, and the optical bandgap of solid films is around 1000 nm, which suffice for achieving high optical transparency at broad NIR wavelengths (Fig. S54, ESI†). At a higher loading level of $2.87 \times 10^{20} \text{ cm}^{-3}$ with efficient poling, very large r_{33} values, up to 106.1 pm V^{-1} with $n_{\text{TE}}/n_{\text{TM}}$ of 1.6149/1.7092 at 1304 nm and 88.7 pm V^{-1} with $n_{\text{TE}}/n_{\text{TM}}$ of 1.6000/1.6725 at 1541 nm, have been achieved in the poled films. The values are among the highest reported non-resonant r_{33} values at two major telecom

wavelengths in poled slab waveguides and comparable to the performance from push-pull tetraene chromophores.^{33,37,43} Given the blue-shifted λ_{max} at 719 nm and the band edge of the **APTBD-5** chromophore in the series, the results are extremely encouraging for the future development of low-loss waveguide modulators at both the second and third telecom windows.

Conclusions

We report the synthesis and systematic study of the linear and EO properties of new thiophene-bridge-based push-pull chromophores by alternating the sequence of π -conjugation units to boost the EO activities of poled polymers. The relationship between molecular and bulk NLO responses of these chromophores has been analyzed and compared with thiophene-stilbene-based chromophores as one of the benchmarks for EO devices. Through ROGM and DFT calculations, we find that alternating the sequence of π -conjugation units of thiophene-bridge-based chromophores can significantly affect the dipole moment components of molecules, which are the key molecular parameters in improving the $\mu_{\text{total}}\beta_{\mu}$ product of chromophores for higher r_{33} values of poled polymers. We also obtained counter-intuitive results of orthometric site-isolators in affecting the properties of these chromophores in terms of solubility, NLO property, and geometric isomerism, which is highly dependent on the strength and structural asymmetry of TCF and CF₃-TCF acceptors.

Our study leads to the remarkable performance of new **APTBD**-based chromophores in their molecular hyperpolarizabilities and very effective translation to large r_{33} values up to 106.1 pm V^{-1} at 1304 nm and 88.7 pm V^{-1} at 1541 nm in poled polymers, which is over 30% better than the optimal results from thiophene-stilbene-based chromophore isomers. It demonstrates that optimizing the $\mu_{\text{total}}\beta_{\mu}$ product as the figure of merit of the chromophore and increasing their loading density are critically important in realizing the large bulk NLO response of OEO materials for photonic applications.

Experimental section

Materials and methods

All chemicals were purchased from Energy Chemical or Aldrich and used as received unless otherwise mentioned. High-resolution mass spectrometry (HRMS) was performed using a Thermo Scientific Q Exactive mass spectrometer. For the formulation of EO polymers, the solvents dibromomethane (DBM) and 1,1,2-trichloroethane (TCE) were distilled prior to use. The cyclic voltammetric data were measured using an Electrochemical



Analyzer (CHI 750) with Ag/AgCl as the reference electrode, a platinum wire as the working electrode, platinum gauze ($5 \times 5 \times 0.3$ mm) as the counter electrode and 0.1 M tetrabutylammonium hexafluorophosphate (TBAPF) as the electrolyte in dichloromethane. Thermogravimetric analyses (TGA) were carried out on a PerkinElmer STA 6000 Simultaneous Thermal Analyzer at a heating rate of $20^\circ\text{C min}^{-1}$. The films used in absorption spectrum and EO measurements were spin-coated on glass or ITO glass substrates with SPIN-PROCESS CONTROLLER. The dielectric constants of EO films were measured at low frequencies using a KEYSIGHT E4990A Impedance Analyzer. The UV-vis-NIR spectra of chromophores were recorded using an Ultra-Violet Visible Scanning Spectrophotometer (Shimadzu 1700) and an Ultra-Violet Visible Near Infra-red Spectrophotometer with Integrating Sphere (PE Lambda 750). DFT calculations using the Gaussian 09 package were carried out at the level of B3LYP/6-31G(d,p) for ground-state geometry optimization and CAM-B3LYP/6-31G(d,p) for static hyperpolarizability calculations.

Poling and measurement of r_{33} values and refractive indices

For studying the EO property derived from the chromophores, guest-host EO polymers were formulated by mixing chromophores at a given loading density into the host polymer poly(styrene-co-methyl methacrylate) (P(S-co-MMA)) in the solvent DBM (or TCE). The resulting solutions were filtered through a $0.2\ \mu\text{m}$ PTFE filter and spin-coated onto indium tin oxide (ITO) glass substrates. After soft baking, the films of doped polymers were baked in a vacuum oven at $60\text{--}70^\circ\text{C}$ overnight to ensure the complete removal of residual solvent. The thicknesses of the films were measured using a DektakXT Stylus Profiler and further confirmed on the subsequent optical measurement for refractive indices at the wavelengths of 1304 nm and 1541 nm using a commercial prism-coupler system (Metricon 2010/M). Then using the Desk V HP Sputter Unit (Denton Vacuum LLC), a thin layer (~ 20 nm) of semi-transparent gold was sputtered onto the films as the top electrode for contact poling and the subsequent EO coefficient measurement. The electric field poling of films was conducted at a central processor-controlled Mettler FP82 hot stage. The poling field was set at $100\ \text{V}\ \mu\text{m}^{-1}$ with the help of monitoring the LTC using a Keithley 2657A Source-Meter Unit. The optimal poling temperatures were around $110\text{--}115^\circ\text{C}$. After the poling, the refractive indices and r_{33} values of poled films were measured using the attenuated total reflectance (ATR) method in slab waveguide geometry on Metricon 2010/M, in which modulation voltages were applied to the poled films for EO coefficient measurements. The accuracy of the measurement system is verified with standard EO materials including lithium niobate thin film from NANOLN and AJLZ53/P(S-co-MMA) film from our research group.

Conflicts of interest

There are no conflicts of interest to declare.

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